S.N.D.T. WOMEN'S UNIVERSITY MUMBAI

M. PHARM. SYLLABUS

SPECIALIZATION: PHYTOPHARMACY AND PHYTOMEDICINE

FOUR SEMESTER PROGRAMME

Effective from Academic Year 2014-15

The M. Pharm. (**Phytopharmacy and Phytomedicine**) course, a very unique one has been proposed to introduce in Maharashtra for the FIRST TIME at C.U. Shah College of Pharmacy, by SNDT Women's University after due sanction from the DTE, Govt. of Maharashtra and AICTE. The course is devised with a focus on the aptitude, talents and job potential for women in herbal drug industry and research and development institutes.

This four semester programme has the following specific features.

- Emphasis on modern analytical techniques like UV, spectroflurometry, infrared spectrophotometry, NMR, Spectrometry HPLC, X-ray diffraction analysis and spectral analysis.
- Thrust on good manufacturing practices, quality audits, documentation and validation with a view to create total quality consciousness in herbal drug industry
- 3) Herbal product development and Packaging courses designed to teach current trends in formulation of herbal pharmaceuticals and newer herbal drug delivery systems.
- 4) Understanding of Regulatory affairs, New Drug Application and patenting procedures for herbal products.
- 5) Students work on a research project for two semesters and submit a dissertation at the end of fourth semester for which they are evaluated by subject experts.
- 6) One month in plant training in industry to correlate theory with professional practice.
- 7) Guest lectures and seminars are routinely arranged where visiting faculty impart insights in state-of-art technology and current advances in herbal drug development.

ADMISSION CRITERIA:

Only Female candidates shall be admitted as per rules of Directorate of Technical Education, Maharashtra: (a) Non Sponsored Seats (b) Sponsored Seats (c) Seats for reserved Category candidates. Admission will be given on purely merit basis

to candidates with valid GPAT/CET score. Any additional rules prescribed by the AICTE or SNDT Women's University will be applicable.

PROGRAM OUTCOMES

M.PHARM IN PHYTOPHARMACY AND PHYTOMEDICINE

- This Master's program is designed in such a way that students will have a complete understanding of important research areas of herbal crude drugs.
- 2. The program will impart the students end to end knowledge of important aspects of herbal drug technology.
- 3. It will enable the students to build expertise in various disciplines of herbal medicines, which can be applied in the herbal industry in various departments like manufacturing, research and development, formulation development, drug discovery, quality control, regulatory affairs, intellectual property rights, scientific/medical writing, pharmacovigilance study, sales and marketing.

SCHEME: M.Pharm (Phytopharmacy and Phytomedicine)

Sem	Subject Code	Subject	Hrs/ Week TH	Hrs/ Week PR	Credits TH	Credits PR	Marks TH	Marks PR
First	S1-P&P-1	Modern Analytical Techniques I	4	8	4	4	100	100
	S1- P&P -2	Advanced Pharmacognosy and Phytochemistry I	4	8	4	-	100	100
	S1- P&P -3	TQM,Patent Regulation & Validation	4	-	4	4	100	-
	S1- P&P -4	Herbal Product Development I	4	-	4	-	100	-
		Total	16	16	16	8	400	200
Seco nd	S2- P&P -1	Modern Analytical Techniques II	4	8	4	4	100	100
nu	S2- P&P -2	Advanced Pharmacognosy and Phytochemistry-II	4	-	4	-	100	100
	S2- P&P -3	Herbal Product Development II	4	-	4	-	100	-
	S2- P&P -4	Ayurveda & Allied Plant based therapies	4	8	4	4	100	-
		Total	16	16	16	8	400	200
Thir d	S3 – P&P -1	Industrial Training	One Mon th		2		50	
	S3- P&P -2	Biological Evaluation	4	4	4	-	100	-
	S3- P&P -3	Computing & Statistics	4	4	4	-	100	-
	S3- P&P -4	Research Methodology	4	-	4	-	100	-

	S3- P&P -5	Research Seminar	2	-	2	-	50	-
	S3- P&P -6	Research Project	-	8	8	-	200	-
		Total	14	16	24	-	600	-
Four	S4 - P&P -1	Research Project	32	-	12	-	300	-
th								
		Research Colloquium	-	-	4	-	100	-
		& Viva	-	-	8	-	200	-
		Total		-	24	-	600	-
		Grand Total	78	48	80	16	2000	400

Examination Pattern for M. Pharm. in Herbal Drug Technology

Semester I

SR.	SUBJECT	Exam	Theo	Theory			Exam	Practicals			
NO		Dur.	Int.	Ext.	Total	Credits	Dur.	Int	Ext.	Total	Credits
1	Modern Analytical Techniques I	2	50	50	100	4	6	50	50	100	4
2	Advanced Pharmacognosy and Phytochemistry I	2	50	50	100	4	6	50	50	100	4
3	TQM,Patent Regulation & Validation	2	50	50	100	4	-	-	-	-	-
4	Herbal Product Development I	2	50	50	100	4	-	-	-	-	-

Semester I : Total credits = 24

Semester-II

SR.	SUBJECT	Exam	Theor	y			Exam.	Prac	cticals		
NO		. Dur.	Int.	Ext.	Total	Credits	Dur.	Int	Ext.	Tota	Credits
										1	
1	Modern Analytical	2	50	50	100	4	6	50	50	100	4
	Techniques-II										
2	Advanced	2	50	50	100	4	6	50	50	100	4
	Pharmacognosy										
	and										
	Phytochemistry-II										
3	Herbal Product	2	50	50	100	4	-	-	-	-	-
	Development II										
4	Ayurveda & Allied	2	50	50	100	4	-	-	-	-	-
	Plant based										
	therapies										

Semester II : Total credits = 24

Semester III

SR.	SUBJECT	Exam	Theo	Theory Exam Practicals							
NO		Dur.	Int.	Ext.	Total	Credits	Dur.	Int	Ext.	Total	Credits
1	Biological	2	50	50	100	4					
	Evaluation										
2	Computing &	2	50	50	100	4					
	Statistics										
3	Research	2	50	50	100	4	6	-	-	-	-
	Methodology										
4	Research	1	25	25	50	2	1	-	-	-	-
	Seminar										
5	Research	-	-	-	-	-	-	-	-	200	8
	Project										
6	Industrial Training				50	2					

Semester III: Total credits = 24

Semester IV

SR.	SUBJECT	Exam	Theor	У			Exam	Prac	ticals		
NO		Dur.	Int.	Ext.	Total	Credits	Dur.	Int	Ext.	Total	Credits
1	Research		200	200	400	12	-	-	-	-	-
	Project (Thesis)										
2	Colloquia		100		100	4		-	-	-	-
3	Viva	-		100	100	8		-	-	-	-
	Total		300	300	600	24					

Semester IV: Total credits = 24

Semester I+ II + III + IV = 96 Credits Course

Internal Assessment

Internal Assessment will be carried out in two Steps:

- (a) Formative Assessment that envisages deployment of variety of tools and technique for diagnostic and remedial assessment of students
- **Pen-paper Assessment** in the form of question bank for class tests, home assignments, oral tests, quizzes, and MCQs
- **Practical Assessment** with viva questions based on various skills involved in doing Experiments
- Individual/Group Assessment in the form of activities with questioning tool kit, role plays, seminars, symposium, presentations, group discussions, projects, surveys, interviews, campaigns, site visits, data handling and data interpretation
- (b) Summative Assessment that provides a large number of question types such as,
- VAS (very short answer)
- SA(short answer)
- LA (long answers)
- MCQs(multiple choice questions)
- HOTS(higher order thinking skill) questions

SR. NO.	HEAD	MARKS
1	Assignments	15
2	Presentation	10
3	Unit Test	25
	TOTAL	50

SEMESTER III & IV

Project and Thesis work

Every student for the degree of Master of Pharmacy shall be required to undertake a project involving methodical research under the supervision of an approved guide and submit three copies of the thesis, duly certified by the supervisor to the Head of the Department/Principal.

SEMESTER III

Research Project and Industrial training

The student should complete industrial training for one month during the course work. The research project will be evaluated as follows:

Research Project	Marks
Reference work	100
Experimental work	100
Total marks	200

Research seminar	Marks
Total marks	50

Industrial Training	Marks
Total marks	50

SEMESTER IV

Research Project and Thesis work

The research project will be evaluated as follows:

Internal Assessment

Head	Marks
Experimental work	75
Presentation/ communication	50
Result/ conclusion	75
Research Colloquium	100
Total marks	300

External Assessment

Head	Marks
Viva voce and external assessment	300

Semester I

M. Pharm (Phytopharmacy and Phytomedicine)

SEMESTER I

- 1. Analytical Techniques-I (Theory & Practical)
- 2. Advanced Pharmacognosy and Phytochemistry-I (Theory & Practical)
- 3. TQM and Patent Regulation and Validation Theory
- 4. Herbal Product Development-I Theory

SEMESTER II

- 1. Analytical Techniques-II (Theory & Practical)
- 2. Advanced Pharmacognosy and Phytochemistry-II (Theory & Practical)
- 3. Herbal Product Development-II Theory
- 4. Ayurveda and Applied plant based therapies

SEMESTER III

- 1. Biological Evaluation (Theory & Practical)
- 2. Computing & Statistics
- 3. Research Methodology

SEMESTER I

M.Pharm. S1-[P&P]-1: MODERN ANALYTICAL TECHNIQUES- I [Theory]

OBJECTIVES:

- To make students familiar with the principles of quantitative estimation using UV-visible spectrometry
- To enable students to use spectrometers with proper understanding
- To make students competent for the basic quality control requirements or needs of industries
- To make students familiar with the principles of selective quantitative estimation using instrumental methods
- To enable students to use spectrofluorometer with proper understanding

- To make students familiar with the principles of absorption and emission spectrometry
- To enable students to use atomic absorption spectrometer with proper understanding and make them competent for quality control activities of industry
- To enable students to use IR spectrometers with proper understanding

COURSE OUTCOME:

The subject aims to provide students with the necessary skills for,

- Learning basic concepts, principles and advanced analytical instrumental techniques such as UV, IR, Spectroflorimetry, X-ray diffraction, Atomic absorption and emission spectroscopy and electrophoresis, for identification, characterization and quantification of drugs.
- To make students competent for the basic quality control requirements or needs of the industries.

CONTENTS:

Module 1	UV & IR Spectrometry & Spectrofluorimetry 1	credit
Contents	Topics covered	hrs
	Ultraviolet-Visible Spectrometry	(5)
	General Principles of Spectrometry: Instrumentation, single and double beau	am
	UV spectrometers, Applications of UV spectroscopy, Fisher Woodward ru	les
	for calculation of λ max values. Introduction to Optical rotatory Dispersion a	ınd
	Circular Dichroism. Principle and applications of Derivative UV Spectrosco	ру
	Spectrofluorimetry	
	Principle, definition and types of luminescence, Resonance fluorescence an	ıd

Mechanism of fluorescence and phosphorescence, singlet and triplet states,	(3)
fluorescence, factors affecting fluorescence, intrinsic structure of a molecule	
and fluorescence, instrumentation and applications.	
• IR Spectrometry	
Principle, types of vibrations, Instrumentation Michelson interferometer	(5)
applications, various regions in IR spectrum and their use for	
characterization of functional groups. Interpretation of IR spectrum	
characterization of functional groups. Interpretation of he spectrum	

Module 2	Atomic Absorption And Emission Spectrometry, X-Ray Diffraction	1
	Thermal Methods of Analysis & Electrophoresis	credit
Contents	Topics covered	Hrs
	Atomic Absorption And Emission Spectrometry	
	Principle, Sample atomization techniques & Applications	(3)
	• X-Ray Diffraction Analysis	
	Principle, Bragg's Law, instrumentation, sources of X-rays, Applications	(3)
	Thermal Methods of Analysis	
	Thermogravimetry (TG), Differential thermal Analysis (DTA), Differential	
	Scanning Calorimeter (DSC) - Principles, technique, instrumentation and	
	applications including interpretation of data	(5)
	• Electrophoresis	
	Theory and principles, Zeta potential, classification, instrumentation,	
	moving boundary electrophoresis, Zone Electrophoresis (ZE),	
	Isoelectric focusing (IEF), Immuno-electrophoresis and applications of	(3)
	electrophoresis	

Module 3		1 credit	ţ
Contents	Topics covered		Hrs
	Radio analytical techniques used in pharmaceuticals: Isotopic dilution	on	(6)
	methods, Radioimmunoassay, ELISA etc		
	• Microscopy: SEM, TEM, cryomicroscopy, AFM, confocal microscopy		(5)

(2)
(4)

• Determination of Water

Importance of determination of water or moisture content. Various methods used for determination of water and moisture content

Module 4	Project and Seminar Presentation on some recent research /seminars	1 credit
	based on the above topics	
Assigned	1. A.H. Beckett and J.B. Stenlake, Practical Pharmaceutical chemistry,	Part
Reading/	1 & 2, Athlone Press, London.	
Reference	es 2. Pharmacopoeia of India,6 th Edition, Govt. of India, Ministry of Healt	th &
	welfare, 2010.	
	3. British Pharmacopoeia, General Medicine Commision, UK., 2011.	
	4. United State Pharmacopeia, 34 th Edition, Convention, Inc., Rockville	e, MD
	20852, 2011.	
	5. Vogel's Textbook of Quantitative Chemical Analysis, 6 th Ed	dition,
	Prentice Hall, 2000	
	6. Robert M. Silverstein, Francis X. Webster, David J. Kiemle,	2009.
	"Spectrometric identification of organic compounds". 7th Ed. John	Wiley
	& Sons	
	7. Pavia D. L., 2009. "Introduction to spectroscopy". 4th, Belmont CA	
	8. Munson & Munson, "Pharmaceutical analysis: modern methods".	edited
	by James W. Munson, New York: M. Dekker	
	9. Kenneth A. Connors, 2007. "A Textbook Of Pharmaceutical Ana	alysis"
	3rd Ed. Wiley India-wse	
	10. Jens Thuro Carstensen, 2001. "Advanced pharmaceutical solids" M	Marcel
	Dekker, New York	
	11. Joseph B. Lambert, Scott Gronert, Herbert F. Shurvell, David Lightn	ner,
	Robert Graham Cooks, 2010.	
	12. "Organic structural spectroscopy", 2nd Ed. Pearson Education, Limit	ted.

- 13. Principles of Instrumental Analysis: Douglas A. Skoog (Author), F. James Holler, Stanley R. Crouch, 6th edition, Publisher: Brooks Cole. 2006.
- 14. Instrumental Methods of Analysis: S. S. Mahajan, , Popular Prakashan Pvt. Ltd., Mumbai, 2010.
- 15. Quantitative Analysis of Drugs in Pharmaceutical Formulations: P.D. Sethi, 3rd edition, CBS Delhi. 2008.
- 16. Published articles pertaining to the learnt techniques in reputed journals like Analytical Chemistry, Analytical Communications, The Analyst, Indian Drugs, etc.

M.Pharm. S1-[P&P]-1: MODERN ANALYTICAL TECHNIQUES- I [Practicals] OBJECTIVES:

- To make students familiar with the principles of selective quantitative estimation using instrumental methods
- To enable students to use UV-spectroscopy and spectrofluorometer with proper understanding
- To make students familiar with the principles of absorption and emission spectrometry
- To enable students to use atomic absorption spectrometer with proper understanding and make them competent for quality control activities of industry
- To enable students to use IR spectrometers with proper understanding

COURSE OUTCOME:

The subject aims to provide students with the necessary skills for,

- Skills in selecting the suitable techniques for analysis of drugs and pharmaceuticals products.
- To apply the knowledge learnt in developing new analytical procedures of the research work.

CONTENTS:

Module 1	UV –Visible spectrometry 1 cr	edit
Contents	Experiments	(30)
	1 Calibration of UV -Visible spectrophotometer for absorption and	1 (4)
	wavelength	
	2 Determination of λmax of a compound by Fisher Woodward Rule	(4)
	3 Determination of molecular absorptivity or E ¹ % of a substance	(4)
	4 Determination of range of linearity in accordance with Beer- Lambert Lav	(6)
	5 Analysis of a single component system from crude drugs and herba	(6)
	formulations.	
	6 Analysis of a binary mixture by simultaneous equation method and	l (4)
	absorption ratio method.	
	7 Interpretation of ORD and CD spectrum.	(2)

Module 2	Spectrofluorimetry, DSC and electrophoresis 1	lcredit
Contents	Experiments	Hrs
	8 Plotting of absorption spectrum	(4)
	9 Plotting of emission spectrum	(4)
	10 Plotting of a standard curve for quinine sulphate	(4)
	11 Analysis of fluorescent compounds such as scopoletin, pigments	(6)
	12 Analysis of proteins using electrophoresis	(4)
	13 Determination of melting point & heat of fusion using DSC.	(4)
	14 Determination of glass transition temperature using DSC	(4)

Module 3	IR Spectrometry	netry 1 credi	
Contents	Experiments		Hrs
	17. Calibration of IR spectrometer with polystyrene film		(6)
	18. IR spectrum of a neat liquid		(6)
	19. Preparation of KBr pellet for any one solid sample		(6)

	20.	Preparation of a 'mull' for phytoconstituents with different functional	(10)	
		groups such as amine, nitro, aldehyde, keto, carboxylic, hydroxyl, etc.		
	21.	To identify impurity in the sample	(2)	

Assigned	1. A.H. Beckett and J.B. Stenlake, Practical Pharmaceutical chemistry, Part 1
Reading/	& 2, The Athlone Press, London, 2011.
References	2. Vogel's Textbook of Quantitative Chemical Analysis, 6 th Edition, Prentice
	Hall, 2000
	3. Spectroscopic identification of organic compounds. John Dyer, Willy, NY.
	4. Spectrometric Identification of Organic Compounds: R. M. Silverstein,
	Francis X. Webster and David Kiemle, 7th edition, 2005, Wiley
	Publication, NY.
	5. Instrumental Methods of Analysis: S. S. Mahajan, 2010, Popular Prakashan
	Pvt. Ltd., Mumbai.

M.Pharm. S1-[P&P]-2: ADVANCED PHARMACOGNOSY AND PHYTOCHEMISTRY-I [Theory]

OBJECTIVES:

- To learn various quality control parameters to test quality of crude drugs from natural origin.
- Identify and quantify adulterations in herbal crude drugs,
- To impart insight on WHO guidelines required for cultivation, collection and quality control of herbal drugs.
- To make students understand pharmacopoeial standards and monographs given in various herbal pharmacopoeias, pharmacovigilance study and its importance.

COURSE OUTCOME:

The subject aims to provide students with the necessary skills for,

 Applying standardized quality control parameters to test quality of crude drugs from natural origin.

- Determining the adulterations found in herbal crude drugs,
- To follow pharmacopoeial standards and monographs using various herbal pharmacopoeias.
- Understanding of WHO guidelines required for cultivation, collection and quality control of herbal drugs.

CONTENT:

Module 1	Introduction to Pharmacognosy and WHO guidelines	1 cred	lit
Contents	Topics covered		Hrs
	Adulteration and methods to encounter adulteration of plan	nt drugs.	
	Evaluation of plant drugs. Organoleptic evaluation of drugs i	ncluding	(2)
	gross morphology, sampling, preliminary examination and foreign	matter.	
	• Physical evaluation of plant drugs: Determination of moisture	content,	
	foreign organic matter, ash values, extractive values and swelling	ng index.	(2)
	Refractive index, optical rotation and their applications in standa	rdization	
	of plant drugs.		
	• Microscopic evaluation of plant drugs: T.S./L.S./Surface v	views of	
	selected Plant drugs - Use of microtome and preparation of his	stological	(4)
	slides, Quantitative microscopy, vein islet number, vein ter	mination	(4)
	number, stomatal number, stomatal index, palisade ratio. Mic	crometry,	
	measurement of fibers, trichomes, starch grains and calcium	oxalate	
	crystals. Lycopodium spore analysis. Fluorescence analysis		
	• Pharmacopoeial evaluation of plant drugs - Determination of	f various	
	diagnostic features of identification of different plant drugs as per	different	(3)
	herbal pharmacopoeias.		
	WHO Guidelines for cultivation, collection and quality control of	of Herbal	
	Drugs		(2)

Module 2	Cultivation and Extraction of Plant drugs	1credit

Contents	Topics covered	Hrs
	Selection, identification and authentication of herbal materials, drying and	(2)
	processing of herbal raw material	
	Cultivation technology, post harvest care and processing of medicinal and	
	aromatic plants: Profile of some high trade value plants: Chirata, Giloe,	(5)
	Gudmar, Isapgol, Jatamansi, Kalmegh, Kesar, Mulethi, Sarpagandha and	
	Tulsi, Ashwagandha, Belladona, Ginger, Turmeric, Aloes, Digitalis, Vinca,	
	Ephedra, Senna, etc	
	• Extraction of Herbal Materials, Different methods of extraction-	
	conventional (soxhlet, reflux, decoction, percolation, infusion), and	(5)
	novel/green methods (microwave assisted, ultrasonic assisted, supercritical	
	fluid extraction, pressurized extraction), Choice of solvents for extraction	
	Safety of herbals/ herbal pharmacovigilance	
	Herbs as raw materials: Definition of herb, source of herbs, herbal	(1)
	medicine, herbal medicinal product, herbal drug preparation	(2)

Module 3	Isolation of phytoconstituents	1 cred	lit
Contents	Topics covered		Hrs
	• General methods of isolation and separation of phytoconstituents.		(3)
	•Isolation of Phytoconstituents: chemical properties, character	terization	
	(excluding synthesis) and therapeutic uses of some me	edicinally	(6)
	important class of Plant Phenolics (Tannins & flavonoids),	Alkaloids	
	(Quinine,, Atropine, Solasodine Vincristine, Vinblastine, Str	ychnine),	

 Glycosides (Sennoside, Digoxin, Diosgenin,), Terpenoides, Steroids and Resinous substances (Podophyllotoxin), Fixed oils, Volatile oils, Carbohydrates, taxol. Herbal Remedies - Toxicity & Regulations: Importance of Herbal Therapies, Herbal versus Conventional drugs, Efficacy of herbal therapies, safety in herbal drugs, toxicity in Herbals and their interaction, Herbal drug regulations in India 	(2)
Edible dyes sweeteners, perfumery and cosmetic agents from plants Engagement from plant origin.	(1) (1)
 Enzymes from plant origin. Marine plants: Introduction, chemistry and biology of marine products of plant origin 	(1)

Module 4	Nutraceuticals & Project and Seminar Presentation on some recent	1 credit
	research /seminars based on the above topics	
	Nutraceuticals and herbal health supplements	(4)

Assigned 1. W.C. Evans, 2002. "Trease& Evan's Pharmacognosy". WB.Saunders& Reading/ co., London. References 2. T. Swain, 1963. "Chemical plant Taxonomy". Academic Press, London. 3. C.A Stace, 1985. "Plant Taxonomy and Biosystematics". Edward Arnold, London. 4. C.K. Atal, "Cultivation and Utilization of Medicinal plants". R.R.L. 5. H.E. Street, 1997. "Plant Cell and Tissue Culture". Blackwell Scientific, London. 6. N. Takashashi, 1986. "Chemistry of Plant Hormones" CRC Press Inc., Florida. 7. A.R. Gennaro, 2000."Remington: The Science & Practice of Pharmacy". Lippincott Williams & Wilkins, Philadelphia. 8. Kaufmann, "Natural products for plants". CRC press New York. 9. K. Nakanishi, 1977. "Chemistry of Natural Products". Kodansha Book Publishing Company, Osaka (Japan). 10. V. Rajpal, 2002. "Standardization of Botanicals", Eastern Publishers, New Delhi. 11. J.B. Harborne, 1998. "Phytochemical methods", Chapman and Hall. 12. K. Paech, 1956. "Modern methods of plant analysis"., Springer-Verlag 13. Guidelines for the Assessment of herbal medicines, 1991, WHO Report, Geneva. 14. Quality Control Methods for Medicinal Plant material, 1992, WHO Guidelines. 15. Indian Pharmacopoeia, 1996, Govt. of India, Ministry of Health and family welfare, Delhi. 16. A.N. Kalia, Textbook of Industrial Pharmacognosy, 2005, CBS Publishers, New Delhi. 17. Dr.C.K. Kokate, Practical Pharmacognosy, 1988, Vallabh Prakashan, Delhi. 18. Dr.P.Mukherjee, Quality control herbal drugs, 2005, Business Horizons,

M.Pharm. S1-[P&P]-2: ADVANCED PHARMACOGNOSY AND PHYTOCHEMISTRY-I [Practicals]

OBJECTIVES:

- To learn various quality control parameters to test quality of crude drugs from natural origin.
- Identify and quantify adulterations in herbal crude drugs,
- To learn to handle various conventional and Novel extraction techniques
- To make students understand pharmacopoeial standards and monographs given in various herbal pharmacopoeias.

COURSE OUTCOME:

The subject aims to provide students with the necessary skills for,

- Identification of medicinal plants using various macroscopic and microscopic parameters.
- Knowledge of different conventional and novel methods of extraction of crude drugs.
- Application of pharmacovigilance study for herbal drugs.
- Knowledge and skills of selecting suitable techniques for separation and isolation of bioactive phytoconstituents using various chromatographic techniques and spectral analysis.

CONTENT:

		1 credit	
Contents	Experiments		Hrs

- 1. Microscopical evaluation of plant drugs (at least five) listed in theory- Vein islet number, vein termination number, stomatal index, stomatal number, palisade ratio, trichomes, starch grains, calcium oxalate crystals)
- 2. Determination of moisture content, foreign organic matter, ash values, extractive values, swelling index of plant drug.
- 3. Extraction by different methods Conventional (Soxhlet, Maceration, Reflux, Percolation, Stirring etc.) & Novel (Microwave Assisted, Ultrasonic method, Supercritical fluid extraction etc)
- 4. Physico-chemical, phytochemical evaluation of some plant drugs containing alkaloids, terpenoids, glycosides phenolics, steroids.
- 5. Estimation of total solid content and alcohol content in Asava/Arista
- 6. Pharmacopoeial evaluation of natural products
- 7. Determination of vitamin C in some crude drugs

Assigned Reading/ References

- 1. W.C. Evans, Trease and Evans Pharmacognosy, 15th edition2002, W.B. Sounders & Co., London.
- 2. S.S. Handa and M.L. Kaul, Supplement to cultivation and utilization of medicinal plants, 1996, R.R.L Jammu, India.
- 3. Ram P Rastogi, Compendium of Indian Medicinal Plants, 1998, Vol. I-V, CSIR, Lucknow, New Delhi.
- 4. T. Fleming, PDR for Herbal Medicine, 2nd edition, 2000, Medical Economics compant, Mountvale, New Jersy.
- 5. M.J. Cupp, Toxicology and Clinical Pharmacology of Herbal Products, 2000, Humana Press, New Jersy.
- 6. Wealth of India- Raw Materials, 1985, CSIR Publication, New Delhi.
- 7. Dr.P.Mukherjee, Quality control herbal drugs, 2005, Business Horizons, New Delhi.
- 8. V.D.Rangari, Pharmacognosy & Phytochemistry,1st edn.2004,Career Publications, Nasik.

M.Pharm. S1-[P&P]-3: TQM, PATENT REGULATION AND VALIDATION (THEORY) OBJECTIVES:

- To learn preparation, applications and importance of documentation in herbal industry.
- To teach students importance of quality in pharmaceutical products.
- To impart knowledge of total quality management and concepts of GMP, GLP and GCP.
- To give an understanding on quality audits, ICH guidelines, statistical analysis and validation processes, regulatory aspects and Intellectual property rights for herbal products.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Understanding the importance of quality in pharmaceutical products.
- Knowledge of total quality management and concepts of GMP,GLP and GCP.
- Preparation, applications and importance of documentation in herbal industry.
- Knowledge of quality audits, ICH guidelines and statistical analysis.
- Application based knowledge of validation processes, Regulatory aspects and Intellectual property rights for herbal products.

CONTENT:

Module 1	I. To understand basic principles of total quality 1 cred	lit
	management and quality audits	
Contents	Topics Covered	Hrs
	Concept of Total Quality Management, Four M's responsible for Quality	(3)
	variation in pharmaceutical products.	
	• Concepts of GMP, GLP and GCP, Quality control laboratory responsibilities,	(3)
	routine controls on instruments and reagents. Standard test procedures, non-	
	clinical testing, controls on animal house, Data generation and storage.	
	• Documentation and its importance, Manufacturing documents, Standard	
	Operating Procedures, Finished product release documentation.	(2)
	• Quality Audits: Auditing of manufacturing processes and facilities, Quality	
	Review, Compliance reports and handling of Non –compliance.	
	• ICH guidelines: Q1-Q10, Guidelines with special reference to quality by	(2)
	design and risk management.	(3)

Module 2	II. To understand validation of processes and	1 credit
	equipment	1
Contents	Topics Covered	(2)
	• Qualification, validation and calibration of equipment .	
	• Validation of processes like mixing, granulation, drying, compression,	
	filtration, filling, etc.	(3)
	• Validation of sterilization methods and equipment, dry heat steril	ization,
	autoclaving, membrane filtration.	(3)
	Validation of manufacturing processes, Equipment, Environment a	nd Water
	supply systems and analytical methods.	(2)
	• Validation and audits of analytical procedures such as HPLC, UV,	GC,
	HPTLC.	(3)
	Validation of personnel handling the analytical instruments	
		(2)
Module 3	III. Regulatory aspects of herbal pharmaceuticals and IPR	1 credit
Contents	Topics Covered	hrs
	Regulatory aspects of herbal pharmaceuticals in India, US, European Company (1997).	ope and (6)
	other countries, US-FDA and WHO Approval, Clinical trial a	pproval,
	dossier preparation for herbals.	
	• Intellectual Property Rights, Patent search and awareness,	Patent (4)
	applications and filling procedures in India and in other countries	
	• International treaties and conventions on IPR - Paris convention	on, PCT (3)
	- an introduction, PCT application & general rules, WTO / GATT	
	& Uruguay TRIPS, WIPO.	
	Patent infringement, exploitation of patent, abuse of patent	(2)
Module 4	To study safety procedures and health hazards &	1
	Project and Seminar Presentation on some recent research /sem	ninars credit
	based on the above topics	
	To learn safety procedures of pharmaceutical and herbal industries.	. (2)

Assigned Reading/ References

- 1. S.H. Willing, M.M Tucherman and W.S. Hitchings IV, Good Manufacturing
 Practices for Pharmaceuticals, Marcel Dekker,
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- 3. Andrew A . Signore and Terry Jacobs Good Design Practices for GMP Pharmaceutical Facilities Informa Healthcare 2005
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- 8. S. H. Willig, M.M.Tuckerman and W.S.Hitchings, "Good Manufacturing Practices for Pharmaceuticals", Drugs and Pharm. Sci. Series, Vol. 16, Marcel Dekker Inc., N.Y.
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- 11. Report of CIOMS (Council for International Organisations of Medical Sciences) Working Group III, Guidelines for Preparing Core Clinical-Safety Information on Drugs, Geneva. 1995
- 12. Process validation in manufacturing of Biopharmaceuticals-Guidelines current practices, and industrial case studies by Anurag Singh Rathod, Gail Sofer, G.K. Sofer. Informa Healthcare Publisher 2005.
- 13. Remington's the science and practice of pharmacy 21st Edition by Alfonso R. Gennaro. Lippincott Williams & Wilkins Publisher 2006.
- 14. Pharmaceutical Process Validation by Robert A. Nash, Alfred H. Wachter Marcel Dekker Publisher 3rd edition 2003.
- 15. CDSO publications and updates of drug and Cosmetics act and rules (Govt. of India).
- 16. CDER Publications and Guidance
- 17. EMEA Publications and Guidance
- 18. Orange Book, ICH guidelines, Indian Patents Act
- 19. Country specific Regulatory Guidelines (available from internet)
- 20. Govt. Publications on issues affecting sales, distribution, manufacturing, excise, etc.
- 21. J. D. Nally, "Good manufacturing Practice for Pharmaceuticals" Informa Healthcare.
- 22. I. Kanfer& L. Shargel, "Generic Product Development BE issued" Informa Healthcare.
- 23. R. A. Guarino, "New Drug Approval Process. The Global challenges". Informa Healthcare.
- 24. Watcher and Nash, "Pharmaceutical Process Validation". Marcel Dekker.

M.Pharm. S1-[P&P]-4: HERBAL PRODUCT DEVELOPMENT-I (THEORY)

OBJECTIVES:

- To give students an indepth understanding of various aspects of product development such as pre-formulation study design, different methods to identify drugexcipient interactions and herbal drug stability in Herbal product development.
- To impart knowledge to formulate solid dosage forms such as tablets and coating technology.
- To teach preparation and standardization of herbal formulation.
- To impart knowledge on pharmaceutical polymers for novel drug delivery system,
 Drug Dissolution and Diffusion studies, Pharmacokinetic modeling.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Pre-formulation study design, different methods to identify drug-excipient interactions and herbal drug stability in Herbal product development.
- Application based knowledge to formulate solid dosage forms such as tablets and coating technology.
- Preparation and standardization of herbal formulation.
- Study of Pharmaceutical polymers for novel drug delivery system [NDDS].
- Drug Dissolution and Diffusion studies, Pharmacokinetic modeling of the herbal products.

CONTENT:

Module 1	I. Preformulation, Chemical Kinetics and Herbal	Instr. hrs	S
	Drug Stability		
Contents	Topics Covered		Hrs
	• Preformulation Studies with respect to herbal pharmaceuticals:	,	
	pka and solubility kinetics, pH profile, partition coefficient morphology, polymorphism, powder flow, surface characteristics, solublization techniques, drug –excipients compatibility	cteristics,	(6)

protocol for performulation studies	
• Chemical Kinetics & Herbal Drug stability:	
Pathways of drug degradation, Rate & order of reactions, Factors affecting	(4)
reaction kinetics, stability testing, Accelerated studies and shelf life	(4)
assignment as per ICH guidelines.	
Dosage form consideration in preformulation:	
Solid dosage form, solution formulations, emulsion, suspension, freeze	(4)
dried products, topical, pulmonary, evaluations and its regulatory	(4)
considerations, antioxidants, chelating agents, impurity, GMP related to	
drugs from herbal origin.	

Module 2	II. Principles and Techniques of Solid Dosage Forms	1 credit
	and Coating Technology & Study of Pharmaceutical	
	polymers	
Contents	Topics Covered	Hrs
	Solid dosage forms:	(4)
	Recent advances in tablet and capsule technology like double	
	compression, direct compression, capsule filling machine,	
	Excipients (binding agents, super disintegrants, lubricants and	
	diluents) from herbal origin	
	Coating of solid dosage forms:	
	Various types of functional coatings, polymers, Advances in	(2)
	process controls, coating equipments, coating pans, Accela cota,	
	Hi-coater, Driacoater, fluid bed coating equipments, Coating	
	application and metering equipment, particle coating methods,	
	pelletization. Technology.	
	Topical drug delivery systems:	
	Various types of topical drug delivery systems such as creams,	
	gels, nanogels, nanoemulsions, ointments and their evaluation.	(2)
	Pharmaceutical polymers:	

Module 3	III) Drug Dissolution and Diffusion Studies & Study of	1 cred	it
	Pharmacokinetic modeling		
Contents	Topics Covered		Hrs
	• Dissolution Studies:		(3)
	Steady state diffusion-procedure and applications, drug dissoluti	on, drug	
	release, diffusion principles in biological systems, thermodyna	amics of	
	diffusion, Fick's law. Devices for dissolution rate testing viz	., forced	
	convection, non-sink devices, and continuous flow through method	ds; effect	
	of environmental factors in dissolution testing; test apparatus for	r various	
	drug delivery systems.		
	• Pharmacokinetics:		(6)
	Compartmental & Non compartmental analysis, Pharmacokinetic i	modeling	
	approaches, Biopharmaceutical classification of drugs, ab	sorption,	
	permeability and solubility limited drugs, Biowavers for bioequ	iivalence	
	studies, Concepts of in vitro & in-vivo Correlation, One a	nd Two	
	Compartmental Modeling, Statistical Moment Analysis, Non-linear	kinetics.	
	Characterization of biopharmaceutical drugs and phytomedicnes		(3)
Module 4	Project and Seminar Presentation on some recent	1 cred	it
	research /seminars based on the above topics		

Assigned Reading/

References

- Carstensen, Thuro J., "Pharmaceutical principles of solid dosage forms",
 Volume 110, Marcel Dekker New York, 2001, CRC
- 2. Ray and Weller, "Handbook of Pharmaceutical Excipients", Pharmaceutical Press, 2009.
- 3. Lachman, Lieberman, "Pharmaceutical dosage forms: Dispersed systems", Vol. I, II, Marcel-Dekker New York, 2008.
- 4. Lisbeth, Illum & Stanley S. Davis, "Polymers in Controlled Drug Delivery", Wright, Bristol, 1987.
- 5. ICH Guidelines available at: http://www.ich.org
- 6. Rawlins, E. A., "Bentley's text book of Pharmaceutics" 8th edition, London: Bailliere Tindal.1995.
- 7. Rubinstein, M. H. M. E. Aulton, "Pharmaceutics: the science of dosage form design", 3rd edition, pp. 304-321, London: ELBS Longman Group Ltd., 1988
- 8. Rudnic, E. M., & Schwartz, J. D. "Remington: The Science and Practice of Pharmacy" Philadelphia: Lippincott Williams & Wilkins. 2006.
- Saha, S., & Shahiwala, A. F., "Multifunctional coprocessed excipients for improved tabletting performance". Expert Opinion on Drug Delivery ,pp 197-208, 2009.
- 10. David K Platt, Biodegradable Polymers, iSmithers Rapra Publishing, 2006.
- 11. Catia Bastioli, Handbook of biodegradable polymers, iSmithers Rapra
- 12. Shaikh R., Sial A., "Stability of pharmaceutical formulations", Pak. J. Pharm. Sci., 2nd edition, pp 83-86 1996,.
- 13. ICH Q1A (R2), "Stability testing of new drug substances and products", International Conference on Harmonisation, IFPMA, Geneva, 1996 Milo Gibaldi and Donald Perrier, "Pharmacokinetics", Drugs and Pharm. Sci. Series, Vol. 15, Marcel Dekker Inc., N.Y.

- 14. Shargel L, Susanna Wu-Pong, Andrew B. C. Yu. "Applied Biopharmaceutics and pharmacokinetics", 3rd edition, McGraw-Hill, Medical Pub. Division, 2005
- 15. Malcolm Rowland, Thomas N. Tozer., "Clinical Pharmacokinetics: Concept and Application"; 3rd Edn. B. I. Lea & Febiger, 1989.
- 16. Shargel, "Generic Drug Product Development Specialty Dosage Form", 1st Edition, 2010.

Semester II

SEMESTER II

M.Pharm. S2-[P&P]-1: MODERN ANALYTICAL TECHNIQUES II [Theory] OBJECTIVES:

- To impart students an in-depth understanding of various types of planar chromatographic techniques such as Paper Chromatography, Thin Layer Chromatography and High Performance Thin Layer Chromatography HPTLC.
- To impart students an understanding of various column chromatographic techniques such as HPLC, GC and its industrial applications.
- To give an understanding of structure elucidation of pure isolated phytoconstituents— Theory and Problem solving, using spectral analysis such as UV, IR, Mass spectroscopy, NMR etc. which can be used for characterization of bioactive phytoconstituents from herbal sources.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Learning principles and techniques of various types of planar chromatography such as PC, TLC and HPTLC,
- Understanding of principles and techniques of various types of column chromatography such as HPLC, GC etc.
- Structure elucidation of pure isolated phytoconstituents Theory and Problem solving, using spectral analysis such as UV, IR, Mass spectroscopy, NMR etc. which can be used for characterization of bioactive phytoconstituents from herbal sources.

CONTENT:

Module 1	Principles and Techniques of planar chromatography	1 credit	
Contents	Topics covered		Hrs

	General principles, theory and the applications of planar	
	chromatographic techniques	
	• Techniques and instrumentation of thin layer chromatography (TLC)	(4)
	• Techniques and instrumentation of Paper chromatography (PC)	(2)
	• Techniques and instrumentation of High performance thin layer	
	chromatography (HPTLC) Analytical method development and its	(5)
	validation as per ICH guidelines. Quantification using HPTLC.	
	 Applications of TLC, PC, HPTLC for analysis of herbals. 	(2)
	• Comparison of planar chromatography and column chromatography	(1)
Module 2	Principles and techniques of column chromatography 1 cr	edit
Contents	Topics covered	Hrs
	General principles, theory and the application of column	
	chromatographic techniques:	
	Principle and techniques of conventional column chromatography	(1)
	Techniques, instrumentation and Applications of High performance	
	liquid chromatography (HPLC) – Theory of HPLC-Van Deemter	(6)
	Equation, various detectors used, derivatisation in HPLC. Analytical	
	method development, validation as per ICH guidelines and	
	troubleshooting. Quantification methods used in HPLC. Ultra pressure	
	liquid chromatography.	
	Techniques, instrumentation and Applications of Gas chromatography	
	(GC)-Theory of GC, packed column, Capillary column, carrier gases	(5)
	used.	
	Techniques, instrumentation and Applications of Supercritical Fluid	(1)
	Extraction	
	Techniques, instrumentation and Applications of Flash Chromatography	(1)
	Techniques, instrumentation and Applications of Size exclusion	
	chromatography and ion pair chromatography	(2)

Module 3	Structure elucidation of organic compounds- Theory and 1 cred	it
	Problem solving	
Contents	Topics covered	Hrs
	General principles, theory and the application of techniques for	
	structure elucidation of organic compounds	
	• Theory, principle, instrumentation of Mass spectrometry: use of isotopic	(7)
	abundance in molecular formula calculation. Different ionization	
	techniques like EI, CI, FD, FI, MALDI, API, ESI. Fragmentation of	
	molecule using these techniques. Tandem mass spectrometry and its	
	applications for pharmaceuticals and herbals, different types of Mass	
	spectrometry	
	• High Resolution 1H &13C Nuclear magnetic resonance (NMR)	(7)
	Spectrometry Theoretical calculation of chemical shifts of various carbon	
	atomsTheory, principle of NMR spectroscopy, instrumentation, Different	
	1D & 2D NMR correlation spectrometric techniques such as COSY,	
	NOESY, HETCOR, INADEQUATE, HSBC, HMQC etc. Use of this	
	technique in determination of absolute configuration.	

Module 4	Problem solving in structure elucidation & Project and Seminar	1 credit
	Presentation on recent research /seminars based on above topics	
	Problem solving in structure elucidation of phytosterols,	(7)
	flavonoids and terpenoids. organic compounds using UV, IR,	
	NMR, 1HNMR, 13CNMR and Mass spectroscopy	

Assigned Reading/ References

- 1. E. Stahl, Thin-Layer Chromatography, A Laboratory Handbook. 2nd Edition Springer-Verlag Berlin–Heidelberg–New York 1969.
- 2. Wagner & S. Bladt, Plant Drug Analysis by H., 2nd Edition, Springer 2001.
- 3. Jiri Gasparic and Jaroslav Churacek, A Laboratory Handbook of paper and thin layer chromatography, Ellis Horwood limited 1979.
- 4. P.D. Sethi, HPTLC Quantitative Analysis of Pharmaceutical Formulations. CBS Publishers and. Distributors, New Delhi, 1996.
- F. James Holler, Stanley R. Crouch Douglas A. Skoog. Principles of Instrumental Analysis, , Publisher: Brooks/Cole Pub Co; 6th edition, 2006
- 6. Analytical Chemistry by open learning series, Wiley Publisher, 2nd Edition
- 7. David G. Watson Pharmaceutical analysis: a textbook for pharmacy students and pharmaceutical chemists Elsevier/Churchill Livingstone, 2005
- 8. Bernard Fried, Joseph Sherma, Thin-layer chromatography 4th Edition Marcel Dekker 2005
- 9. Instrumental methods of Analysis by Higuchi, CBS Publishers. 1997
- 10. High Performance Liquid Chromatography: Analytical Chemistry by open learning series, Wiley Publisher, 2nd Edition 1992.
- 11. W. John Lough, High performance liquid chromatography: fundamental principles and practice Blackie Academic & Professional Publisher, 1995
- 12. HPLC: High Performance Liquid Chromatography: Volume 2, by P.D. Sethi and Rajat Sethi CBS Publisher, 2008
- 13. F. James Holler, Stanley R. Crouch Douglas A. Skoog. Principles of Instrumental Analysis, , Publisher: Brooks/Cole Pub Co; 6th edition, 2006
- Frank A. Settle , Brian D. Lamp , David L. McCurdy, Mark F,
 Vitha , Brian W. Gregory, Yinfa Ma Instrumental Methods of Analysis Wiley-Interscience; 8th edition, 2011
- 15. R.M. Silverstein, G.C., Bassler, T.C. Morrill Spectroscopic identification of organic compounds John Wiley and Sons, New York, 5th Edition. 1991.
- William. Kemp Organic Spectroscopy 3 edition. W.H. Freeman & Company; 1991

- 17. J.R. Dyer, Applications of absorption Spectroscopy of Organic compounds Prentice Hall, London 2009
- 18. Pavia D. L., 2009. "Introduction to spectroscopy". 4th, Belmont CA
- 19. Munson & Munson, "Pharmaceutical analysis: modern methods". edited by James W. Munson, New York: M. Dekker
- Kenneth A. Connors, 2007. "A Textbook Of Pharmaceutical Analysis"
 3rd Ed. Wiley India-wse
- 21. Jens Thuro Carstensen, 2001. "Advanced pharmaceutical solids" Marcel Dekker, New York
- 22. Joseph B. Lambert, Scott Gronert, Herbert F. Shurvell, David Lightner, Robert Graham Cooks, 2010.
- 23. "Organic structural spectroscopy", 2nd Ed. Pearson Education, Limited.

M.Pharm. S2-[P&P]-1: MODERN ANALYTICAL TECHNIQUES II [Practicals] OBJECTIVES:

- To impart students a hands-on expertise of various types of planar chromatographic techniques such as Paper Chromatography, Thin Layer Chromatography and High Performance Thin Layer Chromatography HPTLC.
- To train students to perform various column chromatographic techniques such as HPLC, GC and its industrial applications.
- To give an understanding of application of structure elucidation of pure isolated phytoconstituents- Theory and Problem solving, using spectral analysis such as UV, IR, Mass spectroscopy, NMR etc. which can be used for characterization of bioactive phytoconstituents from herbal sources.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

 Applications of PC, TLC, HPTLC, HPLC and GC which can be applied for identification and analysis of herbal crude drugs and products and for separation, isolation and analysis of marker compounds, extracts and herbal formulations. • Structure elucidation of pure isolated phytoconstituents - Theory and Problem solving, using spectral analysis such as UV, IR, Mass spectroscopy, NMR etc. which can be used for characterization of bioactive phytoconstituents from herbal sources.

Module 1	Techniques of planar chromatography- TLC, PC,	1 credit
	HPTLC.	
Contents	Experiments	Hrs
	1. Development of suitable solvent system for the separation of mixture	s of (6)
	organic compounds by TLC.	
	2. Development of suitable solvent system for the separation of herbal	(6)
	extracts by TLC.	
	3. Quantitative separation of phytoconstituents from herbal extracts by	(4)
	Preparative thin layer chromatography.	
	4. Use of various derivatising agents for detection of phytoconstituents	by
	TLC and PC.	(4)
	5. Separation of sugars/ amino acids by TLC and PC.	(4)
	6. Demonstration and hands on training on High performance thin layer	(2)
	chromatography.	
	7. Analytical method development for three component mixture using	(4)
	HPTLC	
Module 2	Techniques of column chromatography- HPLC, GC, Flash	1credit
	chromatography, Super critical fluid chromatography.	
Contents	Experiments	Hrs
	8. Demonstration of High performance liquid chromatography (HPLC).	(2)
	9. Plotting of a standard curve for caffeine / betaine/ catechin by HPLC.	(6)
	10. Quantitative estimation of caffeine in cola drinks and tea extract by	(6)
	HPLC.	(4)
	HPLC.	(4)

11.	To check the effect of alteration of various parameters on retention times	(6)
	(RT) of compounds by HPLC.	(6)
12.	Determination of HETP value, selectivity factor, resolution, tailing factor	(4)
	by HPLC.	(4)
13.	Demonstration of Gas liquid chromatography/ Flash chromatography/	(2)
	Supercritical fluid extraction.	(2)

Module 3	Structure elucidation of organic compounds- Problem solving	1 credit
Contents	Experiments	Hrs
	14. Identification of various functional groups (amine, nitro, aldehyde, ke	eto, (6)
	carboxylic, hydroxyl, etc.) by UV and IR	
	15. Identification of different functional groups by PNMR.	(6)
	16. Identification of different types of carbons and carbon containing gro	oups (4)
	by 13 CNMR	
	17. Identification of Molecular ion peak, base peak in a mass spectrum of	f (6)
	small molecular weight phytoconstituents.	
	18. Structure elucidation of some small molecular weight phytoconstitue	nts (6)
	by UV, IR, NMR and MS spectral data.	

Assigned	6. Thin-Layer Chromatography, A Laboratory Handbook by E. Stahl, Second.
Reading/	Edition, Springer-Verlag Berlin–Heidelberg–New York 1969.
References	7. Plant Drug Analysis by H. Wagner & S. Bladt, Second Edition, Springer.
	8. A Laboratory Handbook of paper and thin layer chromatography by Jiri
	Gasparic and Jaroslav Churacek, Publisher-Ellis Horwood limited.
	9. Manual of HPTLC applicator, scanner and photodocumentation system by
	CAMAG
	10. HPLC: High Performance Liquid Chromatography: Volume 2, by P.D.
	Sethi and Rajat Sethi CBS Publisher, 2008

M.Pharm. S2-[P&P] -2: ADVANCED PHARMACOGNOSY AND PHYTOCHEMISTRY-I [Theory]

OBJECTIVES:

- To impart students' knowledge on standardization of herbal drugs, qualitative and quantitative phytochemical evaluation of herbal extracts using various analytical techniques from industry point of view.
- To teach drug discovery and development of novel phytoconstituents from natural sources.
- To handle various aspects of regulatory requirements/documentation required in herbal industry..

COURSE OUTCOMES

The subject aims to provide students with the necessary skills for,

- Industrial application of standardization of herbal drugs, qualitative and quantitative phytochemical evaluation of herbal extracts using various analytical techniques.
- Understanding of drug discovery and development of novel phytoconstituents from natural sources such as Taxol, Artemisin etc.
- Handling regulatory requirements/documentation required for herbal products...

Module 1	Standardization of Herbal drugs 1	credit
Contents	Topics covered	Hrs
	Chemotaxonomy: Significance in classification of medicinal plants, distribution of chemotaxonomic groups of constituents in plant kingdom like alkaloids, glycosides, terpenoids etc.	(2)
	• Phytochemical evaluation of plant drugs: Qualitative and Quantitative evaluation of phytoconstituents such as Alkaloids, steroids, terpenoids, flavonoids, glycosides, tannins etc. Chemoprofiling, TLC & HPTLC fingerprinting of crude drugs.	(2)

Bioactivity guided fractionations, Cytotoxicity tests	(2)
• Drug discovery and development from natural products with special emphasis on drugs derived from some of the following Plants: Ashwagandha, Digitalis, Artemesia, <i>Atropa belladonna, Catharanthus roseus</i> , Podophyllum, Taxus species.	(6)

Module 2	Plant Biotechnology 1 cr	1 credit	
Contents	Topics covered	Hrs	
	Historical perspectives, prospects for development of plant biotechnology	(1)	
	as source of medicinal agents. Applications in pharmacy and allied fields.		
	 Types, techniques, nutritional requirements and growth of plant tissue cultures. Organogenesis and embryogenisis. Protoplast fusion and cultures, artificial seeds, micropropogation of medicinal and aromatic 	(2)	
	plants. Genetic stability of tissue cultures.		
	• Secondary metabolism in tissue cultures with emphasis on production of		
	medicinal agents. Screening and selection of high yielding cell lines Effect of cultural practices, precursors and elicitors on production of biomedicinals.		
	 Plant finger print analysis: Methods used in gene identification localization and sequencing of genes. Application of PCR to plant genome analysis 		
	 Biotransformation, bioreactors, industrially potential tissue cultures systems for pilot and large scale cultures of plant cells, cellular totipotency, cryopreservation and retention of biosynthetic potential in cell 	(2)	

	cultures.	
•]	Immobilized plant cell culture systems, immobilization techniques, effect	
	of immobilization on secondary metabolism and realization of	(2)
	chemosynthetic potential in immobilized cells. Genetic transformation	
1	methods, Hairy root cultures and their applications.	
•	Basic metabolic pathways and techniques employed in elucidation of	(2)
1	biosynthetic pathway.	
•]	Biogenesis of tropane, quionoline, Imidazole, Isoquinoline and Indole	(2)
	alkaloids; Sterols, Anthraquinone and Saponin glycosides; Flavanoids;	
	and Isoprenoid compounds of pharmaceutical significance	

Module 3	Regulatory aspects of herbals	1credit	
Contents	Topics covered		Hrs
	• Recent Methods (UV, HPLC, HPTLC, etc.) of assay of Androgra	pholide,	(4)
	Amarogenin, Asiaticisides, Atropine, Solasodine, Bacoposide, C	Caffeine,	
	Cubebol, Citral, Curcumin, Digitoxin, Diosgenin, Embelin, I	Emetine,	
	Ergometrine, Eugenol, Gingerol, Gycerrhetinic acid, Hesp	peridine,	
	Kutkosides, Piperine, Plumbagin, Quinine, Quinidine, Recinol	ic acid,	
	Sennosides, Taxol, Vinca alkoloids, Withaferin, etc. in ex	xtract /	
	formulations.		
	• Qualitative and quantitative estimations exemplified by the me	ethod of	(2)
	preparation of at least two standardized extracts. Single an	d multi	
	components herbal formulations		
	Stability studies for extracts (Predictable chemical and galenical cha	anges)	(2)
	Pharmaceutical aids: Profile for manufacture and commerce of	Papain,	
	Pectin, Pharmaceutical gums, Starch, Absorbent cotton and	Gelatin.	(2)
	Methods of preparation of different conventional solid and liquid	l dosage	

forms incorporating herbal extracts.	
• GMP and other regulatory and safety requirements (ICH,OECD),	
Schedule Y, Drug and Cosmetic Act and Rules for Herbal, Ayurvedic and	(4)
other Drugs of traditional origin	

research /seminars based on the above topics	Module 4	Project and Seminar Presentation on some recent	1 credit
		research /seminars based on the above topics	

Assigned	1. W.C. Evans, 2002. "Trease& Evan's Pharmacognosy". WB.Saunders &	
Reading/	co., London.	

References

- 2. T. Swain, 1963. "Chemical plant Taxonomy". Academic Press, London.
- 3. C.A Stace, 1985. "Plant Taxonomy and Biosystematics". Edward Arnold, London.
- 4. C.K. Atal, "Cultivation and Utilization of Medicinal plants". R.R.L. Jammu
- 5. H.E. Street, 1997. "Plant Cell and Tissue Culture". Blackwell Scientific, London.
- 6. N. Takashashi, 1986. "Chemistry of Plant Hormones" CRC Press Inc., Florida.
- 7. A.R. Gennaro, 2000."Remington: The Science & Practice of Pharmacy". Lippincott Williams & Wilkins, Philadelphia.
- 8. Kaufmann, "Natural products for plants". CRC press New York.
- 9. K. Nakanishi, 1977. "Chemistry of Natural Products". Kodansha Book Publishing Company, Osaka (Japan).
- V. Rajpal, 2002. "Standardization of Botanicals", Eastern Publishers, New Delhi.
- 11. J.B. Harborne, 1998. "Phytochemical methods", Chapman and Hall.
- 12. K. Paech, 1956. "Modern methods of plant analysis"., Springer-Verlag
- 13. Guidelines for the Assessment of herbal medicines, 1991,WHO Report, Geneva.
- 14. Quality Control Methods for Medicinal Plant material, 1992, WHO Guidelines.
- 15. Indian Pharmacopoeia, 1996, Govt. of India, Ministry of Health and family welfare, Delhi.
- 16. A.N. Kalia, Textbook of Industrial Pharmacognosy, 2005, CBS Publishers, New Delhi.
- 17. Dr.C.K. Kokate, Practical Pharmacognosy, 1988, Vallabh Prakashan, Delhi.
- 18. Dr.P.Mukherjee, Quality control herbal drugs, 2005, Business Horizons,

M.Pharm. S2-[P&P]-2: ADVANCED PHARMACOGNOSY AND PHYTOCHEMISTRY-I [Practical]

OBJECTIVES:

- To develop expertise in various techniques used in isolation of some important phytoconstituents from the crude drugs.
- To teach bioactivity guided fractionations, phytochemical fingerprinting and structure elucidation of phytoconstituents.
- To learn various quality control parameters to test quality of herbal formulations..

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Application based knowledge about recent trends and advances in the field of phytochemistry.
- Expertise in isolation of various important phytoconstituents from the crude drugs.
- In-depth understanding of bioactivity guided fractionations, phytochemical fingerprinting and structure elucidation of phytoconstituents.
- Applying standardized quality control parameters to test quality of herbal formulations.

		1 credit	
Contents	Experiments		Hrs

- 1. Isolation and separation of phytoconstituents by Column chromatography
- 2.Isolation and Chemical Evaluation of Phytochemical Constituents: Curcumin, Caffeine, Quinine, Strychnine, glycyrrhizin and sennosides
- 3.Isolation of volatile oils from various plant drugs and their TLC Characterization
- 4. Isolation of piperine from pepper.
- 5. Isolation & TLC of reserpine from Rauwolfia root.
- 6. Isolation of Hespiridine from orange peel.
- 7. Isolation & TLC of Menthol from mentha oil
- 8. Qualitative and quantitative estimation of phytoconstituents in crude drugs & commercial herbal formulations.
- 9. Preparation of detailed monograph of at least one plant drug covering Pharmacognosy and Phytochemical investigation with its use in traditional system of medicine.
- 10. Preclinical studies of some herbal extracts like analysesic antiinflammatory and antianxiety.

Assigned Reading/

References

- 1. W.C. Evans, Trease and Evans Pharmacognosy, 15th edition2002, W.B. Sounders & Co., London.
- 2. S.S. Handa and M.L. Kaul, Supplement to cultivation and utilization of medicinal plants, 1996, R.R.L Jammu, India.
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- 4. T. Fleming, PDR for Herbal Medicine, 2nd edition, 2000, Medical Economics compant, Mountvale, New Jersy.
- 5. M.J. Cupp, Toxicology and Clinical Pharmacology of Herbal Products, 2000, Humana Press, New Jersy.

- 6. Wealth of India- Raw Materials, 1985, CSIR Publication, New Delhi.
- 7. Dr.P.Mukherjee, Quality control herbal drugs, 2005, Business Horizons, New Delhi.
 - 8. V.D.Rangari, Pharmacognosy & Phytochemistry,1st edn.2004,Career Publications, Nasik.

M.Pharm. S2-[P&P]-3: HERBAL PRODUCT DEVELOPMENT-II (Theory)

OBJECTIVES:

- To give an understanding of designing and evaluation of various site specific drug delivery such as ocular and transdermal drug delivery system and advances in Oral, Mucosal, Intrauterine & Parenteral drug delivery system with respect to herbal drug delivery systems to safely achieve desired therapeutic effect of the herbal drugs with suitable drug delivery system.
- To impart learnings on concepts of rate controlled and site specific drug delivery systems and particulate carrier systems.
- To teach various aspects of packaging materials and product-package compatibility for herbal dosage forms.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Learning concepts of rate controlled and site specific drug delivery systems and particulate carrier systems.
- Understanding the need, concept, design and evaluation of various site specific drug
 delivery such as ocular and transdermal drug delivery system and advances in Oral,
 Mucosal, Intrauterine & Parenteral drug delivery system with respect to herbal drug
 delivery systems to safely achieve desired therapeutic effect of the herbal drugs with
 suitable drug delivery system.
- Knowledge of packaging materials and product-package compatibility for herbal dosage forms.

Module 1	To study concepts of rate controlled and site specific drug 1 credit	
	delivery systems and particulate carrier systems	
Contents	Topics Covered	Hrs.
	• Concepts and systems design for rate controlled delivery: Rate	(3)
	preprogrammed, Activation modulated and Feedback regulated drug	
	delivery systems.	
	Particulate carrier systems: microspheres, liposomes and nanocarriers.	(3)
	• Site specific drug delivery: Active and passive targeting, monoclonal	
	antibodies for drug targeting.	
	1 Ocular delivery of drug: Anatomy & physiology of eye, development	
	of ocular controlled release therapeutic systems, safety & toxicity	(3)
	evaluation.	
	2 Transdermal drug delivery: Permeation through skin, permeation	
	enhancers, technologies for developing transdermal drug delivery	(3)
	systems like gels, patches and sprays and evaluation thereof.	

Advances in Oral, Mucosal, Intrauterine & Parenteral 1	credit
with respect to Herbal Drug Delivery Systems	
Topics Covered	Hrs
• Oral Herbal Drug Delivery Systems: Osmotic pressure controlled,	(3)
membrane permeation controlled, pH controlled, Ion-exchange	
controlled, gel diffusion controlled and hydro dynamically balanced	
systems, modulation of gastro intestinal transit time and release kinetics	
and evaluation thereof.	
• Mucosal herbal drug delivery systems: Mechanism of transmucosal	(3)
permeation and mucosal membrane models, Buccal, Nasal, Pulmonary,	
Rectal and Vaginal drug delivery systems.	
• Intrauterine drug delivery systems: Medicated IUDS, copper IUD,	(3)
	 with respect to Herbal Drug Delivery Systems Topics Covered Oral Herbal Drug Delivery Systems: Osmotic pressure controlled, membrane permeation controlled, pH controlled, Ion-exchange controlled, gel diffusion controlled and hydro dynamically balanced systems, modulation of gastro intestinal transit time and release kinetics and evaluation thereof. Mucosal herbal drug delivery systems: Mechanism of transmucosal permeation and mucosal membrane models, Buccal, Nasal, Pulmonary, Rectal and Vaginal drug delivery systems.

Hormone releasing IUD, long acting contraceptive formulations.	
• Parenteral drug delivery systems: Injectable controlled release	(3)
formulations, long acting depot formulations, implantable drug delivery	

Module 3	III. Packaging materials for herbal dosage forms 1	credit	
Contents	Topics Covered		Hrs
	.Glass containers for Herbal Pharmaceuticals: Glass types,	, their	(2)
	manufacture chemical composition, Performance testing and quality co	ontrol,	
	Defects.		
	Plastics containers for Herbal Pharmaceuticals: Classification	ion of	(2)
	plastics, plastic polymers and their physico-chemical, mechanical	al and	
	biological properties, Quality control testing and biological toxicity.		
	Metal containers: Aluminum and tinplate drums collapsible	tubes.	(2)
	Aerosol containers, Lacquering, coating and lining.		
	Flexible packaging: Types of films, Co-extruded films, foils, coating	ng and	(2)
	laminates, shrink and stretch films, blisters and Strip Packaging.		
	Paper and paperboard: Types of paper, folding cartons, quality c	control	
	testing of paper and paperboard and their common defects		(1)
	Corrugated and solid fibre boards and boxes: Types of corrug	gation,	
	methods, types of box design and Quality control.		(1)
	Caps and Closures: Types of caps, closures, liners, child resistant	t caps.	
	Elastomeric closures for parenterals, classification of Elastomers, ph	nysical	(1)
	chemical and biological properties and their quality control.		
	• Labels and labeling: Types of labels, adhesives, inject and bar codin	ng and	
	printing of labels, Quality control and common defects in printi	ing of	(2)
	labels.		

Module	4	IV. Study of Product package compatibility &	1 credit
		Project & Seminar	

Contents	Topics covered	hrs
	Product–Package compatibility: Stability of product, package selection	(4)
	and development criterion, Line clearance and packaging operation in pharma	
	and herbal industry.	
	• The Seminar topic will be given to the student based on the above content	(8)
	& they have to present the same	

Assigned Reading/ References

- 1. Yie W. Chien, "Novel Drug Delivery Systems", Drugs and Pharm. Sci. Series, Vol. 14, Marcel Dekker Inc., N.Y,1992.
- 2. Robinson J.R and Lee V.L, "Controlled Drug delivery fundamentals and applications", 2nd edition, Marcel Dekker, Inc., *New York*, 1987.
- 3. S.D. *Bruck*, "Controlled Drug Delivery", Vol.1 (Basic Concepts) CRC Press.1983.
- 4. Davis, S.S. and Illum, L., "Colloidal delivery systems- opportunity and challenges, in site specific drug delivery cell biology", John Wiley and Sons, Chichester, , 1986.
- 5. Vyas, S.P. and Khar, R.K., "Targeted and controlled drug delivery novel carriers", CBS, 1st edition, 2002.
- 6. Micheal Roberts, "Dermal Absorption & Toxicity Assessment", 2nd Edition, Vol 177, 2007.
- 7. G.S.Banker & C.T.Rhodes, "Modern Pharmaceutics", Drugs and Pharm. Sci. Series, Vol. 7, Maracel Dekker Inc., N.Y. 1994.
- 8. Jain S., Jain N., "Liposomes as drug carriers", In: Controlled and novel drug delivery", CBS Publishers and Distributors, 1997.
- 9. Carstensen, Thuro J., "Pharmaceutical principles of solid dosage forms" New York Marcel Dekker Volume 110, 2001.
- 10. Lachman, L., Lieberman, H. A., & Kanig, J. L. "The Theory and Practice of Industrial Pharmacy" 3rd edition, Mumbai: Varghese Publishing House,1991
- 11. Rawlins, E. A. (1995). Bentley's text book of Pharmaceutics, 8th edition, London: Bailliere Tindal.
- 12. Micheal Rathbone, "Modified Drug Release Drug Delivery Technology", 2nd Edition, Vol 1, 2008.
- 13. Chilukuri D.," Pharmaceutical Product Development: In Vitro-In Vivo Correlation", Vol 165, 2007.
- 14. Rubinstein, M. H.,. Aulton M. E., "Pharmaceutics: the science of dosage form design", pp. 304-321., London: ELBS Longman Group Ltd. 1988.
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Dekker, Inc.1993

- 16. Rudnic, E. M., & Schwartz, J. D. "Remington: The Science and Practice of Pharmacy", Philadelphia: Lippincott Williams & Wilkins. 2006.
- 17. Saha, S., & Shahiwala, A. F. "Multifunctional coprocessed excipients for improved tabletting performance", 2nd edition, 2009
- 18. K.E.Avis, "Pharmaceutical Dosage Forms: Parental Medication", Vol. I Marcel Dekker Inc., N.Y, 2008.
- 19. S. Turco and R.E. King, "Sterile Dosage Forms", 2nd edition, 1998.

M.Pharm. S2-[P&P] -4:AYURVEDA AND ALLIED PLANT BASED THERAPIES (THEORY)

OBJECTIVES:

- To impart knowledge on primary concepts and principle of various traditional system of medicines such as Ayurveda, Unani, Homeopathy and Siddha.
- To give an understanding of formulation development and standardization of various formulations used in alternative systems of medicines.
- Application based usage of monographs of medicinal plants in various pharmacopoeias
 for studies, salient features of the techniques of preparation of some of the important
 class of formulations as per Ayurveda, Siddha, Homeopathy and Unani Pharmacopoeia
 and other texts.
- A thorough understanding of standardization, shelf life and stability studies of different Indian systems of medicines.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Understanding primary concepts and principle of various traditional system of medicines such as Ayurveda, Unani, Homeopathy and Siddha.
- Gaining knowledge of preparation and standardization of various formulations used in alternative systems of medicines.
- Using monographs of medicinal plants in various pharmacopoeias for studies, salient features of the techniques of preparation of some of the important class of formulations as per Ayurveda, Siddha, Homeopathy and Unani Pharmacopoeia and other texts.

• Understanding of standardization, shelf life and stability studies of different Indian systems of medicines.

Module 1		Principles of Ayurvedic system of medicine 1 credit	:
Contents		Topics Covered	Hrs.
	•	Basic principles of treatment in Ayurvedic System of medicine	(2)
	•	Plants used in Ayurveda for various disorders.	(2)
	•	Preparation of Ayurvedic formulations like Asava, Arista, Bhasma, Churna,	(4)
		Ghrita, Vati /Gutika/pills, Avaleha, Kshar, Kashaya, Taila etc as per	
		Ayurvedic Pharmacopoeia and texts	
	•	Introduction to Ayurvedic Pharmacy, some aspects of standardization of	(4)
		Ayurvedic formulations	
	•	Clinical trials, introduction to "Reverse Pharmacology" and other models	(3)

Module 2		Principles of Allied systems of medicine 1 credit	
Contents		Topics Covered	Hrs.
	•	Basic principles of treatment in Unani system of medicine - Common plants used and types of formulations.	(3)
	•	Basic principles of treatment in Homeopathic systems of medicine - Common plants used and types of formulations.	(4)
	•	Basic principles of treatment in Siddha systems of medicine - Common plants used and types of formulations.	(3)
	•	Ethnoplants, plants used in folklore and tribal medicines	(3)

Module 3	Monographs of plants in various pharmacopoeias 1 credit	
Contents	Topics Covered	Hrs.
	 Monographs of various medicinal plants and products as per Indian Herbal Pharmacopoeia (IHP) 	(2)
	 Monographs of various medicinal plants and products as per Ayurvedic Pharmacopoeia of India 	(2)
	 Monographs of various medicinal plants and products as per Indian Pharmacopoeia 	(3)
	 Monographs of various medicinal plants and products as per British Herbal Pharmacopoeia, Comparison of IHP,IP,BHP and Ayurvedic Pharmacopoeia Monographs of various medicinal plants and products as per Homeopathic 	(2)
	Pharmacopoeia of India and American Homeopathic Pharmacopoeia	(2)
	 Monographs of various medicinal plants and products as per Siddha Pharmacopoeia of India 	(1)
	 Monographs of various medicinal plants and products as per Unani Pharmacopoeia of India 	(1)

Module 4	Project and Seminar Presentation on some recent	1 credit
	research /seminars based on the above topics	

Assigned Reading/ References

- 1. Indian Herbal Pharmacopoeia (IHP), Revised edition 2002, published by Indian Drug Manufacturers Association.
- 2. Ayurvedic Pharmacopoeia of India, Vol. 1-6, 2007, Government of India, Ministry of Health and Family Welfare, CCRAS, Department of AYUSH, New Delhi,
- 3. Indian Pharmacopoeia, 6th edition Vol.1-3, 2010, Government of India, Ministry of Health and Family Welfare
- 4. British Herbal Pharmacopoeia, 1996, published by Scientific Committee of the British Herbal Medicine Association.
- Homeopathic Pharmacopoeia of India, Vol. 1-7, 1986, Government of India, Ministry of Health and Family Welfare
- American Homeopathic Pharmacopoeia, 1882, Boericke and Tafel, Kessinger Publishing's Legacy.
- 7. Siddha Pharmacopoeia of India, Part 1, vol. 1, first edition, 2008, published by Department of AYUSH, Government of India
- 8. Unani Pharmacopoeia of India Part 1, vol. 1-6, Part 2, vol. 1-2, 2008, published by Department of AYUSH, Government of India

Semester III

SEMESTER III

M.Pharm. S3-[P&P] -2: BIOLOGICAL EVALUATION (Theory)

OBJECTIVES

- To train in pre-clinical drug evaluations and recent experimental techniques in the drug discovery and development.
- To impart knowledge of maintenance of laboratory animals as per the guidelines.
- To impart students in-depth knowledge of various in-vitro and in-vivo preclinical evaluation processes and the regulations and ethical requirement for the usage of experimental animals.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Understanding of pre-clinical drug evaluations and recent experimental techniques in the drug discovery and development.
- Knowledge of maintenance of laboratory animals as per the guidelines.
- In-depth knowledge of various in-vitro and in-vivo preclinical evaluation processes and the regulations and ethical requirement for the usage of experimental animals.

Module 1	Pre-clinical drug evaluation	1 credit
Contents	Topics Covered	Inst. hrs
	New Drug Discovery and development process- Leads from plants.	(3)
	• Pre-clinical drug evaluation for biological activity as per Schedule	
	Y, ICH and OECD guidelines: Safety studies in animals- for acute,	(5)
	sub-acute and repeat dose toxicity studies, ED50 and LD50	
	determination, Special toxicity tests like teratogenecity,	
	mutagenicity, carcinogenicity and reproduction toxicity studies,	
	genetic toxicology,	
	• Clinical Trials, Phase I, II & III	
	• Screening of drugs, Transgenic and Knockout animals, Biostatistics	(3)

	and calculation of doses in experimental pharmacology	(3)
Module 2	II. Biological standardization and Molecular biology	1 Credit
Contents	Topics Covered	Hrs
	• General principles, scope and limitations of bioassay, bioassay of official drugs	(3)
	Molecular biology of receptors such sodium, calcium and potassium	
	ion channels as well as GPCR.	(2)
	Biological evaluation of drugs- Screening and evaluation (including	
	principles of screening, development of models for diseases: <i>in vivo</i> models/ <i>in vitro</i> models/ cell line study) techniques of the following:	(8)
	1.Cardiovascular drugs(antihypertensive, antiaarythmic ischemic heart diseases, atherosclerosis, cardiotonic)	
	2.Drugs acting on CNS(anesthetics, sedatives, hypnotics, antiepileptics, antiparkinsonism,	
	3. Drugs used in metaboliic syndrome- diabetes	
	4. Analgesic drugs and anti-inflammatory drugs	
	5. Gastro intestinal(peptic ulcers, IBS, hormone and endocrine	
	disorders)	
	6. Adrenegric and cholinergic drugs	
	7.Drugs used in respiratory disorders	
	8.Antifertility drugs and diuretics	
Module 3	Microbiological evaluation and Pyrogen Science	1 credit
Contents	Topics Covered	
	1. Microbiological limit tests with special emphasis on Crude drugs	(2)
	2. Bacterial, fungal, microbial counts – Principle & Methodology	(2)
	3. Sterility tests: Methodology & Interpretation	(2)
	4. Tests for effectiveness of antimicrobial preservatives.	(2)
	5. Sources, chemistry and properties of bacterial Pyrogens and	(2)

	endotoxins,	(2)
	6. Official Pyrogen tests- IP, BP & USP, Interpretation of data, and	(2)
	comparison of LAL.	
	7. Microbiological assay of antibiotics and vitamins.	(3)
Module 4	Project and Seminar Presentation on some recent research	1 credit
	/seminars based on the above topics	

M.Pharm. S3-[P&P] -2: BIOLOGICAL EVALUATION [Practicals]

OBJECTIVES:

- To impart knowledge of various in-vitro and in-vivo preclinical evaluation processes and the regulations and ethical requirement for the usage of experimental animals.
- To teach maintenance of laboratory animals as per the guidelines.
- To build expertise in microbiological testing of various herbal formulations.

COURSE OUTCOMES:

The subject aims to provide students with the necessary skills for,

- Knowledge of maintenance of laboratory animals as per the guidelines.
- In-depth knowledge of various in-vitro and in-vivo preclinical evaluation processes and the regulations and ethical requirement for the usage of experimental animals.
- Microbiological testing of various herbal formulations.

Module 1	Introduction to Animal Models	Credit 1
Contents	Experiments	Practical
		hrs
	1. To Study Effect of diazepam on locomotor activity by actophotometer.	(4)
	2. To Study Effect of diazepam on muscle grip by rota-rod apparatus	(4)
	3. To Study Effect of diazepam on behavioral activity by hole-board technique and elevated plus maize.	(6)
	4. To Study of chlorpromazine induced catatonia.	(4)
	5. Effect of analgesic by hot plate technique	(6)

Module 2	Microbiological testing of various formulations.	Credit 1
Contents	Experiments	
	1. Effectiveness of antimicrobial agent by cup plate method	(4)
	2. Effectiveness of antimicrobial agent using Ditch plate method	(6)
	3. To perform Sterility test of given sample.	(4)
	4.To determine minimum inhibitory concentration of given	(6)
	antibiotic	(4)
	5. To perform the Microbial limit tests of Crude drugs	
Module 3	Demonstrations	Credit 1
Contents	Experiments	
	1. Demonstration of animal handling techniques.	(4)
	2. Demonstration of various routes of administration	(4)
	3. Demonstration of Bioanlayser	(6)
	4. Demonstration of Blood Cell Counter	(4)
	5. Demonstration of Water Maize Apparatus	(6)

M.Pharm. S3-[P&P]-3: COMPUTING & STATISTICS [Theory]

OBJECTIVES

- To impart the students understanding on application of computers in pharmaceutical sciences, stores management, inventory control, drug information systems and hospital information systems.
- Introduce the students to computer-aided drug design (CADD), QSAR.
- To teach them statistical techniques in solving the pharmaceutical problems.
- To build an expertise in statistical data analysis & applications of spreadsheet to pharmacy.
- To teach concepts of Statistics Probability, internet & application of softwares in data interpretation.

COURSE OUTCOMES

The subject aims to provide students with the necessary skills for,

• Application of computers in pharmaceutical sciences, stores management, inventory

control, drug information systems and hospital information systems.

- Knowledge of the statistical techniques in solving the pharmaceutical problems.
- Introduction to computer-aided drug design (CADD), QSAR various softwares and molecular modeling in CADD.
- Understanding concepts of Statistics Probability, internet & application of softwares in data interpretation.
- Understanding the statistical data analysis & application of spreadsheet to pharmacy.

Module 1	I. Basics of computers	1 cred	it
Contents	Topics Covered		hrs
	• Application of computers in pharmaceutical sciences, stores mana	agement,	(3)
	inventory control, drug information systems and hospital info	ormation	
	systems		
	• Access to and retrieval of information: Smart search using interne	et, use of	(3)
	search engines and web sites, drug information sources.		
	• Computer applications in pharmacy, with special reference to form	mulation	(3)
	development, production, quality assurance, and validation.		
	• Modeling and simulation of data with application in pharmacokinetic	ics	(3)

Module 2	II. Applications in Pharmacy 1	credit
Contents	Topics Covered	
	• Introduction to computer-aided drug design (CADD), QSAR various software's and molecular modeling in CADD	(3)
	• Importance and generation of physico-chemical descriptors using various software's.	(3)
	• Correlation methods and generation of molecular models using computer	(3)

I	software's. Interpretation and statistical significance of molecular models	
	developed using softwares.	(3)
	• Structure based and pharmacophore based drug designing using CADD.	
	Importance of docking studies in drug development.	

Module 3	III. Concept of Statistics	1 cre	dit
Contents	Topics Covered		Hrs
	Probability: Laws of probability, types of distribution.		(4)
	 Hypothesis testing: Types of errors, tests for significance: one-tailed two- tailed tests, t test, z test, chi-square test. 	d and	(4)
	 Correlation and regression: definition and calculation of correlation coefficient, regression coefficient, least square, method, linear regre 		(4)

Module 4	IV. Application of Statistics	1 cred	lit
Content	Topics Covered		Hrs
	Analysis of variance: Completely randomized design randomized completely randomized design randomized completely randomized design randomized completely randomized design randomized design randomized completely randomized design randomized d	omplete	(4)
	block design, Factorial design, and response surface graphs.		
	• Non-parametric tests: The sign test, The Mann-Whitney U test, T	he Runs	(4)
	test, Spearman's rank correlation.		
	Role of statistics in design of pharmaceutical and biomedical exper-	iments	(4)
	specially controlled clinical trials.		

M.Pharm. S3-[P&P]-3: COMPUTING & STATISTICS [Practical]

OBJECTIVES

- To impart the students understanding on statistical techniques in solving the pharmaceutical problems.
- To teach them applicationS of computers in pharmaceutical sciences, stores management, inventory control, drug information systems and hospital information

systems.

- To build an expertise in statistical data analysis & applications of spreadsheet to pharmacy.
- Introduce the students to computer-aided drug design (CADD), QSAR.
- To teach concepts of Statistics Probability, internet & application of softwares in data interpretation.

COURSE OUTCOMES

- The subject aims to provide students with the necessary skills for,
- Hands-on knowledge of the statistical techniques in solving the pharmaceutical problems.
- Introduction to computer-aided drug design (CADD), QSAR various softwares and molecular modeling in CADD.
- Understanding concepts of Statistics Probability, internet & application of softwares in data interpretation.
- Understanding the statistical data analysis & application of spreadsheet to pharmacy.

Module 1	1. Basics of computers -	
Contents	Experiments	(20)
	Major Commands For Windows Operating System	(6)
	• Introduction To Word Processing (MS word)	(4)
	• Presentation Tool: Introduction to presentation tool, features and functions,	
	Creating presentation, Customizing presentation, Showing presentation.	(6)
	Tools used may be Microsoft Power Point, Open Office or similar tool.	

Module 2	II. Use of internet & application of softwares in c	ata -	
	interpretation		
Contents	Experiments	•	(20)

Introduction to Internet, Use of Internet and www	(6)
Applications of Software-QSAR, CADD, Pharmacokinetics, Factorial	(6)
design.	
• Using search engines like Google, Yahoo etc, Using advanced search	(6)
techniques. Literature search using various search engines like google,	
pubmed, science direct, freepatentsonline.	

Module 3	III. Statistical Data Analysis & Application of -				
	Spreadsheet to Pharmacy				
Contents	Experiments	(20)			
	• Spreadsheet Tool: Introduction to spreadsheet application, features and				
	functions, Using formulas and functions, Data storing, Features for				
	Statistical data analysis, Generating charts/ graph and other features. Tools				
	used may be Microsoft Excel, Open office or similar tool.				
	R-Project: Statistical package.	(6)			

M.Pharm. S3-[P&P]-4: SUBJECT 3: RESEARCH METHODOLOGY

OBJECTIVES

- To inculcate an understanding of research methodology and study various aspects and ethics associated with it.
- To impart an understanding of problem identification, its implementation and evaluation and also introduce various research funding agencies for pharmacy.
- To introduce students to different methods of assessment and concepts of basic research and give a brief overview of formation of research problem.
- To teach in detail concepts of mathematical modeling and types involved in processes of formulation of model based on simulation.
- To give an understanding of experimental modeling, general model of process and introduce risk assessment and uncertainty associated with experimental modeling.
- To inculcate an understanding of research deliverables in form of various publications, thesis writing and presentations.

• To develop a learning of principles on ethical consideration involving research and issues related to plagiarism.

COURSE OUTCOMES

The subject aims to provide students with the necessary skills for,

- Understanding various aspects and ethics associated with research methodology.
- Research Problem identification, its implementation and evaluation.
- Applying to various research funding agencies which provide grants for the research projects.
- Defining of research problem and building hypothesis which will be helpful in industrial R&D projects.
- Knowledge of risk assessment and uncertainty associated with experimental modeling can be applied in industrial projects.
- Understanding of research deliverables in form of various publications, thesis writing
 and presentations and principles on ethical consideration involving research and issues
 related to plagiarism will help the candidate to design and work on an innovative and
 ethical research work.

Module 1	Introduction of Research Methodology	1 credit
Objectives	 To inculcate an understanding of research methodology To understand various principles of learning & 	
	theory based on it.	
	3. To know various government & other research funding agencies.	
	4. To understand various methods and sources of literature.	
Contents	Topics Covered	15

	Learning and instruction	5				
	Principles of Instructional design and learning					
	theory, Merrill's five principles and Gagne's					
	condition of learning. Active learning, group					
	learning, collaborative learning, problem-based					
	learning, team-based learning, Experiential					
	learning model of Kolb.					
	Basics of Research	6				
	Definition, objectives, motivation, types of					
	research and approaches: descriptive research,					
	conceptual, theoretical, applied and experimental.					
	Literature review	3				
	Important methods and sources to search for literature					
	(Primary and secondary sources), referencing and search					
	from Journals and Patents, Literature search using internet					
	and web based interfaces, suitable search engines,					
	advanced search techniques & data bases.					
	Review and compilation of the collected matter					
	Funding & Scholarship	1				
	Agencies funding research in pharmaceutical					
	sciences, Scholarship, types of scholarships in					
	education.					
Module 2	Basics of Research	1 credit				
Objectives	To learn about various assessment techniques.					
	2. To understand basics of research.					
	3. To study various research problems & develop					
	research plan					
	4. To learn planning, execution and implementation					
	of the schedule					

Assessment	3
Definition and methods, Georges Millers pyramid,	
Assessment, measurement and tests, Types of	
numbers, Formative and summative assessment.	

	Formation of Research Problem	6
	Research Process: To determine what type of research to	
	be done, plan of research work	
	Selection of research area, prioritization of research.	
	Objectives and scope of work, Developing Research Plan	
	and Schedule: Scheduling Constraints, steps, problems in	
	scheduling, limitations.	
	Implementation and Documentation	6
	Collecting the requisites of the experiments to be	V
	performed, maintaining the records of all the experiments,	
	maintenance of equipments/instruments and log books for	
	all the instruments, to come out with innovative ideas.	
Module 3	Mathematical Modelling & Analysis of Data	1
Wiodale 5	Manifest Manifest of Data	credit
Objectives	1. To acquaint research students with various	
	mathematical & experimental modeling	
	techniques used to draw conclusions in	
	Experimental Research.	
	2. To be able to identify, analyze and solve problems	
	related to research using software.	
	3. To study the various software used in pharmacy	
	for data analysis.	
	Mathematical Modeling and Simulation	5
	Concept of modeling, classification of mathematical	
	models, modeling with ordinary differential equations,	
	difference equations, partial differential equations, graphs,	
	simulation: concept, types (quantitative, experimental,	
	computer, fuzzy theory, statistical) processes of	
	compater, ruzzy meory, statistically processes of	

	Formulation of model based on simulation. Variables and	
	measurement	
	Experimental Modeling	5
	a) Definition of experimental design, examples,	•
	single factor experiments blocking and Nuisance	
	factors, guidelines for designing experiments.	
	b) General model of process: Input factors/	
	variables, Output parameters / variables	
	controllable / uncontrollable variables, dependent /	
	independent variables, experimental validity.	
	Analysis of data	5
	T	
	a) Types of data: parametric and nonparametric,	
	descriptive and inferential data, b) Collection of data, normal distribution, calculation of	
	b) Collection of data: normal distribution, calculation of co-relation coefficient	
	c) Data processing: analysis, error analysis, meaning,	
	and different methods: analysis of variance,	
	significance of variance, analysis of covariance,	
	multiple regression, testing linearity/nonlinearity of	
	model, testing adequacy of model.	
	d) Test to be used in data exploration and their choice	
	e) Introduction of software used in data analysis.	
Module 4	Ethics In Pharmacy & Research Deliverables	1
	•	credit
	To learn techniques used in the professional	
	presentations.	
	• To learn about research publications, thesis	
	writing and presentations.	

To understand ethical consideration in	volving
research and issues related to plagiarism	n.
Research Deliverables	6
a) Various Forms of Publication: Thesis, Pap	er,
Research proposal	
b) Thesis Writing: Introduction, Literature	Review or
State-of-the-Art, Research Approach (me	ethodology),
Results or findings, Discussions, Conclus	sions, Scope
for future work References, Appendices,	
c) Presentation: Poster, thesis, proposal, and	paper
Ethical issues in research	6
Historical perspectives, General pr	inciples on
ethical consideration involving	g human
participation, General ethical evaluation	on of drugs/
device/ diagnostics/ vaccines/ herba	l remedies.
Statement of specific principles	for human
genetics and genomic research. I	international
Conference on Harmonization. Go	od Clinical
Practices norms, Ethical principles rela	ited to
animal experiments.	
Plagiarism	3
Issues related to plagiarism, copy	right laws,
acknowledging the sources, format for	manuscript
writing, documentation, organization of	f reference
material, bibliography, end note.	

Assigned Reading/ References

- **1.** B.D. John, A.L. Brown and R.R. Cocking, 1999. "How People Learn: brain, mind, experience and school". Washington, DC: National Academy Press.
- **2.** J.R. Fraenkel, N.E. Wallen, 2008. "How to Design and Evaluate Research in Education", 7th Ed. Boston: McGraw-Hill.
- **3.** K.E. David, 2009. Curriculum Development for Medical Education: *A Six-Step Approach*, 2nd Ed. The John Hopkins University Press. ISBN 0-8018-9367-4.
- **4.** N. Peter, 2009. "Leadership: Theory and Practice." 3^{rd} *Ed.* Thousand Oaks: Sage Publications.
- **5.** G. Bordage, B. Dawson, 2003. Experimental study design and grant writing in eight steps and 28 questions. *Medical Education*, *37*(4): 376-385.
- **6.** B.J. Avolio, F.O. Walumbwa, T.J. Weber, 2009. Leadership: Current theories, research, and future directions. *Annual Review of Psychology*, 60: 421-449.
- **7.** C.R. Kothari, 2004. "Research Methodology". 2nd Ed. New Age International (p) Limited, Publishers.
- **8.** D. Montgomary, 2000. "Design of Experiments". 5th Ed. Wiley Interscience.
- **9.** K.P. Willkinsion, L. Bhandarkar, "Formulation of Hypothesis". 3rd Ed. Himalaya publishing, Mumbai.
- **10.** Schank Fr, 2008. "Theories of Engineering Experiments". 2nd Ed. Tata McGraw Hill.
- **11.** D.C. Montgomery, 2009. "Introduction to SQC" 6th Ed. John Willy & sons.

S3-[P&P]-5: RESEARCH SEMINAR

SEME	ESTER		SUBJ	ECT	
III Research Seminar					
WEEKLY ASSIGNMENT		CRE	DITS	MAI	RKS.
TH	PR	TH	PR	TH	PR
2		2		50	

COURSE OUTCOMES:

Research seminar aims to provide students,

- Ability to do literature survey on the research topic given by the research guide, interpret and compile the data into a scientific presentation.
- Ability to efficiently prepare more focused and professional power point presentation.
- Develop good communication skills
- Confidence to present information clearly and effectively.

S3-[P&P]-6: MINOR RESEARCH PROJECT

SEME	ESTER	SUBJECT			
III		Research Project			
WEEKLY ASSIGNMENT		CREDITS MARK		RKS.	
TH	PR	TH	PR	TH PR	
	24	-	8	200	

COURSE OUTCOMES:

Minor research project aims to provide students,

- Ability to do literature search, build a rationale, collect, analyze, interpret and evaluate the information that is related to the specific area of research.
- Ability to efficiently plan a research project.
- Applying the concept of research methodologies, methods and analytical techniques.
- Do research work independently in the laboratory.
- Efficiently solve the research problems.
- Ability to compile, present and defend the research report.

S3-[P&P]-1: INDUSTRIAL TRAINING

SEMESTER		SUBJECT				
III		Industrial Training				
ONE MONTH		CREDITS		MARKS.		
TH	PR	TH	PR	TH	PR	
		2		50		

COURSE OUTCOMES:

One month industrial training aims to provide students, industrial application of the theory and practical based research knowledge on various research areas of medicinal plants that the students gain through different subjects studied in the three semesters.

Semester IV

S4-[P&P]-1: RESEARCH PROJECT

SEMESTER		SUBJECT				
III		Research Project				
WEEKLY ASSIGNMENT		CREDITS		MARKS.		
TH	PR	TH	PR	TH	PR	
32		24			600	

COURSE OUTCOMES:

Minor research project aims to provide students,

- Ability to review scholarly literature collected from various scientific sources critically
 for the project and formulates a research rationale in the research area of medicinal
 plants.
- Efficiently conduct research to achieve the objectives.
- Propose new ideas/ methodologies or procedures in the research area of medicinal plants.
- Ability to compile the findings into a research thesis.
- Ability to prepare and present the research work.
- Ability to defend the research findings in front of scholarly audience.