

UNIVERSITY OF MUMBAI



Revised Syllabus

Program: M.Pharm

Semester I to IV

(Credit Based Semester and Grading System
with effect from the academic year 2013–2014)

M. Pharm. Credit Based System

Scheme Examination Semesters I to IV

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Syllabus Semesters I to IV

For ALL BRANCHES OF M. PHARM.

M. Pharm.

Semester I (For All Branches of Study)

No.	Title of the subject	Type of Course	Credits	Contact hours/week			ESE (hour)		Weightage (%)		
				L	I.L	P	T	P	CIA		ESE
		C/E/S							ST	IA	
1	Modern Pharm. & Med. Chem.	C	4	3	1	-	3	-	15	10	75
2	Modern Pharmaceutics	C	4	3	1	-	3	-	15	10	75
3	Modern Pharmacology	C	4	3	1	-	3	-	15	10	75
4	Modern Pharmaceutical Analysis	C	4	3	1	-	3	-	15	10	75
5	Study of Natural Products	C	4	3	1	-	3	-	15	10	75
6	Seminar	C	4		4	-	-	-	-	100	-

CIA Continuous Internal Assessment; **ESE** End Semester Examination; **ST** Sessional Tests; **IA** Internal Assessment, **L** Lectures, **I.L.** Integrated Learning involving Tutorials, Group Discussions, Assignments, Field Work; **P** Practicals, Lab. work, Project, **C** Core, **E** Elective, **S** Self Study.

Total Credits: Semester I – 24

In Semester I learners of all Branches will take the above 5 subjects, irrespective of their area of specialization in M. Pharm. The Evaluation pattern for every subject will be 25 marks for Continuous Internal Assessment (CIA) and 75 marks for the End Semester Examination. The 25 marks for CIA will be divided into 5 marks for Attendance and 5 marks for surprise quizzes and student-teacher interaction and 15 marks for a Periodic test held mid-semester.

Every learner will deliver a Seminar. This will be evaluated at the college level by a Committee consisting of the Principal, and faculty of the Department in which the student will be doing his research work. After the conduct of the Seminar the marks will be forwarded by the college/institution to the University.

Semester II (Branch wise)

No.	Title of the subject	Type of Course	Credits	Contact hours/week			ESE (hour)		Weightage (%)		
				L	I.L	P	T	P	CIE		ESE
		C/E/S							ST	IA	
1	Biostatistics and Research Methodology	C	4	3	1	-	3	-	15	10	75
2	Core Subject 1	C	4	3	1	-	3	-	15	10	75
3	Core Subject 2	C	4	3	1	-	3	-	15	10	75
4	Elective 1	E	4	3	1	-	3	-	15	10	75
5	Elective 2	E	4	3	1	-	3	-	15	10	75
6	Experimental Techniques in Pharmaceutical Sciences	C	4	-	-	8	6	-	15	10	75

CIA Continuous Internal Assessment; **ESE** End Semester Examination; **ST** Sessional Tests; **IA** Internal Assessment, **L** Lectures, **I.L.** Integrated Learning involving Tutorials, Group Discussions, Assignments, Field Work; **P** Practicals, Lab. work, Project, **C** Core, **E** Elective, **S** Self Study.

In the second semester, there will be one common paper Biostatistics and Research Methodology and a practical Experimental Techniques in Pharmaceutical Sciences for all the branches of specialization. At present the syllabus for Experimental Techniques in Pharmaceutical Sciences has been prepared for the main branches of specialization – namely Pharmaceutical Chemistry/Medicinal Chemistry, Pharmaceutics, Pharmacognosy, Pharmaceutical Analysis/Quality Assurance and Pharmacology. Students of other branches of specialization like Clinical Pharmacy, Medicinal Natural Products, Biopharmaceutics and Pharmacokinetics etc. can choose to do the exercises prescribed for any one of the above mentioned branches, preferably selecting something closest to their branch of specialization. Each Branch of Specialization will also have two Core subjects in the Branch of Specialization and two Electives that may be selected from the List of Electives specified in the Syllabus.

The ESE in the subject Experimental Techniques in Pharmaceutical Sciences will be conducted by the college/institution on behalf of the university. After the conduct of the examination, the marks will be forwarded by the college/institution to the University

Total Credits: Semester II – 24

Semesters III and IV

Research work related to the title of the thesis that has been registered with the University.

The learner should be allotted a Research Guide in Sem. I. The Guiding Teacher (Research Supervisor) along with the learner may plan the Research area to be pursued during Sems. III and IV. The title of the thesis should be communicated to the University before the commencement of Sem. III. The learner is expected to work a minimum of 40 hrs/week in Research to be entitled for 24 Credits each Semester. The Guiding Teacher (Research Supervisor) will sign a statement to this effect at the conclusion of Semesters III and IV, which may be communicated to the Controller of Examinations at the conclusion of each semester. Before completing the course, the learner will be required to give a Colloquium on the research work carried out by him/her during Sems. III and IV. The Colloquium will follow an open structure and will be assessed besides others by the Head of the Department, the Guide and the Principal of the College. A Statement that the learner has delivered a Colloquium must be sent to the University and will be mandated before the conduct of the *viva-voce* examination by the University.

Learners should be encouraged to attend conferences, seminars where they may present their research work, and to publish the findings of their research.

At the end of Sem. IV the learner will submit a thesis to the university. This will jointly be evaluated by the guiding teacher and an external examiner appointed by the university from academia or from the pharmaceutical industry. The evaluation will be for a total of 100 marks, of which 50 marks will be given by the guiding teacher and 50 marks by the external examiner. The parameters on which the marks will be given are: a. Literature Survey (10 marks) b. Presentation (8 marks) c. Methodology (7 marks) d. Results and Discussion (10 marks) and e. Viva-voce (15 marks). This makes a total of 50 marks to be given by the guiding teacher and by the external examiner separately.

Total Credits for M. Pharm. 96

Semester I

ALL BRANCHES

Modern Pharmaceutical and Medicinal Chemistry

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Drug Discovery	5
1.1	Historical perspective	1
1.2	Lead Discovery	1
1.3	Lead Modification – identification of the pharmacophore, functional group modification, privileged structures and drug-like molecules, modifications to increase potency and the therapeutic index, modifications to increase oral bioavailability	3
2	Receptors	10
2.1	Basic ligand concepts – agonist, antagonist, partial agonist, inverse agonist, efficiency and potency	1
2.2	Interactions (Forces) involved in drug-receptor complexes	2
2.3	Receptor theories – occupancy theory, rate theory and activation theory	1
2.4	Receptor classification – the four superfamilies	2
2.5	Receptor binding assays- measurement of K_d , B_{max} and IC_{50}	2
2.6	Topographical and stereochemical considerations in drug –receptor interactions	2
3	Prodrugs and Drug Delivery Systems	13
3.1	Enzyme activation of drugs, utility of prodrugs – aqueous solubility, absorption and distribution, site specificity, instability, toxicity, poor patient acceptability, formulation problems.	2
3.2	Carrier-linked prodrugs – carrier linkages for various functional groups, carrier-linked bipartite prodrugs, macromolecular drug carrier systems, tripartite prodrugs, mutualprodrugs, bioprecursor prodrugs (hydrolytic activation, elimination activation, oxidative activation, reductive activation, nucleotide activation, phosphorylation activation, sulfation activation and decarboxylation activation).	6
3.1	<i>Self study of specific examples of drugs that have been converted to prodrugs for solving problems related to ADME and their release mechanisms. Self study of prodrugs involving specific tissue targeting or specific activation at the target tissue.</i>	5
3.2		
4	Drug Metabolism	18
4.1	Introduction to xenobiotic/drug metabolism and its relation to other defence systems (Physical barriers, excretion, immune system).	0.5
4.2	Types of reactions (I and II), consequences of drug metabolism (DM) [inactivation, bioactivation, prodrugs], organs of DM, localization of drug metabolizing enzymes,	0.5

	factors affecting drug metabolism.	
4.3	Cytochrome P450s: Introduction to the family of enzymes, their classification and nomenclature.	1
4.4	CYP450 catalytic cycle, different types of reactions catalyzed by CYP450s and the mechanisms of catalysis.	4
4.4	Human CYP450s involved in DM, their distribution and properties, typical substrates, specific probe substrates, specific inhibitors, induction of CYPs and specific inducers	2
4.5	Discussion of glucuronosyltransferases, sulfotransferases, glutathione S-transferases, N-acetyl transferases, and FMO [on lines similar to that specified for CYPs as listed above].	4
4.5	<i>Self study of alcohol/aldehyde dehydrogenases, xanthine and aldehyde oxidase, epoxide hydrolase, esterases, azo and nitro reductases (reactions catalyzed by these enzymes, mechanisms of the reactions, typical substrates/inhibitors/inducers)</i>	6
5	Enzymes	14
5.1	Introduction to enzymes, binding site, specificity of enzyme catalyzed reactions and rate acceleration, Michaelis Menten kinetics and methods for plotting enzyme kinetic data	4
5.2	Mechanisms of enzyme catalysis – covalent catalysis, acid-base catalysis, electrostatic catalysis, some examples of the mechanisms of enzyme catalysis	2
5.3	Coenzyme catalysis – pyridoxal 5'-phosphate (racemases, decarboxylases, aminotransferases), nicotinamide and flavin (two-electron mechanism, one-electron mechanism and hydride transfers), folic acid and thiamine (one carbon transfer reactions).	4
5.1	<i>Self study of Hanes plot, Cornish-Eisenthal Bowden plot,</i>	1
5.2	<i>Self study of roles of coenzymes – biotin, coenzyme A, cyanocobalamin, vitamin K</i>	3
	Total	60

Books:

1. The Organic Chemistry of Drug Design and Drug Action, Silverman R. B., Academic Press.
2. Textbook of Drug Design and Discovery, Eds. Krogsgaard-Larsen P., Liljefors T., Madsen U., Taylor & Francis.
3. Drug Discovery – A History, Sneader W., Wiley.
4. Medicinal Chemistry: An Introduction, Thomas G, Wiley.
5. Drug Discovery – A History, Sneader W, John Wiley & Sons, Ltd.
6. Comprehensive Medicinal Chemistry, Series Ed., Hansch C., Pergamon Press.
7. Wilson and Gisvold's, Textbook of Organic Medicinal and Pharmaceutical Chemistry, Lippincott-Raven
8. Foye's Principles of Medicinal Chemistry, Lippincott Williams and Wilkins.
9. Drug Metabolizing Enzymes-Cytochrome P450 and Other Drug Metabolizing Enzymes in Drug Discovery and Development, Lee JS, Obach SR and Fisher MB, Marcel Dekker, Fontis India, 2003
10. Pharmaceutical Profiling in Drug Discovery for Lead Selection, Borchardt RT, Kerns EH, Lipinski CA, Thakker DR and Wang B, AAPS Press, 2004
11. Drug Metabolism – Current Concepts, Ionescu C and Cairns MR, Springer International Edition
12. Handbook of Drug Metabolism, Woolf TF, Marcel Dekker, 1999.

Unit	Course Contents (Topics)	Hours
1	Drug Stability:	9
1.1	Importance and need for stability testing	1
1.2	Revision of degradation pathways, kinetics, physical stability	2
1.3	Solution and Solid state stability, pH stability profiles, v and u graphs, package evaluation, ICH guidelines, statistical aspects in derivation of shelf life.	3
1.2	<i>Self study- Calculations for shelf life based on degradation kinetics</i>	3
2	Solubilization and Dissolution:	14
2.1	Importance of aqueous solubility of drugs, particularly NCEs, surfactant systems and phase diagrams, polymeric surfactants, cosolvents, complexation, solid state manipulations, cyclodextrins, drug derivatization, salt screening.	5
2.2	Revision of equations of dissolution and factors affecting dissolution, intrinsic solubility and dissolution rate, validation of testing, different equipments (emphasis on USP apparatus 4), Dissolution of TDDS, particulates, gels & ointments, comparison of profiles by f2 analysis, development of dissolution method, relevance of dissolution testing in ANDAs , bio-relevant media, BCS classification, IVIVC- study design and interpretation	5
2.3	<i>Self study- Calculations based on various solubility parameters and equations of dissolution. Pharmacopeial dissolution apparatus, data treatment of dissolution profiles.</i>	4
3	Excipients and introduction to polymers:	7
3.1	Role of excipients, purity, safety and toxicity with reference to routes of exposure-oral, inhalational, parenteral, others; regulatory aspects, risk assessments, Harmonization of excipient standards like residual solvents class 1,2,3.	2
3.2	Different classes of excipients - surfactants, special lipids, superdisintegrants, gelling agents, colours and flavours, sweetening agents, co-processed excipients.	2
3.3	Definition of polymers, classification; concept of properties used in characterisation, methods of polymerisation, biocompatibility evaluation, applications	2
3.4	<i>Self study: sources and brand names of various excipients</i>	1
4	Optimisation Techniques:	8
4.1	Definition, Need, Advantages, description of terms such as independent variable, response parameters, response surface, contour plots, polynomial equations	2
4.2	Simplex and factorial designs in optimisation	3
4.3	Application of optimisation techniques in QbD in product development	1
4.5	<i>Self study: Placket-Burman design, central composite designs</i>	2
5	Preformulation:	12
5.1	Scope of Preformulation-Role & importance in New Drug Discovery & Approval process-Lead optimization, Steps in Designing the preformulation evaluation of a new drug, critical issues and	3

	problems/constraints	
5.2	Key Areas in Preformulation research- Bulk Characterization, Solubility Analysis, Stability Analysis, Compatibility with common excipients	4
5.3	Preformulation aspects for Tablets, Injectables, Liquid preparations, Protein & peptide drugs.	3
5.4	<i>Self study: case study of drug exhibiting various polymorphic forms, drug excipient compatibility</i>	2
6	Powder Technology (Micromeritics):	10
6.1	Revision of following topics: <ul style="list-style-type: none"> • Important definitions & Units • Importance of particle size in pharmaceutical development. • Fundamental & derived properties of powders • Particle size reduction –comminution mechanisms & equipments • Methods of particle size determination (emphasis on basic principles & interpretation of data) 	2
6.2	Comminution- Theory of comminution, milling rate (various mathematical relationships), concept of milling/grinding index, energy for comminution, distribution and limit of comminution	2
6.3	Compaction of powders- definitions of compression & consolidation, deformation mechanisms of matter, steps in compaction of tablets (in detail), theoretical aspects- Force Volume relationships/porosity –pressure equations (Heckel’s Law & equation), Granulation of powders –theory, Effect of compaction pressure on various tablet properties, Energy for compaction & effect of lubrication of granules, instrumentation of tablet presses (principles)	3
6.4	<i>Self study: case studies on compaction behaviour of two excipients</i>	3
	Total	60

Books :

1. Drug Stability Principles and Practices by Carstensen J, Marcel Dekker, 3rdedn, Vol 107, 1990.
2. Pharmaceutical Stress testing by Baertschi SW, Taylor and Francis, Vol 153, 2005.
3. Pharmaceutical characterisation of Pharmaceutical Solids by Brittain HG, Marcel Dekker, Vol 70, 1995
4. Preformulation in Solid Dosage Form Development by Adeyeye MC, Brittain HG, Informa Healthcare, Vol 178, 2008.
5. Dissolution, Bioavailability and Bioequivalence by Abdou HM, Ed A. Gennaro, B. Migdalof, Mack Printing Company, 1st edn, 1989.
6. Pharmaceutical Bioequivalence by Welling PG, Francis LST, Dighe SV, Marcel Dekker, Inc., Vol. 48, 1991.
7. Pharmaceutical Dissolution Testing by Banaker U, Marcel Dekker, Vol 49, 1992.
8. Excipient toxicity and safety by Weiner M L, Kotkoski LA , Vol 103, Marcel Dekker, 1999.
9. Martin's Physical Pharmacy and Pharmaceutical Sciences, by Sinko PJ, Ed Lea & Feiger, Lippincott Williams & Wilkins, 6thedn, 2010.
10. Modern Pharmaceutics by Banker GS, Ed Banker GS & Rhodes CT, Marcel Dekker, 4th edn, Vol 121, 2003.
11. Pharmaceutical Statistics by Bolton S, Marcel Deckker, 3rdedn, Vol 80, 1997.
12. The Theory and Practice of Industrial Pharmacy by Lachman L, Lieberman HA, Kanig JL, Varghese Publishing House, 3rdedn, 1990.
13. Pharmaceutical Dosage Forms: Tablets, Unit Operations and Mechanical Properties Ed Augsburger LL, Hoag SW, Informa Healthcare USA, Inc., 3rdedn, Vol 1, 2008.
14. Techniques of Solubilization of Drugs by Yalkowsky SH, Marcel Dekker, Vol 12, 1985.
15. Pharmaceutical Dissolution Testing by Dressman J. Ed Dressman J, Kremmer J, Tylor & Francis, 2005.
16. Controlled Drug Delivery: Clinical Applications, by Bruk SD, CRC Press Inc., Vol 2, 1983.
17. Handbook of Pharmaceutical Granulation Technology by Parikh DM , Informa healthcare, 2nd edition, Vol 154, 2007.
18. Pharmaceutical Powder Compaction Technology by Alderborn G, Nystrom C, Marcel Dekker, Vol 71, 1996.

Modern Pharmacology

4 hrs/week

Unit	Course Contents (Topics)	Hours
1		11
1.1	Drug Absorption, distribution, metabolism and excretion.	5
1.2	<ul style="list-style-type: none"> Mechanisms of transport of drug across membranes. Transporters involved in drug absorption, distribution and excretion processes. 	3
1.3	<ul style="list-style-type: none"> <i>Self study-Drug efflux pathways and experimental methods to study drug transport.</i> <i>Pharmacokinetic factors affecting drug action</i> 	3
2	Mechanism of drug action	11
2.1	Classification of receptors and description of each class with examples.	1
2.2	<ul style="list-style-type: none"> Signal transduction mechanisms. Detailed description of signal mediation through cascades after adrenergic, muscarinic, GABAergic, insulin receptor stimulation. 	4
2.3	Regulation of receptors, their involvement in various biological processes including diseases resulting from receptor malfunction and their role in pharmacotherapeutics.	1
2.4	Regulation of intracellular calcium.	2
2.5	Pharmacodynamic interactions in a multicellular context e.g. Vascular wall (interactions of physiological ligands and drugs in pathophysiological setting).	1
2.6	<i>Self study- Classification and characterization of receptors-IUPHAR (Eg. 5-HT receptors)</i>	2
3	Functions of sodium and potassium channels and therapeutic potential of channel modulators.	3
4	<p>Factors affecting drug responsiveness.</p> <ul style="list-style-type: none"> Alteration in concentration of drug that reaches receptors. Variation in concentration of an endogenous receptor ligand. Alteration in number and function of receptors. Clinical selectivity: Beneficial vs. toxic effects of drugs. <p>a. Beneficial and toxic effects mediated by the same receptor - effector mechanism.</p> <p>b. Beneficial and toxic effects mediated by identical receptors but in different tissues or by different effector pathways.</p> <p>c. Beneficial and toxic effects mediated by different types of receptors.</p> <ul style="list-style-type: none"> Desensitization, tachyphylaxis. Drug tolerance. 	3
5	Cellular and molecular mechanisms of	4
5.1	Drug dependence (Eg. Morphine).	
5.2	Microbial resistance.	
6	Advances in therapy of	18
6.1	CNS: Depression, Alzheimer's disease, Psychosis, Parkinson's disease, Epilepsy.	5

6.2	CVS: Hypertension, Angina Pectoris, Congestive cardiac failure, Arrhythmia.	5
6.3	Management of Diabetes Mellitus.	2
7	Apoptosis	4
7.1	Molecular biology, physiological, pharmacological implications and therapeutic prospects.	2
7.2	<i>Self study – Interaction between cell, growth factors and extracellular matrix.</i>	2
8	Immunopharmacology	6
8.1	Introduction to immunopharmacology, immunomodulators, Immunostimulants and Immunosuppressants.	4
8.2	<i>Self study-Autoimmunity</i>	2
	Total	60

Books:

1. Rang and Dale's pharmacology-- Elsevier Churchill Livingston.
2. Lange's Basic and clinical pharmacology, Katzung B.G. Masters S.B., Trevor A.G. Tata McGraw Hill.
3. Goodmann and Gilman's pharmacological basis of therapeutics, Edited by Laurence Brunton, Bruce Chabner and Bjorn Knollman, McGraw Hill.
4. Pharmacological reviews, Annual reviews Inc.
5. Advances in pharmacology, Academic Press.
6. Trends in Pharmacological Sciences, Cell Press Elsevier Publication.

Modern Pharmaceutical Analysis

4 hrs/week

Unit	Course contents (Topics)	Hours
1	Multicomponent analysis of drugs using UV- Vis. spectroscopy:	6
1.1	Simultaneous equation method, Absorbance ratio method, Difference spectroscopy, Derivative spectroscopy and Introduction to Ratio derivative spectroscopy,	4
1.2	<i>Self study-Pharmaceutical applications of above techniques (1.1)</i>	2
2	F.T.I.R spectroscopy:	6
2.1	Construction and working, Newer sampling techniques.	2
2.2	Interpretation of I.R. spectra in mid I.R. region (aliphatic and aromatic compounds for simple compounds such as amines, alcohols, amides, nitriles, ketones, aldehydes, esters, acids, nitro and anhydrides).	2
2.3	<i>Self study-Interpretation of recorded I.R spectra of drugs and organic compounds.</i>	2
3	NMR spectroscopy:	10
3.1	¹H- NMR: Basic theoretical concepts-(<i>Self study-chemical shift, splitting pattern and coupling constant-2 hrs</i>), Non-first order spectra, methods to make complex spectra simple, FT-NMR.	6

3.2	¹³C-NMR: Theory and Principle.	2
3.3	Applications of 2D-NMR (only COSY and HETCOR)	2
4	Mass Spectrometry:	10
4.1	Different ionisation techniques-EI, CI, FD, FI, MALDI, API (APPI, APCI, ESI).	4
4.2	Different analysers-Quadrupole, TOF, QTOF, Ion cyclotron, Ion trap.	2
4.3	Concepts for interpretation of mass spectra-Molecular ion peak, base peak, Isotope abundance, fragmentation pathways- α fission, β fission, MacLaffarty rearrangement, Retro Diels Alder rearrangement, Tandem mass (MS-MS).	4
5	Terminologies of chromatography: <i>Self study-Theoretical plate, HETP, Plate theory, Rate theory, Van Deemter equation, Isocratic elution, Gradient elution, capacity factor, selectivity factor, Resolution, tailing factor, asymmetry factor.</i>	3
6	Advances in chromatography:	11
6.1	HPLC-Ion pair chromatography, Chiral chromatography (Chiral stationary phases, use of mobile phase additives, precolumn derivatisation, chiral detectors), UPLC, <i>Self study-Advances in HPLC detectors (1 hr).</i>	5
6.2	Supercritical Fluid chromatography-Principle, Instrumentation and pharmaceutical applications.	2
6.3	<i>Self study-HPTLC-Principles, Instrumentation and applications including fingerprint analysis.</i>	1
6.4	Gas chromatography-Headspace analysis.	1
6.5	Gel electrophoresis-Principle, Instrumentation and applications.	2
7	Hyphenated techniques:	4
7.1	Interfaces used in and applications- GC-MS, LC-MS, LC-MS-MS	3
7.2	Introduction to LC-NMR and MALDI-TLC.	1
7	Thermoanalytical techniques: Principle, instrumentation and applications including interpretation of data in pharmacy for:	5
7.1	<i>Self study-DSC and TGA</i>	3
7.2	TMA (Thermo mechanical analysis).	1
7.3	<i>Interpretation of DSC and TG curves of suitable compounds/drugs (Self study)</i>	1
8	Microscopy: Principle, Instrumentation, sample preparation and pharmaceutical applications of- Scanning Electron Microscopy, Transmission Electron Microscopy, Atomic Force Microscopy,	5

	Confocal microscopy.	
		Total 60

Books:

1. Chromatographic methods by A.Braithwaite & S.J.Smith, Kluwer Academic publishers, Netherlands, London, USA.
2. Thermal Analysis of Pharmaceuticals by Craig, Informa, CRC Press, Indian Reprint.
3. Practical Pharmaceutical Chemistry by A.H.Beckett and J.B.Stenlake, fourth edition, part two, CBS Publishers and Distributors.
4. Spectrometric Identification of Organic compounds by R.M.Silverstein, F.X.Webster,D.J.Kiemle , Latest edition, John Wiley & Sons
5. Applications of absorption spectroscopy of organic compounds by John Robert Dyer
6. Organic Spectroscopy by William Kemp, PALGRAVE.
7. Textbook of Pharmaceutical Analysis by K.A.Connors, Wiley Interscience Publications.
8. Introduction to Spectroscopy by D.L.Pavia, G.M.Lampman & G.S.Kriz.
9. Remington: The Science & Practice of Pharmacy, 20th edition, Vol. 1, Lippincot Williams & Wilkins
10. Introduction to Modern Liquid Chromatography by L.R.Snyder, J.J.Kirkland 3rd edition.
11. Chiral separations by Liquid Chromatography and Related Technologies Chromatographic Science Series by Hassan Y., Imran Ali, Vol. 90.
12. Static head space gas chromatography Theory & practice by Bruno Kolb & L.S.Ettre.
13. Encyclopedia of Chromatography, by Jack Cazes, 3rd edition, Vol.1,2 & 3.
14. Online LC-NMR and Related techniques by Klasu Albert, John Wiley & Sons
15. LC-MS- A Practical Users guide, by Marvin C. McMaster.

Study of Natural Products

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Introduction to study and research in herbal drugs:	4
1.1	Different approaches to plant selection, collection and processing for herbal drug research (Random selection, Use of ethnobotanical information, Use of chemotaxonomical classification etc).	2
1.2	Recent advances in concept of authentication & standardization - significance of chemotaxonomy and DNA finger-printing with respect to gene expression for secondary metabolites.	2
2	Extraction of phytochemicals	18
2.1	Concepts of extraction with respect to activity guided fractionation & isolation of Markers/Biomarkers.	2
2.2	Recent trends in extraction, optimization of extraction, and analysis of the phytochemicals of different classes.	2
2.3	Detail discussion of large scale extraction of the following: (1) Opium alkaloids (2) Piperine (3) Sennosides (4) Caffeine (5) Cinchona alkaloids (6) Rutin (7) Lemon grass oil (8) Patchouli oil (9) Steroids (Diosgenin from all sources)	9

2.4	<i>Self study- preparation of flow chart and discussion of physicochemical principles for all large scale extractions</i>	5
3	Natural products in drug discovery and drug development	8
3.1	Role of natural products as leads to the design of new drugs with case history with examples e.g., artemisinin, taxane, camptothecin and a few others.	2
3.2	Natural products derived combinatorial libraries and their significance in drugs discovery programme (HITS and leads).	2
3.3	<i>Self study- Discussion of lead molecules in drug discovery</i>	4
4	Study of following excipients of natural origin in NDDS with respect to sources, preparation, composition and application	16
4.1	Natural dyes & colorants, sweeteners, flavours and fragrant materials	8
4.2	Kappa carrageenans, galactomannans, glucomannans, cellulose derivatives, lecithin, & alginates.	4
4.3	<i>Self study- Role of excipients mentioned above, in formulations, with examples</i>	4
5	Application of immunoglobulins from plant sources in diagnosis and therapy.	4
6	Nutraceuticals and their role in health care.	4
6.1	Study of following classes of herbs with two or three suitable examples of each: (1) Antioxidants (2) Immunomodulators (3) Antihyperglycemics (4) Hepatoprotectives	4
7	Status of natural products in official books	6
7.1	Introduction to Herbal Pharmacopoeias of different countries	2
7.2	Monographs of natural products in other official books.	2
7.3	<i>Self study-Discussion of monograph of few substances of natural origin</i>	2
	TOTAL HOURS	60

Books:

1. Pharmacognosy Phytochemistry – Medicinal Plants- Jean Brunetton, Lavoisier Publishing, Paris.
2. Text book of Pharmacognosy- Trease and Evans- 14th edition. Elsevier science
3. Transgenic Plants- R. Ranjan- Agro Botanica, New Delhi.
4. Transgenic Plants-A Production system for Industrial and Pharmaceutical Proteins. by Meran Owen, Jan Pen- John Wiley.
5. Medicinal Plant-Their Bioactivity, Screening and Evaluation- CSIR.
6. Homeopathic Pharmacopoeia of India- Publisher Ministry of Health.
7. The Ayurvedic Formulary of Part I & II- Publisher Ministry of Health.
8. Chinese Materia Medica-You-Ping Zhu- Harwood Academic Publishers.
9. India Materia Medica- Nadkarni A.K. –Bombay Popular Prakashan.
10. Phytochemical Methods - J.B.Harbone - Chapman and hall
11. Cultivation's and Processing of Medicinal Plants-Ed. by L. Hornok-John Wiley.
12. Introduction to Flavonoids-Bohrn Bruce A. – Herwood Academic Publishers.
13. Cultivation and Utilization of Aromatic plants – Ed. By Atal C. K. and Kapur B.M.- CSIR.
14. Plant Tissue and Cell Culture Ed. H.E. Street – Blackwell Scientific publications.
15. Aflatoxin- Leo A. Goldblatt- Academic Press New York.
16. Microbial Toxins- Ciejler, Kadis and Ajl- Academic press.

17. Antimicrobial in Food – Alfred Larry Branen, P. Michael Davidson Publishing house
18. Chemical plant Taxonomy T. Swain, 1963. Academic Press, London.
19. Plant Taxonomy and Biosystematics .C.A Stace, 1985. Edward Arnold, London.
20. Modern methods of plant analysis K. Paech, 1956., Springer-Verlag.
21. Indian Herbal Pharmacopoeia, Vol. 1&2, RRL, IDMA, 1998, 2000.
22. Indian Pharmacopoeia, 2010.
23. Standardization of Botanicals,V. Rajpal, 2002. Eastern Publishers, New Delhi.
24. Natural Compounds as Drugs – Vols. I & II, Editor- Frank Petersen, René Amstutz, Die Deutsche Bibliothek, Germany.
25. Quality control of Herbal Drugs: An Approach to evaluation of Botanicals, Pulok Mukherjee - Riddhi International
26. Chemicals from Plants: Perspectives on Plant Secondary Product, Walton & Braun, Imperial College Press.
27. Towards Natural Medicine Research in the 21st Century H. Ageta, N. Aimi et al Excerpta Medica, International Congress Series 1157.

SEMESTER II

FOR ALL BRANCHES

Biostatistics and Research Methodology

4 hrs/week

Unit	Course Content (Topics)	Hours
	Biostatistics	
1	Collection and Organization of data	8
1.1	Graphical and pictorial presentation of data	1
1.2	Measures of central tendency and dispersion	2
1.3	Variance and standard deviation, relative error, coefficient of variation, precision and accuracy	2
1.4	Sampling techniques: simple random sampling; stratification; estimation of the mean and proportion.	3
2	Probability	6
2.1	Definition. Conditional probability and Bayes' theorem. Probability distributions: binomial, multinomial and Poisson distributions. Normal and lognormal distributions. Use of normal distribution tables.	6
3	Regression	6
3.1	Linear regression and correlation, curvilinear regression, method of least squares, curve fitting, Fiducial limits, probit and logit analysis	6
4	Parametric tests	8
4.1	Testing hypothesis, Types of error. Level of significance. Significance tests and p-value	4
4.2	Tests of significance based on normal distribution, test of significance for correlation coefficients, confidence interval for mean and regression proportion.	4
5	Nonparametric tests	4
5.1	Nonparametric procedures: Chi square goodness of fit test, sign test, Mann-Whitney test; Wilcoxon signed rank test.	4
6	Experimental designs	8

6.1	Randomization, completely randomized, randomized block and Latin square designs, factorial design, cross over and parallel designs	4
6.2	Students should learn use of Minitab / R Software for data summary, correlation, regression analysis, test of hypothesis and experimental design	4
	Total Biostatistics	40
	Research Methodology	
7	Objectives and purpose of Research	2
7.1	Types of research – Educational, clinical, experimental, basic, applied and patent oriented research	2
8	Literature survey	2
8.1	use of library, books and journals, eJournals, retrieving patents and seeking reprints.	2
9	Methods and tools used in research <ul style="list-style-type: none"> • Qualitative and quantitative studies • Simple data organization, descriptive data analysis • Limitations and sources of errors • Inquiries in form of questionnaire, opinionaire or by interview • Statistical analysis of data including variance, standard deviation, standard error, mean, student's <i>t</i> test and annova, correlation of data and its interpretation, computer data analysis 	6
10	Scientific writing and reporting <ul style="list-style-type: none"> • Different types of research papers • Title and author names • Abstract and key words • Methodology 	3
11	Scientific Presentation <ul style="list-style-type: none"> • Importance, types and different skills • Content, format of model, introduction and ending • Skills for oral presentation and types of visual aids • Questionnaire 	3
12	Patents and Trade marks <ul style="list-style-type: none"> • The Indian patent system • Present status of intellectual property rights (IPR) • Product patents and process patent • Requirements and preparation of patent proposal • Registration of patent in foreign countries 	4
	Total Research Methodology	20
	Total (Biostatistics and Research Methodology)	
		60

Books:

1. Pharmaceutical Statistics – Practical and Clinical Applications, Bolton S., Marcel Dekker, Inc. N., USA
2. Biostatistics: A Foundation for Analysis in Health Sciences, Wayne W Daniel, John Wiley & Sons, Inc.
3. Introduction to Statistical Analysis, Dixon W. J. and Massey F. J., McGraw Hill, N.Y., USA.

4. Statistical Methods, Snedecor G. W. and Cochran W. G., Iowa State University Press, Ames, Iowa.
5. Research in Education, John W Best and James V Khan, Prentice Hall of India Pvt. Ltd.
6. Effective Business Report Writing, Brown Leland, Prentice Hall Inc. India.
7. Presentation Skills, Michael Hatton, Indian Society for Technical Education, New Delhi.
8. Thesis and Assignment writing, Anderson Jonathan and Durston Berry H, Wiley Eastern Ltd., Bangalore.
9. Writing a Technical Paper, Donald H Menzel, McGraw Hill Book Company, Inc., New York.

Semester II

BRANCH: PHARMACEUTICAL CHEMISTRY

Core 1

Advanced Pharmaceutical and Medicinal Chemistry

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Enzyme Inhibition	16
1.1	Coverage of basic aspects of enzyme kinetics, catalysis, transition-state theory.	2
1.2	Drug Resistance through alterations of drug uptake, overproduction of enzyme, alterations of the enzyme active site, overproduction of the substrate or new pathways for formation of the product	1
1.3	Drug synergism, concepts and mechanisms.	1
1.4	Reversible enzyme inhibitors – competitive inhibition, non-competitive inhibition, uncompetitive inhibition with suitable examples. Detection of type of inhibition by suitable plotting methods. Concepts of IC_{50} and K_i .	4
1.5	Slow-tight binding inhibitors, covalent enzyme inhibitors and mechanism-based inhibitors with suitable examples. Concept of K_{inact} and K_i for irreversible inhibitors	4
1.6	<i>Self study of specific examples of different types of inhibitors and their design (some examples like COX inhibitors, ACE inhibitors, RT inhibitors, HIV protease inhibitors, aromatase inhibitors, DHFR inhibitors, viral DNA polymerase inhibitors, thymidylate synthase inhibitors and others)</i>	4
2	QSAR	14
2.1	Historical Aspects	1
2.2	Electronic Effects- the Hammett equation, lipophilic effects, experimental measurement of lipophilicity, logP and logD, effect of ionization on logP, calculation of logP and logD, Steric effects- the Taft equation	3
2.3	Hansch Analysis, Free-Wilson method, Topliss operational scheme	2
2.4	Basics of regression analysis - linear and multilinear regression, introduction to PCA, PCR, PLS, ANN and GFA. Correlation coefficients (r^2 and r^2_{pred}), F-test, standard error, validation methods like cross-validation by calculation of q^2 , boot-strap analysis and randomization. Application domain for predictions using a QSAR model.	6
2.5	Design of training and test sets using factorial design	2
2.5	<i>Self Study – Different types of descriptors reported in literature that account for the steric, electronic and lipophilic effects.</i>	2
3	Peptides and Peptidomimetics	14
3.1	Coverage of peptide structure, biosynthesis of peptides and solid-phase/solution synthesis of peptides.	4
3.2	Design of peptidomimetics by manipulation of the amino acids, modification of the peptide backbone, incorporating conformational constraints locally or globally, α -helix, β -sheet, β -and	4

	γ -turn mimetics	
3.3	<i>Self study of examples of peptidomimetics for some enzymes and receptors like ACE, CCK, bradykinin</i>	2
4	Antisense therapeutic agents	6
4.1	History and principles	2
4.2	Design of antisense oligonucleotides and small interfering RNAs (siRNAs) with some examples	4
5	Molecular Biology, Genetic engineering and Biotechnology in production of biologicals as drugs.	6
5.1	<i>Self study of biotechnology based drugs, vaccines and diagnostic agents with respect to their biological source, their design and the mechanism of their actions</i>	4
	Total	60

Books

1. The Organic Chemistry of Drug Design and Drug Action, Silverman R. B., Academic Press.
2. Textbook of Drug Design and Discovery, Eds. Krosggaard-Larsen P., Liljefors T., Madsen U., Taylor & Francis.
3. Medicinal Chemistry: An Introduction, Thomas G, Wiley.
4. Peptide and Protein Design for Biopharmaceutical Applications, Ed Jensen K. J., Ch. 3 Aspects of Peptidomimetics by Maes V., Tourwé D., John Wiley & Sons, Ltd, Chichester, UK.
5. Comprehensive Medicinal Chemistry, Series Ed., Hansch C., Pergamon Press.
6. Burgers Medicinal Chemistry, Drug Discovery and Development, Wiley.

Core 2

Advanced Organic Chemistry

4 hrs/week

Unit	Course Content (Topics)	Hours
1.0	Advanced Stereochemistry	12
1.1	<i>Self Study - Coverage of the basic concepts in stereochemistry – optical activity, specific rotation, racemates and resolution of racemates, the Cahn-Ingold-Prelog sequence rule, meso compounds, pseudo asymmetric centres, pro-R, pro-S, axes of symmetry, Fischers D and L notation, cis-trans isomerism, exo-endo, syn-anti nomenclature. Stereoselective and stereospecific reactions. Conformational isomerism in acyclic systems. Shape of six membered rings and effect of substituents and reactivity.</i>	2
1.2	Chirality in systems lacking a stereogenic carbon atom	2
1.2.1	Point chirality – tertiary amines and phosphines	1
1.2.2	Axial chirality – allenes, biphenyls and binaphthyls	1

1.2.3	Helical structures – polynucleotides, polyamino acids, biaryls and allenes	1
1.3	Methods for estimating ratios of stereoisomers in a mixture, separation and identification of the individual components by NMR spectroscopy, X-ray crystallography.	1
1.4	Nucleophilic attack on acyclic carbonyl compounds – Cram’s rule, Felkin-Ahn rule. Locking effects in nucleophilic reactions at carbonyl groups	2
1.5	Stereochemistry of important reactions leading to formation of alkenes – Wittig and related reactions	2
2.0	Catalysis & Organometallics in Organic Synthesis	12
2.1	Types of catalysis, heterogeneous and homogenous catalysis, advantages and disadvantages, catalytic cycles	1
2.2	Heterogeneous catalysis – preparation, characterization, kinetics, supported catalysts, catalyst deactivation and regeneration, some examples of heterogeneous catalysis used in synthesis of drugs.	1.5
2.3	Homogenous catalysis, hydrogenation, hydroformylation, hydrocyanation, Wilkinson catalysts, chiral ligands and chiral induction, Ziegler-Natta catalysts, some examples of homogenous catalysis used in synthesis of drugs	1.5
2.4	Phase transfer catalysis - theory and applications	1.5
2.5	Introduction, Classification of organometallic compounds based on hapticity and polarity of the M-C bond. Nomenclature and general characters. Synthesis, stability and decomposition pathways.	1.5
2.6	Transition metal π -complexes with unsaturated organic molecules, carbon monoxide, alkenes, alkynes, allyl, dienes, cyclopentadienyl, arene complexes, preparation, properties, nature of bonding and structural features, important reactions relating to nucleophilic attack on ligands and to organic synthesis. Basic organometallic reactions covering oxidative reactions, migratory reactions, insertions, extrusion, additions, eliminations – their mechanisms and stereochemistry.	3
2.7	<i>Self Study - Basic organometallic reactions covering oxidative reactions, migratory reactions, insertions, extrusion, additions, eliminations – their mechanisms and stereochemistry</i>	2
3.0	Synthon Approach and Applications	13
3.1	Retrosynthesis and its advantages, rules for dissection of molecules, meaning of the term, disconnection, FGI, FGA and synthons, guidelines for the order of events	1
3.2	C-X disconnections; C-C disconnections – alcohols and carbonyl compounds; 1,2-, 1,3-, 1,4-, 1,5-, 1,6-difunctionalized compounds	4
3.3	Strategies for synthesis of three, four, five and six-membered rings	3
3.4	Strategies for synthesis of aromatic and saturated heterocycles	3
3.5	<i>Self Study – Strategies for synthesis of saturated heterocycles.</i>	2
4.0	Asymmetric Synthesis	6

4.1	Introduction and need; chiral synthesis using chiral pool, chiral auxiliaries, chiral catalysts	2
4.2	Enzymes, chiral solvents and whole organisms	2
4.3	Analytical methods of determining purity of stereoisomers	1
4.4	<i>Self Study - Applications in industry</i>	1
5.0	Combinatorial Chemistry	11
5.1	Introduction, advantages and planning combinatorial synthesis	1
5.2	Solid phase and solution phase synthesis	5
5.3	Supports, linkers, and tags	1
5.4	Deconvolution and iteration	1
5.5	Parallel synthesis, multistep – convergent and sequential synthesis	2
5.6	<i>Self Study – Multicomponent reactions</i>	1
6.0	Green Chemistry	5
6.1	History, need and the goals of green chemistry	1
6.2	Basic principles of green chemistry, illustrated with examples to discuss issues of prevention of waste or minimize by-products, atom economy, prevent and minimize formation of hazardous or toxic products, design of safer chemical equivalents, selection of appropriate solvents, media, separation agents, improve economy and efficiency of reactions, by use of microwaves, ultrasound etc., and use of renewable starting materials.	3
6.3	<i>Self Study – Reactions carried out using Microwave and ultrasound.</i>	2
	Total	60

Books

1. Stereochemistry of carbon compounds, Eliel E, Wilen S H, Manden L N, Wiley.
2. Stereochemistry of Organic Compounds, Nasipuri D, Wiley Eastern.
3. Advanced Organic Chemistry, Carey FA and Sundberg RJ, Part A and B, Springer
4. Introduction to Green Chemistry, Ryan M. A., Tinneland M., American Chemical Society (Washington).
5. Combinatorial Chemistry; Synthesis and Application, Eds., Wilson S. R. Czarnik A. W., Wiley: New York.
6. Organic Chemistry, Clayden J, Greeves N, Warren S, Wothers P, Oxford University Press.
7. Stereoselective Synthesis, Atkinson R S, John Wiley & Sons.
8. The Organometallic Chemistry of the Transition Metals, Crabtree R. H., John Wiley
9. Transition Metals in Synthesis of Complex Organic Molecules, Hegedus L., University Science Books.
10. Homogenous Transition Metal Catalysis, Masters C., Chapman & Hall.
11. Principles and Practice of Heterogenous Catalysis, Thomas J. M., Thomas M. J., John Wiley
12. Principles of Asymmetric Synthesis, Gawley R. E., Aubrey J, Elsevier.
13. Greene's Protective Groups in Organic Synthesis, Wuts, P. G. M., Green T. W., Wiley
14. Organic Synthesis – The Disconnection Approach, Stuart, W., Wiley.
15. The logic of chemical synthesis, Corey E J and Cheng X-M, John Wiley and Sons.

Semester - II

BRANCH: PHARMACEUTICS

Core 1 **Advanced Pharmaceutics – I**

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Solids – oral SR system	13
1.1	<i>Self study-Overview of Single oral unit SR systems .</i>	3
1.2	<i>Self study-Structure and physiology of GIT.</i>	1
1.3	Mechanism of Release & Release kinetic equations. Types – Diffusion controlled, Dissolution controlled, Reservoir, Matrix, Osmotic systems, Ion exchange systems Mucosal drug delivery systems- buccal, gingival, sublingual.	4
1.4	Multiparticulate systems-pelletization (emphasis on extrusion and spheronization). Orodispersible systems. Pulsatile Drug delivery systems.	5
2	Parenteral SR systems	12
2.1	Need and concept, routes employed	1
2.2	Approaches- aqueous systems (complexation, use of polymers), aqueous suspensions (depot injections, microspheres, magnetic microspheres), Oily solutions & suspensions, Emulsions (Microemulsions, multiple emulsions,), Implants (in detail-concept, properties desired, various approaches), prodrugs (chemical modifications), infusion pumps.	6
2.3	<i>Self study-Biopharmaceutical aspects, Sterilization & stability issues</i>	2
2.4	Characterization-special emphasis on release studies	1
2.5	Issues related to Safety, Toxicity & Tissue Injury	2
3	Specialized Emulsions	9
3.1	Microemulsions, Multiple emulsions, Self Emulsifying Drug Delivery systems & SMEDDS; Formulation and phase behaviour; Preparation & Characterization; Bioavailability Aspects; Applications.	6
3.2	<i>Self study-Theories of Emulsification, Factors influencing type of emulsion formed.</i>	3
4	Gastro-retentive Drug Delivery Systems	8
4.1	<i>Self study –Introduction; concept of absorption window; need for GRDDS, gastric motility; principles of Gastro-retention; Factors controlling performance of GRDDS.</i>	3
4.2	Different Approaches- High density systems, floating systems, muco-adhesive systems, Expandable systems, Magnetic systems, Superporous Hydrogels	4

4.3	Evaluation.	1
5	Ocular drug delivery systems.	7
5.1	<i>Self study-Structure and physiology of eye; Drug absorption and disposition in the eye.</i>	2
5.2	Methods to prolong ocular drug residence with emphasis on mucoadhesive systems.	1
5.3	Intraocular inserts; Nonerodible inserts / Erodible inserts. Novel ophthalmic drug delivery systems, Nanoparticles , liposomes and prodrugs. Ocular penetration enhancers.	4
6	Transdermal Drug Delivery Systems	7
6.1	<i>Self study-Structure and physiology of skin.</i>	1
6.2	Principles of skin permeation. Kinetics of skin permeation & penetration enhancers. Types (Gels, Patches/films) , Pressure sensitive adhesives.	3
6.3	Development & evaluation – <i>in vitro, in vivo.</i>	1
6.4	Iontophoresis.	1
6.5	Recent advances –use of microneedles in transdermal drug delivery.	1
7	Introduction to Pharmaceutical Processing Development. (As per ICH guidelines)	4
7.1	Elements in Pharmaceutical development <ul style="list-style-type: none"> • Target product profile • Critical Quality Attributes. • Linking Material Attributes & process parameters to CQA's Risk Assessment • Design space • Control Strategy • Product Lifecycle management & continual improvement. 	3
7.2	Submission of Pharmaceutical Development and related information in CTD format. <i>Relevant Examples.</i>	1
Total		60

Books:

1. Targeted and Controlled Drug Delivery: Novel Carrier Systems by Vyas SP, Khar RK, CBS Publishers and Distributors, 1stedn, 2002.
2. Controlled and Novel Drug Delivery by Jain NK, CBS Publishers and Distributors, 2008.
3. Controlled Drug Delivery: Fundamentals and Applications by Robinson JR, Lee VHL, Dekker, 2ndedn, Vol 29, 1987.

4. Novel Drug Delivery System by Chien YW, 2nd edn, Vol 50, Informa Healthcare, 2003.
5. Progress in Controlled and Novel Drug Delivery Systems by Jain NK, CBS Publishers and Distributors; 2004.
6. Ophthalmic Drug Delivery Systems, Mitra AK, 2nd edn., Drugs and Pharmaceutical Sciences Series, Vol. 130, Marcel Dekker, 2003.
7. Polymeric drug delivery system, Kwon GS, Marcel Dekker, Vol 148, 2005.
8. Nanoparticulate Drug Delivery System by Thassu D, Deleers M, Pathak Y, Marcel Dekker, Vol 166, 2007.
9. Controlled Drug Delivery- Challenges and Strategies by Park K, American Chemical Society, 1997.
10. Colloidal Drug Delivery System by Kreuter J, Marcel Dekker Vol 66, 1994.
12. www.ich.org
13. Pharmaceutical Dosage Forms: Disperse Systems by Lieberman HA, Rieger MM, Banker GS, Marcel Dekker, Vol 3, 2nd edn, 2005.
14. Pharmaceutical Emulsions and Suspensions by Nielloud F, Marti- Mestres G, Marcel Dekker, Vol 105, 2000.
14. Controlled Release Systems Fabrication Technology by Dean STH, CRC Press, Vol 1, 1988.
15. Bioadhesive Drug Delivery Systems by Mathiowitz.E , Chickering DE , Lehr CM, Marcel Dekker Vol 98 ,1999.
16. Pharmaceutical Skin Penetration Enhancement by Walters. K A, Hadgraft J, Marcel Dekker, Vol 59, 1993.
17. Percutaneous Absorption by Bronaugh RL, Maibach HI, Taylor and Francis, 3rd edn, Vol 97, 2005.
18. Transdermal Controlled Systemic Medication by Chien YW, Marcel Dekker, Vol. 31, 1987.
19. Oral Mucosal Drug Delivery by Rathbone MJ, Marcel Dekker, Vol 74, 1996.
20. Modified Release Drug Delivery Technology by Rathbone MJ, Hadgraft J, Roberts MS, Lane ME, Informa Healthcare, 2nd edn, Vol 183(1), 2008.
21. Pharmaceutical Pelletization Technology, Ghebre-sellassie. J, , Marcel Dekker, Vol. 37

Core 2

Advanced Pharmaceutics II

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Targeted systems:-Active and Passive approaches:	6
1.1	Tumour targeting, Molecular targets for cellular targeting, Ligands as delivery and targeting tools, Concept of receptor mediated endocytosis.	3
1.2	<i>Self Study: Concepts and rationale of targeting: active and passive targeting, Cellular biochemistry and molecular events in drug targeting</i>	3
2	Pulmonary and nasal drug delivery systems:	14
2.1	Nasal drug delivery: Nasal administration – dosage forms, Strategies for enhancement in nasal absorption, Animal	4

	models for nasal absorption studies, Nasal preparations for systemic effect	
2.2	Pulmonary drug delivery : Factors affecting particle disposition in the lungs, Dosage forms for pulmonary drug delivery (Nebulizer, Metered dose inhalers, Dry powder inhalers), Drug targeting to the respiratory tract, Pulmonary receptor targeting	7.
2.3	<i>Self Study: Anatomy and physiology of the respiratory system, Airway physiology and disposition patterns</i>	3
3	Protein and peptide drug delivery systems:	11
3.1	Physical and chemical stability aspects, protein degradation pathways, techniques of stabilization of proteins and peptides, barriers to transport and approaches to circumvent metabolic barriers.	4
3.2	General protein formulation and delivery system strategies.	1
3.3	Routes for delivery of proteins and peptides with emphasis on oral and mucosal delivery, pulmonary delivery, nasal delivery and parenteral delivery	3
3.4	<i>Self Study: Structure of proteins and peptides, analysis of proteins and peptides.</i>	3
4	Colloidal drug delivery systems:	22
	NOTE that for every colloidal drug delivery system the following aspects to be included: Introduction, comparison with other colloidal drug carriers, Advantages/limitations, constituents and mechanism of formation, method of preparation and drug loading, characterisation and evaluation, stability, long circulating / modified form of colloidal drug carrier, bio distribution and application. The following drug delivery system to be studied with respect these aspects –	
4.1	Liposomes	5
4.2	Niosomes	2
4.3	Nanoparticles	5
4.4	Polymeric micelles	3
4.5	Solid Lipid nanoparticles.	4
4.6	<i>Self Study: An overview colloidal Drug Delivery with respect to Physicochemical & Biopharmaceutical aspects</i>	3
5	Brain targeting:	7
5.1	Introduction, Transport through BBB, Factors affecting drug permeation through BBB	1
5.2	Brain drug delivery strategies: <ul style="list-style-type: none"> • Invasive- Intracerebral implants, Intraventricular infusion, BBB disruption, • Non-invasive techniques- Chemical method, Colloidal drug carrier, receptor/ vector mediated approach. • Miscellaneous techniques- Intranasal etc 	3
5.3	<i>Self Study: Blood brain barrier, CSF barrier, limitations in brain uptake of drug, desired</i>	3

	<i>physicochemical characteristics of drugs.</i>	
	Total	60

Books:

1. Targeted and controlled drug delivery: Novel carrier systems- by S. P Vyas and R. K Khar, CBS publishers and distributors pvt Ltd.
2. Advances in controlled and novel drug delivery edited by N.K.Jain, CBS publishers and distributors pvt Ltd.
3. Robinson J.R and Lee- controlled and novel drug delivery.
4. Controlled drug delivery: Concepts and advances, S.P.Vyas, R.K. Khar, Vallabh Prakashan.
5. Chien- Y. W- Novel Drug Delivery System, Drug and pharmaceutical science series, Vol 14, New York Inc, Marcell Dekker.
6. Controlled and novel drug delivery edited by N. K .Jain, CBS publishers and distributors pvt Ltd.
7. Advances in pharmaceutical sciences – vol-1 to 5, by H. S. Bean and A. H Beckett.
8. Glen S. Kwon, Polymeric drug delivery system- Marcell Dekker Series , Vol 148, P.No.533-560
9. Thassu D “Nanoparticulate Drug Delivery System” Vol 166, Marcell Dekker Series 2007.
10. Park K, Control Drug Delivery- Challenges and Strategies, CRC, Washington DC 1997
11. MacNally E “Protein Formulation and Delivery” 2nd edition, Vol75, 2008
12. Kreuter J, Colloidal Drug Delivery System, Vol.66, Marcell Dekker, Inc New York,1994

Semester-II

BRANCH: PHARMACEUTICAL ANALYSIS

Core 1

Analytical Method Development and Validation Techniques

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Calibration & Validation of analytical instruments: a. HPLC. b. UV-VIS spectrophotometer. c. FTIR. d. Dissolution test apparatus.	4
2	HPLC assay method development for API and drug products:	20
2.1	Preliminary investigations- Nature of sample, its composition and properties. (It should also include significance of pK _a , partition coefficient and current methods to determine the same), separation goals, sample pretreatment and detection, developing separation.	5

2.2	Basics of separation-Resolution, Resolution as a function of- solvent strength, selectivity and column plate number; and sample size effect.	1
2.3	<i>Self study-Detection-Comparison of sensitivity, selectivity, advantages, disadvantages and applications with respect to detectors such as U.V, Fluorescence, PDA, Refractive Index, Evaporative light scattering detector and electrochemical detectors.</i>	2
2.4	Sample preparation and pretreatment for solid, liquid, semisolid samples; column switching and pre and post column derivatisation.	2
2.5	<i>Self study-Columns-characteristics of column and column packing and column specifications.</i>	1
2.6	Method development for Reverse-phase, Ion pair and ion exchange chromatography, Gradient elution-principle and development of gradient separation. <i>Self study- pharmaceutical examples for these methods-(1 hr).</i>	3
2.7	Quantitation analysis- measurement of signals, quantitation methods, sources of errors, procurement, storage and use of reference standards and working standards.	2
2.8	ICH guidelines for analytical method validation (Q2 with latest revision) .System suitability testing as per USP, IP.	3
2.9	<i>Self study-One detailed HPLC analysis of any API by USP or IP (1hr)</i>	1
3	HPTLC : Method development and validation for fixed dose combination drugs and herbal analysis.	3
4	Impurity profiling:	9
4.1	1. <i>Self study-Sources of impurities and ICH terminologies-Organic impurities, Inorganic impurities, Residual solvents, Isolation and characterisation methods for impurities (3 hrs).</i> 2. Analytical method development and quantitation of impurities.	5
4.2	ICH guidelines-Q3A, Q3B, Q3C with latest revisions.	4
5	Bioanalytical method development and validation:	13
5.1	Steps followed-sample preparation, liquid-liquid extraction, precipitation, solid-phase extraction, sintered column solid phase extraction.	3
5.2	Bioanalytical method validation including full, partial, cross validation, selectivity, accuracy, calibration curve, stability (freeze-thaw and mobile phase), recovery.	4
5.3	CDER and ICH guidelines for bioanalytical method validation.	4
5.4	<i>Self study- Examples of bioanalytical method development and validation for a specified drug estimated in urine/ plasma/serum samples.</i>	2
6	Stability testing:	11
6.1	Drug development cycle and stability testing.	2
6.2	Stress testing of drug substances.	1
6.3	Stability indicating assays (specific and selective), Role of kinetic studies.	3
6.4	Stability testing protocols.	1

6.5	Retest period / Shelf-life determination of drug substances / phytopharmaceuticals / biotechnological products and equipments.	2
6.6	ICH guidelines-Q1A and Q1B with latest revisions.	2
	Total	60

Books:

1. Practical HPLC Method Development by L.R.Snyder, 2nd edition, John Wiley & Sons,
2. Analytical Method Validation and Instrument Performance Verification by Chung Chow Chan, Herman Lam, Y.C. Lee, Xue-Ming Zhang, Wiley Interscience, John Wiley & Sons, Incorp Publications.
3. United States Pharmacopoeia and Indian Pharmacopoeia.
4. Handbook of Isolation and Characterisation of Impurities in Pharmaceuticals by Satinder Ahuja & Karen Mills Alsante, Volume 5, Academic Press, USA.
5. Handbook of Bioanalysis & Drug metabolism by Gary Evans, CRC Press (2004),
6. HPLC method development by Satinder Ahuja
7. Sethi's Quantitative Analysis of Pharmaceutical Formulations by P.D.Sethi, fourth edition, volume-1, CBS Publishers and Distributors, New Delhi.
8. Remington-The Science and Practice of Pharmacy, 20th edition Remington-The Science and Practice of Pharmacy, 20th edition.
9. Validation and Qualification in Analytical Laboratories, 2nd edition, Ludwig Huber; Informa Healthcare.
10. Handbook of stability testing in pharmaceutical development - Regulations, Methodologies and Best practices; Editor Kim Huynh-Ba, Springer.
11. J.T. Carstensen, C.T. Rhodes, "Drug stability: principles & Practices". Latest Edition. Marcel Dekker Inc., New York

INTERNET REFERENCES:

1. US FDA (CDER) and ICH guidelines for Bioanalytical method validation.
2. ICH guidelines- Q1A(R), Q3A(R), Q3B, Q3C, Q6A.
3. ICH guidelines for analytical method validation.

CORE 2

Spectroscopic Structural Elucidation

4 hrs / week

Problems of structural elucidation involving the following techniques:		
<ul style="list-style-type: none"> • UV spectroscopy. • IR spectroscopy. • ¹H-NMR spectroscopy. • ¹³C-NMR spectroscopy. • Mass Spectrometry. 		
The problems should cover the following aspects:		
Unit	Course Contents (Topics)	Hours
1	Calculation of λ max for dienes, α,β –unsaturated ketones by UV spectroscopy. <i>Self study- practice problems (1 hr)</i>	5
2	Prediction of characteristic IR bands, NMR spectra (¹ H NMR) –chemical shift, splitting pattern and ratio of proton intensity, (¹³ C NMR)-number of signals, chemical shift and splitting pattern, mass fragmentation patterns. <i>Self study-practice problems (2 hrs)</i>	10
3	Distinguishing compounds using UV / IR / ¹ H NMR/ ¹³ C NMR and /or Mass spectrometry. <i>Self study-practice problems (2 hrs)</i>	10

4	Interpretation of mass spectra with explanation of fragmentation patterns. <i>Self study-practice problems (2 hrs)</i>	9
5	Problems involving structure elucidation by- UV / IR / ¹ H NMR / ¹³ C NMR and / or Mass spectrometry, <i>Self study-practice problems (8 hrs)</i>	26
	Total	60

Books:

- 1 Introduction to Spectroscopy by D.L.Pavia, G.M.Lampman & G.S.Kriz, Latest edition, Thomson Brooks/Cole, United States.
- 2 Spectrometric Identification of Organic compounds by Robert.M.Silverstein & Francis.X.Webster, D.J.Kiemle, Latest edition, John Wiley & Sons.
- 3 Organic Spectroscopy by William Kemp.
- 4 Applications of absorption spectroscopy of organic compounds by John Robert Dyer, Prentice Hall, London.

Semester-II

BRANCH: QUALITY ASSURANCE

Core 1

Quality Assurance Systems

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Regulatory basis for validation: US FDA guidelines (cGMP guidelines, 21 CFR 210-211), EU guidelines, WHO guidelines.	5
2	Terminology and validation overview:	10
2.1	<i>Self Study: Validation versus verification, testing, calibration and qualification.</i>	3
2.2	Concepts of DQ, IQ, OQ and PQ.	3
2.3	Concepts of Prospective validation, retrospective validation, Concurrent and revalidation. Validation Master Plan.	4
3	Validation of Equipment	10
3.1	Dry Powder Mixers	1
3.2	Fluid Bed and Tray dryers	1
3.3	Tablet Compression Machine	2
3.4	<i>Self study :Dry Heat Sterilization/Tunnels</i>	1
3.5	Autoclaves	2
3.6	Capsule filling machines.	1
3.7	Validation of Integrated lines by media fill test.	2
4	Utilities Validation	7
4.1	Validation of Pharmaceutical Water System & pure steam,	2
4.2	Validation of HAVC system	3
4.3	Validation of Compressed air	2
5	Cleaning Validation <i>Self study: Cleaning of Equipment, Cleaning of Facilities.</i>	4
6	Analytical Method Validation: General principles of analytical method validation, Validation of following analytical Instruments	06
6.1	HPLC	2
6.2	Dissolution test apparatus	2
6.3	U.V./Visible spectrophotometers	2

7	Process Validation	13
7.1	<i>Self study : Prospective, concurrent, retrospective & revalidation Self-study</i>	1
	Process validation of following formulations	
7.2	Uncoated / Coated tablets	2
7.3	Hard gelatin Capsules	2
7.4	Ampoules & Vials	2
7.5	<i>Self study: Ointment/Creams</i>	2
7.6	<i>Self study : Liquid Orals</i>	2
7.7	Transdermal patches (Matrix systems)	2
8	<i>Self study-Computer system validation in controlling the manufacturing process.</i>	2
9	Process Analytical Technologies (PAT) and Quality by Design (QbD) (US FDA)	3
	Total	60

Books:

1. Validation and Qualification in Analytical Laboratories by Ludwig Huber, Second edition (2007), Informa Health Care, New York, London.
2. Pharmaceutical Process Validation by R.Nash and Wachter, Volume 129, Latest edition, Marcel Dekker Inc, New York.
3. GMP for Pharmaceuticals by Sidney H. Willing, Fifth edition, Marcel Decker Series, New York.
4. United States Pharmacopoeia & Indian Pharmacopoeia.
5. Validation of Pharmaceutical process, F. J. Carleton and J. Agalloco, Marcel Dekker Inc.

INTERNET REFERENCES:

1. www.fda.gov (US FDA guidelines for PAT and QbD).
2. www.ich.org
3. WHO publications on related topics.
4. EMEA guidelines

Core 2

Pharmaceutical Quality Management

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Concept of-	8
1.1	Total Quality Management (TQM),	2
1.2	Quality control and quality assurance,	2
1.3	Quality control laboratory responsibilities,	2
1.4	<i>Self study: Good laboratory practices</i>	2
2	GMP	8
2.1	Organization of pharmaceutical manufacturing unit, production management,	4
2.2	<i>Self study : Revised schedule M.</i>	4
3	Personnel:	12

Unit	Course Contents (Topics)	Hours
3.1	<i>Self study: Introduction, Human resource development, Qualification Experience and Training, Responsibilities, Personal Hygiene and Gowning.</i>	6
3.2	Location, Plant layout, Lighting, Sewage, Water handling-Sewage, Refuse and Disposal, Washing and toilet facility, Sanitation, Controls of contamination and Environmental controls.	6
4	Materials Management:	8
4.1	API's, raw materials & packaging materials, Purchase specifications, Selection of vendors, Intermediates & Finished products, Rejected and Recovered materials, Recalled products, Reagents & culture media, Reference standards, Waste materials.	6
4.2	Warehousing- Good Warehousing Practices, distribution and records.	2
5	Manufacturing Operations and Control:	8
5.1	<i>Self study: Sanitation of Manufacturing Premises, Line clearance, Mix-ups and Cross contamination, Processing and holding of Intermediates and Bulk Products</i>	3
5.2	Packaging, I.P.Q.C., Release and storage of Finished Product, Process Deviations and Incidents, Drug product inspection, Yield calculations	3
5.3	Expiry dating, Manufacturing record review and approval.	2
6	Documentation and Records: In-process and Product Release Specifications, Master production and control record, Batch production and control record, Standard Operating Procedures (SOP), Change Control, Site master file.	6
7	Post Operational Activities:	5
7.1	Distribution, Complaints and recalls, evaluation of complaints, Recall procedures, related records and documents.	2
7.2	Outsourcing: Facility audit, Manufacturing, Packaging, Analytical, Clinical and other services outsourcing.	3
8	Site and Plant security: Security personnel, Entry procedures to site & plant, Internal security, Vehicle parking, Fuel storage, Canteen & cooking, Garden & horticulture.	2
9	Audits: Principle of Quality audit, Plant level, Department wise documentation.	3
	Total	60

Books

1. Quality Assurance of Pharmaceuticals, Vol. 2, Updated Edition, World Health Organization, Geneva.
2. S.H. Willing, Good Manufacturing Practices for Pharmaceuticals; A plan for total Quality control, Latest Edition, Marcel Dekker.
3. Regulatory guidelines related to GMP by
 - a. 21 Code of Federal Regulation, Parts 210, 211&58 (USFDA guidelines)
 - b. EU, MHRA, UK Guidelines on GMP
 - c. Schedule M of Drug & Cosmetics Act.
4. Quality Planning & Analysis by J. M. Juran and F. M. Gryna, Tata Mcgraw Hill, India.
5. Quality Assurance Guide by Organization of Pharmaceutical Producers of India.

Semester-II

BRANCH: PHARMACOGNOSY AND PHYTOCHEMISTRY

Unit	Course Contents (Topics)	Hours
1	Factors affecting occurrence of compounds of natural origin	6
1.1	Discussion of different factors contributing to the variation in the composition and proportion of secondary metabolites.	2
1.2	Concept of variation of phytochemicals with respect to ecotype, phenotype and genotypic variables. Study of phytoalexins, allelochemicals and aflatoxins, natural Pesticides(cover the topics with at least two examples of each).	4
1.3	<i>Recent advances and applications of phytoalexins, allelochemicals& aflatoxins.</i>	3
2	Chemistry , sources & uses of following classes of phytochemicals (1) Alkaloids – Opioids & Purines (2) Iridoids (3) Coumarins (4) Xanthones	8
2.1	<i>Self study- Update on traditional uses and recent applications of few examples of the above mentioned classes</i>	5
3	Chemistry, classification, sources, uses and structure- elucidation by spectral methods of Flavanoids.	8
3.1	<i>Self study – Comparative spectral analysis & recent application of different classes of flavanoids.</i>	4
4	Study of following therapeutic classes of agents of plant and animal origin with respect to sources, applications and chemistry (any two examples of each class). (1) Anti-bacterial (2) Hepatoprotective & Hypolipidemic agent (3) Anti-virals	6
5	Marine drugs of different therapeutic classes 1) Anti-cancer 2) Cardiovascular drugs 3) Anti-virals 4) Anthelmintics 5) Marine toxins	6
6	Study of photosensitises of natural origin such as porphyrins, psoralenes, thiophenes, quinines and their significance in Photodynamic therapy (PDT) and phtotoxicity.	6
6.1	<i>Self study-Role of PDT in health care with examples.</i>	3
7	Introduction to plant tissue culture (PTC) and plant biotechnology.	5
7.1	Genetic engineering in plants for development of plants resistant to pests, viruses, microbes and diseases. Alteration in ripening of fruits. Advantages and disadvantage of BT crops.	3
7.2	Definition, Methodology, & application of biotransformation of Phytochemicals with suitable examples.	2
	Total	60 hrs

Books

1. Pharmacognosy Phytochemistry – Medicinal Plants- Jean Brunetton, Lavoisier Publishing, Paris.
2. Text book of Pharmacognosy- Trease and Evans- 14th edition. Elsevier science
3. Transgenic Plants- R. Ranjan- Agro Botanica, New Delhi.
4. Transgenic Plants-A Production system for Industrial and Pharmaceutical Proteins. by Meran Owen, Jan Pen- John Wiley.
5. Medicinal Plant-Their Bioactivity, Screening and Evaluation- CSIR.
6. Homeopathic Pharmacopoeia of India- Publisher Ministry of Health.
7. The Ayurvedic Formulary of Part I & II- Publisher Ministry of Health.
8. Chinese Materia Medica-You-PingZhu- Harwood Academic Publishers.
9. India Materia Medica- Nadkarni A.K. –Bombay Popular Prakashan.
10. Phytochemical Methods - J.B.Harbone - Chapman and hall
11. Cultivation's and Processing of Medicinal Plants-Ed. by L. Hornok-John Wiley.
12. Introduction to Flavanoids-Bohrn Bruce A. – Herwood Academic Publishers.
13. Cultivation and Utilization of Aromatic plants – Ed. By Atal C. K. and Kapur B.M.- CSIR.
14. Plant Tissue and Cell Culture Ed. H.E. Street – Blackwell Scientific publications.
15. Aflatoxin- Leo A. Gold Blatt- Academic Press New York.
16. Microbial Toxins- Ciejler, Kadis and Ajl- Academic press.
17. Antimicrobial in Food – Alfred larry Branen, P. Michael Davidson Publishing house
18. Chemical plant Taxonomy T. Swain, 1963. Academic Press, London.
19. Plant Taxonomy and Biosystematics .C.A Stace, 1985. Edward Arnold, London.
20. Modern methods of plant analysis K. Paech, 1956., Springer-Verlag.
21. Indian Herbal Pharmacopoeia, Vol. 1&2, RRL, IDMA, 1998, 2000.
22. Indian Pharmacopoeia, 2010.
23. Standardization of Botanicals,V. Rajpal, 2002. Eastern Publishers, New Delhi.
24. Natural Compounds as Drugs – Vols. I & II, Editor- Frank Petersen, René Amstutz, Die Deutsche Bibliothek, Germany.
25. Quality control of Herbal Drugs: An Approach to evaluation of Botanicals, Pulok Mukherjee - Riddhi International
26. Chemicals from Plants: Perspectives on Plant Secondary Product, Walton & Braun, Imperial College Press.
27. Towards Natural Medicine Research in the 21st Century H. Ageta, N. Aimi et al Excerpta Medica, International Congress Series 1157.

Core 2

Natural Product Technology

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Detailed study of WHO guidelines for quality control of crude drugs	5
2	Study of herbal formulations	12
2.1	Classification and different stages in preparation of herbal formulations for therapeutic and cosmetic applications	5
2.2	Standardization and evaluation of quality control and safety of herbal formulations	5
2.3	Introduction to different Ayurvedic dosage forms	2
2.4	<i>Self study- Any two marketed herbal formulations with respect to constituents and uses.</i>	4
3	Quantitative assays to determine extraction efficiency	8
3.1	General methods of estimation of alkaloids, terpenoids and flavonoids	4
3.2	Analysis of Rutin, lycopene, curcuminoids, artemisinin, enzymes and lectins by different method such as UV, HPTLC, GC, gel electrophoresis etc., determination of percentage purity.	4

3.3	<i>Self study- Merits & demerits of different methods of estimation of alkaloids, terpenoids & flavanoids.</i>	5
4	Introduction to herbal product based industry	6
4.1	Types, forms, scope and application of herbal industries	4
4.2	Type of infrastructure involved in making standardized extract and different dosages forms	2
4.3	<i>Self study – Equipment and machinery used in large scale extraction</i>	3
5	Bioavailability and pharmacokinetic aspect of herbal drugs with examples of well known documented herbal drugs. Introduction to concept of phytoequivalence and pharmaceutical equivalence.	6
6	IPR issues related to herbal and natural products. EMEA and ESCOP guidelines for herbal medicinal products. Preparation of DMF for herbal medicines.	8
6.1	<i>Self study – Any one patent related to natural products.</i>	3
	Total	60 hrs

Books

1. Pharmacognosy Phytochemistry – Medicinal Plants- Jean Brunetton, Lavoisier Publishing, Paris.
2. Text book of Pharmacognosy- Trease and Evans- 14th edition. Elsevier science
3. Transgenic Plants- R. Ranjan- Agro Botanica, New Delhi.
4. Transgenic Plants-A Production system for Industrial and Pharmaceutical Proteins. by Meran Owen, Jan Pen- John Wiley.
5. Medicinal Plant-Their Bioactivity, Screening and Evaluation- CSIR.
6. Homeopathic Pharmacopoeia of India- Publisher Ministry of Health.
7. The Ayurvedic Formulary of Part I & II- Publisher Ministry of Health.
8. Chinese Materia Medica-You-PingZhu- Harwood Academic Publishers.
9. India Materia Medica- Nadkarni A.K. –Bombay Popular Prakashan.
10. Phytochemical Methods - J.B.Harbone - Chapman and hall
11. Cultivation's and Processing of Medicinal Plants-Ed. by L. Hornok-John Wiley.
12. Introduction to Flavanoids-Bohrn Bruce A. – Herwood Academic Publishers.
13. Cultivation and Utilization of Aromatic plants – Ed. By Atal C. K. and Kapur B.M.- CSIR.
14. Plant Tissue and Cell Culture Ed. H.E. Street – Blackwell Scientific publications.
15. Aflatoxin- Leo A. Gold Blatt- Academic Press New York.
16. Microbial Toxins- Ciejler, Kadis and Ajl- Academic press.
17. Antimicrobial in Food – Alfred larry Branen, P. Michael Davidson Publishing house
18. Chemical plant Taxonomy T. Swain, 1963. Academic Press, London.
19. Plant Taxonomy and Biosystematics .C.A Stace, 1985. Edward Arnold, London.
20. Modern methods of plant analysis K. Paech, 1956., Springer-Verlag.
21. Indian Herbal Pharmacopoeia, Vol. 1&2, RRL, IDMA, 1998, 2000.
22. Indian Pharmacopoeia, 2010.
23. Standardization of Botanicals,V. Rajpal, 2002. Eastern Publishers, New Delhi.
24. Natural Compounds as Drugs – Vols. I & II, Editor- Frank Petersen, René Amstutz, Die Deutsche Bibliothek, Germany.
25. Quality control of Herbal Drugs: An Approach to evaluation of Botanicals, Pulok Mukherjee - Riddhi International
26. Chemicals from Plants: Perspectives on Plant Secondary Product, Walton & Braun, Imperial College Press.
27. Towards Natural Medicine Research in the 21st Century H. Ageta, N. Aimi et al Excerpta Medica, International Congress Series 1157.

BRANCH: PHARMACOLOGY

Core 1

Advanced Pharmacology

4 hrs/week

Unit	Course Contents (Topics)	Hours.
1	General aspects of drug discovery and development	2
2	High Throughput Screening	6
2.1	Techniques for High throughput screening. <ul style="list-style-type: none">a. Cell based assaysb. Biochemical assays.c. Radio ligand binding assays.	4
2.2	<i>Self study-Importance of pharmacokinetic studies in drug development.</i>	2
3	Toxicity Studies	8
3.1	<ul style="list-style-type: none">• Acute, subacute and chronic toxicity• Safety pharmacology evaluation.• Genetic toxicity, cytotoxicity, toxicogenomics	5
3.2	<i>Self study-Schedule Y, OECD, ICH guidelines for toxicity studies by various routes.</i>	3
4	Introduction to Chronopharmacology	8
4.1	<ul style="list-style-type: none">• Circadian rhythm, Biological Clock, Location, Neuroanatomy and Neurochemistry.• Rhythms and pharmacokinetics.	5
4.2	<i>Self study- Rhythms and therapeutics of diseases of GIT and asthma.</i>	3
5	Stem Cells and Therapeutic applications	3
6	Novel drug targets	18
6.1	Physiological functions, pharmacological implications and therapeutic potential of the following target sites: <ul style="list-style-type: none">• Rho kinase (ROCK).• Phospho inositide-3-Kinase. (PI3K).• Akt (Protein kinase B).• Caspases.• Proteases.• Poly (ADP ribose) Polymerase (PARP).• Peroxisome proliferator activator receptors (PPAR α,γ).	15
6.2	<i>Self study-Biological functions of Nitric oxide (NO) and therapeutic potential of nitric oxide modulators.</i>	3
7	Transporter proteins	7
7.1	<ul style="list-style-type: none">• Classification and biology of ATP binding cassette (ABC) transporter super family, Solute carrier transporter (SLC).• Multi drug resistance proteins (MDR).• Cystic fibrosis transmembrane regulator (CFTR).	5

7.2	ABC transporters involved in drug absorption distribution and excretion.	2
8	Role of Cytokines, Prostaglandins, TNF- α, Bradykinin, Leukotrienes, PAF, Interferons and Adhesion molecules in various immunological and inflammatory disorders	8
	Total	60

Books:

1. Rang and Dale's pharmacology-- Elsevier Churchill Livingstone.
2. Lange's Basic and clinical pharmacology, Katzung B. G. Masters S. B., Trevor A. G. Tata McGraw Hill.
3. Goodman and Gilman's pharmacological basis of therapeutics, Edited by Laurence Brunton, Bruce Chabner and Bjorn Knollman, McGraw Hill.
4. Pharmacological reviews, Annual reviews Inc.
5. Advances in pharmacology, Academic Press.
6. Trends in Pharmacological Sciences, Cell Press Elsevier Publication.

Core 2

Clinical Research Methodology

4hrs/week

Unit	Course Contents (Topics)	Hours.
1	Clinical trials	14
1.1	Introduction, drug discovery and drug development.	10
1.2	Various phases of clinical trials.	
1.3	Study methodology/designs, Inclusion and Exclusion criteria, objectives and endpoints (efficacy and safety), Methods of allocation, blinding and randomization.	
1.4	Informed consent process.	
1.5	Study monitoring and its importance.	
1.6	Safety monitoring in clinical trials.	
1.7	<i>Self study- BA/BE studies, Post marketing studies.</i>	
2	Documents in clinical study Essential documents in clinical trial-Investigators Brochure (IB), Protocol and amendments in protocol, Case report form (CRF), Informed consent form (ICF), Content of clinical study report(CSR).	4
3	Ethical guidelines in clinical research	9
3.1	History, ICH-GCP and its principles, Indian GCP (CDSCO Guidelines), ICMR guidelines- Ethical guidelines for Biomedical Research on human subjects 2006, Schedule Y 2005, USFDA guidelines for IND, NDA, ANDA applications.	6
3.2	Self study: EMEA organization and its functions, EU regulatory guidelines.	3
4	Roles and responsibility of various clinical trial personnel as per ICH-GCP Sponsor, Investigator, Monitor, Auditors	3

5		5
5.1	Institutional Ethics Committee (IEC)/ Independent Ethics Committee (IdEC)/Institutional Review Board (IRB)	2
5.2	<i>Self study-Ethical theories, Integrity and Misconduct in clinical research.</i>	3
6	Role of Quality assurance in clinical research	2
7	Clinical Data Management and Report Writing	3
8	Pharmacoepidemiology Types, methods, and factors affecting drug utilization, applications of pharmacoepidemiology in health care and rational use of drugs	5
9	Pharmacovigilance	10
9.1	Definition, scope and aims of pharmacovigilance, Adverse drug reactions-Classification, mechanism, predisposing factors and causality assessment, Role of clinical pharmacist in reporting, evaluation, monitoring, prevention and management of ADRs.	5
9.2	<i>Self study: Reporting-CIOMS forms, Periodic safety update reports (PSURs) as per Indian regulatory guidelines.</i>	5
10	Pharmacoeconomics and Outcomes Research Theories and methodologies of Pharmacoeconomics and Outcomes Research. Applications of pharmacoeconomics to pharmacotherapy and managed health care.	5
	Total	60

Books

1. Rick NG. Drugs From Discovery to Approval, second edition, John Wiley & Sons, Inc 2004.
2. Allen Cato, Lynda Sutton. Clinical Drug Trials and Tribulations Second Edition Revised, second edition, Marcel Dekker Inc; 2nd Revised edition March 26, 2002.
3. Ronald D. Mann, Elizabeth B. Andrews. Pharmacovigilance, second edition, John Wiley & Sons Ltd,2002.
4. Shayne C. Gad. Drug Safety Evaluation. A John Wiley & Sons, Inc. Publication, 2000.
5. Sandy Weinberg. Guidebook For Drug Regulatory Submissions, first edition, A John Wiley & Sons, inc.2009.
6. Duolao Wang and Ameet Bakhai Clinical Trials A Practical Guide to Design, Analysis, and Reporting, Remedica 2006.
7. Giovanna di Ignazio, Di Giovanna and Haynes, Principals of Clinical Research, Wrightson Biomedical Pub., 2001
8. R K Rondels, S A Varley, C F Webb, Clinical Datamanagement, Second Edition, John Wiley & Sons Inc, January 1997.
9. Strom BI, Limmel SE. Textbook of Pharmacoepidemiology. Chichester, West Sussex, England: John Wiley & Sons Ltd; 2006.
10. Rascati, Karen L. Essentials Of Pharmacoeconomics. Philadelphia, Pa.: Lippincott Williams & Wilkins, 2009.
11. M. F. Drummond, M. J. Sculpher and G. W. Torrance, Methods for the economic evaluation of health care programmes. Oxford University Press, USA, 2005.

12. Brenda Waning; Michael Montagne; William W McCloskey, Pharmacoepidemiology : principles and practice, New York : McGraw-Hill, 2001

13. Various Guidelines like:

- ICH (International Conference on Harmonisation), GCP for registration of pharmaceuticals for human use. ICH Harmonised Tripartite
- Guideline for Good Clinical Practice, E6, 1996.
- ICMR Guideline – Ethical Guidelines for Biomedical Research on Human Subjects.
- Indian GCP – Central Drugs Standard Control Organization. Good Clinical Practices
- Guidelines for Clinical Trials on Pharmaceutical Products in India. New Delhi: Ministry of Health; 2001.
- Pharmacovigilance Programme of India (PvPI)

SEMESTER II

BRANCH: BIOPHARMACEUTICS AND PHARMACOKINETICS

Core 1

Biopharmaceutics

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Mechanisms of drug release	10
1.1	Diffusion controlled release, chemically controlled release, swelling controlled release of drugs from formulations- Higuchi model and the Power-Law model for drug release and their comparison, discussion of newer mechanistic models described drug release from formulations	6
1.2	<i>Self study of zero, first, and second order release kinetics and their graphical profiles</i>	4
2	Drug Dissolution	18
2.1	Theories of drug dissolution – Noyes Whitney Diffusion model; Hixon Crowell Model; Interfacial barrier model (Continuous and discrete reaction limited dissolution), concepts of solubility versus dissolution rate, physicochemical factors affecting drug dissolution, pharmaceutical factors affecting drug dissolution, physiological factors affecting drug dissolution, , methods for estimation of solubility, methods for determination of dissolution rate	14
2.2	<i>Self study of experimental method design for solubility and dissolution rate determination</i>	4
3	Drug Absorption	16
3.1	Mechanisms of drug absorption, detailed discussion of the variety of transporters and the role of transporters in the GI tract and liver and their role in drug absorption, physicochemical factors affecting drug absorption, pharmaceutical factors affecting drug absorption, physiological factors affecting drug absorption, gut and hepatic metabolism and their role in determination of bioavailability, invitro and invivo methods for estimation of permeability/transport across membranes/absorption, computational methods for prediction of solubility/permeability/absorption	12
3.2	<i>Self study of the experimental design of methods for determination/prediction of drug transport</i>	4
4	Routes of Drug Administration	11

4.1	Discussion of the different routes of drug administration for the perspective of the nature of the absorption barrier/s, mechanisms of drug release, drug permeability/absorption from the site of administration, drug/pharmaceutical/physiological factors affecting drug dissolution/dissolution rate/absorption from the different routes of drug delivery	8
4.2	<i>Self study of the advantages and limitation of the different routes of administration and examples of drug administered by these routes</i>	3
5	Discussion of the traditional and high-throughput approaches towards estimation of solubility, dissolution rate and drug absorption and use of this information in a drug discovery and development setting.	5
	Total	60

Books:

1. Clinical Pharmacokinetics and Pharmacodynamics-Concepts and Applications, Rowland M and Tozer TN, Walters Kluwer – Lippincott Williams and Wilkins.
2. Applied Biopharmaceutics and Pharmacokinetics, Shargel L and Yu ABC, Appleton and Lange, International Edition
3. Handbook of Basic Pharmacokinetics including clinical applications, Ritschel WA and Kearns GL, APhA,
4. Basic Pharmacokinetics, Jambhekar SS and Breen PJ, Pharmaceutical Press.
5. Biopharmaceutics and Pharmacokinetics, Venkateshwarlu V, Pharma Book Syndicate
6. Drug Bioavailability- Estimation of solubility, permeability, absorption and bioavailability, van der Waterbeemd H, Lennernas H and Artursson P, Wiley VCH.
7. Modelling in Biopharmaceutics, Pharmacokinetics and Pharmacodynamics – Homogenous and Heterogenous approaches, Macheras P and Iliadis A, Springer

Core 2

Pharmacokinetics

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Introduction to pharmacokinetics and its utility in drug design and dosage regimen design. Definitions of absorption, distribution, metabolism, excretion, elimination. Different approaches for determination of pharmacokinetics of drugs – non-compartmental, physiological, and compartmental modeling. Assumptions involved in the evolution of single and multi-compartment models.	4
2	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following different dosing methods/protocols [blood/plasma/urine sampling]	40
2.1	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following intravenous bolus dosing [blood/plasma/urine sampling]	5
2.1	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following intravenous multiple bolus dosing	5

	[blood/plasma]	
2.2	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following intravenous constant infusion dosing [blood/plasma].	5
2.3	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following extravascular bolus dosing [blood/plasma]. Discussion of the concepts of bioavailability (absolute and relative) and bioequivalence.	5
2.4	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following extravascular multiple bolus dosing [blood/plasma].	5
2.5	Discussion of approaches to solve problems related to the analysis of pharmacokinetic study data obtained after different types of dosing. Discussion of approaches to problem solving involving data from bioavailability and bioequivalence studies. Discussion of approaches to dosage regimen design	5
2.6	<i>Self study of problems and problem solving related to the theoretical concepts outlined above (blood ad urine data analysis)</i>	10
3	Discussion of the processes of absorption, distribution and elimination with respect to how these processes impact the values of rate constants for absorption/distribution/elimination and the values of bioavailability, volume of distribution and clearance.	10
4	Introduction to drug transporters and their impact on the pharmacokinetics of drugs and pharmacokinetic drug-drug interactions.	3
5	Brief introduction to the concept of dose- and time-dependent pharmacokinetics [non-linear pharmacokinetics] and their impact on drug development and clinical use.	3
	Total	60

Books:

1. Clinical Pharmacokinetics and Pharmacodynamics-Concepts and Applications, Rowland M and Tozer TN, Walters Kluwer – Lippincott Williams and Wilkins.
2. Applied Biopharmaceutics and Pharmacokinetics, Shargel L and Yu ABC, Appleton and Lange, International Edition
3. Handbook of Basic Pharmacokinetics including clinical applications, Ritschel WA and Kearns GL, APhA,
4. Basic Pharmacokinetics, Jambhekar SS and Breen PJ, Pharmaceutical Press.
5. Biopharmaceutics and Pharmacokinetics, Venkateshwarlu V, Pharma Book Syndicate
6. Drug Bioavailability- Estimation of solubility, permeability, absorption and bioavailability, van der Waterbeemd H, Lennernas H and Artursson P, Wiley VCH.
7. Modelling in Biopharmaceutics, Pharmacokinetics and Pharmacodynamics – Homogenous and Heterogenous approaches, Macheras P and Iliadis A, Springer

SEMESTER II

BRANCH: CLINICAL PHARMACY

Core 1

Clinical Pharmacy

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Introduction to Clinical Pharmacy	3
1.1	Scope, Objectives and Goals in Health Care.	
1.2	Practice of Clinical Pharmacy in Hospitals and Community.	
2	Understanding the patient	11
2.1	Pharmacist – Patient Interview, Interview Techniques, communication skills.	8
2.2	Patient oriented medical records (POMR): Medication history and records, habits related to use of OTC medications, foods, allergies and sensitivities.	
2.3	Patient follow- up and discharge interview for hospitalised patients.	
2.4	Pharmacological and Biochemical examinations and their significance.	
2.5	Ethics related to medical record	
2.6	Discharge card	
2.7	<i>Self study-Supervision of therapeutic success, side effects and adverse effects</i>	3
3	Therapeutic use of medicine	10
3.1	Drug selection and administration. Problems associated with concomitant therapy.	
3.2	Patient sensitivities, allergies. Precautions during use, Diet control.	
3.3	Reasons for non-compliance, Strategies for improving compliance.	
3.4	Use of drugs and concerns in geriatric, paediatric patients and in pregnancy	
3.5	Drug-drug interactions and drug interactions with food, alcohol and tobacco.	
	Therapeutic drug monitoring (TDM)	6
4.1	Introduction, individualization of drug dosage regimen (variability-genetic, age, weight, disease and interacting drugs).	
4.2	Indications for TDM, protocol for TDM.	
4.3	Pharmacokinetic-pharmacodynamic correlation in drug therapy.	
4.4	TDM of drugs used in following disease conditions: cardiovascular diseases, CNS conditions etc.	
5	Drug formulary, drug utilization review (DUR) including rational drug therapy	3

6	Drug Information	4
6.1	Introduction to information resources	
6.2	Drug information centre (DIC) and Drug information services.	
6.3	Drug literature utilization, selection, evaluation and communication.	
6.4	Role of DIC in ensuring rational use of drugs (RUD).	
7	Standard treatment protocol of selected non communicable diseases/conditions like diabetes, hypertension, stroke, obesity, arthritis, cardiopulmonary dysfunction and fluid and electrolyte imbalance.	10
8	<i>Self study- General concepts of poisoning and toxicology</i> <i>Critical care management:</i> <i>Common life support systems-Acute and chronic renal failure, cardiac and epileptic attack and respiratory failure.</i>	13
	Total	60

Books:

1. J.T. Dipiro, R.L. Talbert, , G.C. Yee, G.R. Matzke, B.G. Wells, L. Michael Posey (eds.),
2. Pharmacotherapy : A Pathophysiologic Approach, 6th ed., The McGraw Hill Companies, Inc., 2005.
3. E.T. Herfindal and D.R Gourley, Text Book of Therapeutics: Drug and Disease Management, 7th ed.,Lippincott Williams & Wilkins, USA, 2000.
4. T.M. Speight and NHG Holford (ed.), Avery's Drug Treatment: Principals and Practice of Clinical Pharmacology and Therapeutics, 4th ed., ADIS Press, Sydney, Australia, 1997.
5. Dennis L. Kasper, Eugene Braunwald, Anthony S. Fauci, Stephen L. Hauser, Dan L. Longo, J. Larry Jameson, and Kurt J. Isselbacher, (Eds.), Harrison's Principles of Internal Medicine, 16th ed., The McGraw Hill Companies, Inc., 2004.
6. Pharmaceutical Practice- A.J Winfield, R.M.E. Richards, Churchil livingstone publication.
7. Drug Interaction Facts, 2003. David S. Tatro.

Core 2

Clinical Research Methodology

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Clinical trials Introduction, drug discovery and drug development, Various phases of clinical trials, study methodology/designs, Inclusion and Exclusion criteria, objectives and endpoints (efficacy and safety), Methods of allocation, blinding and randomization, Informed consent process, study monitoring and its importance, Safety monitoring in clinical trials, BA/BE studies, Post marketing studies	10
2	Documents in clinical study Essential documents in clinical trial- Investigators Brochure (IB), Protocol and amendments in protocol, Case report form (CRF), Informed consent form (ICF), Content of clinical study report (CSR).	4
3	Ethical guidelines in clinical research History, ICH-GCP and its principles, Indian GCP (CDSCO Guidelines), ICMR guidelines- Ethical guidelines	6

	for Biomedical Research on human subjects 2006, Schedule Y 2005, USFDA guidelines for IND, NDA, ANDA applications	
4	Roles and responsibility of various clinical trial personnel as per ICH-GCP Sponsor, Investigator, Monitor, Auditors	3
5	Institutional Ethics Committee (IEC)/ Independent Ethics Committee (IdEC)/Institutional Review Board (IRB)	2
6	Role of Quality assurance in clinical research	2
7	Clinical Data Management and Report Writing	3
8	Pharmacoepidemiology Types, methods, and factors affecting drug utilization, applications of pharmacoepidemiology in health care and rational use of drugs	5
9	Pharmacovigilance Definition, scope and aims of pharmacovigilance, Adverse drug reactions-Classification, mechanism, predisposing factors and causality assessment, Role of clinical pharmacist in reporting, evaluation, monitoring, prevention and management of ADRs.	5
10	Pharmacoeconomics and Outcomes Research Theories and methodologies of Pharmacoeconomics and Outcomes Research. Applications of pharmacoeconomics to pharmacotherapy and managed health care.	5
	Total	45

Books

1. Rick NG. Drugs From Discovery to Approval, second edition, John Wiley & Sons, Inc 2004.
2. Allen Cato, Lynda Sutton, Clinical Drug Trials and Tribulations Second Edition Revised, second edition, Marcel Dekker Inc; 2nd Revised edition March 26, 2002.
3. Ronald D. Mann, Elizabeth B. Andrews. Pharmacovigilance, second edition, John Wiley & Sons Ltd, 2002.
4. Shayne C. Gad, Drug Safety Evaluation, A John Wiley & Sons, Inc. Publication, 2000.
5. Sandy Weinberg. Guidebook For Drug Regulatory Submissions, first edition, A John Wiley & Sons, Inc. 2009.
6. Duolao Wang and Ameet Bakhai Clinical Trials A Practical Guide to Design, Analysis, and Reporting, Remedica 2006.
7. Giovanna di Ignazio, Di Giovanna and Haynes, Principals of Clinical Research, Wrightson Biomedical Pub., 2001
8. R K Rondels, S A Varley, C F Webb, Clinical Datamanagement, Second Edition, John Wiley & Sons Inc, January 1997.
9. Strom BI, Limmel SE. Textbook of Pharmacoepidemiology. Chichester, West Sussex, England: John Wiley & Sons Ltd; 2006.
10. Rascati, Karen L. Essentials Of Pharmacoeconomics. Philadelphia, Pa.: Lippincott Williams & Wilkins, 2009.
11. M. F. Drummond, M. J. Sculpher and G. W. Torrance, Methods for the economic evaluation of health care programmes. Oxford University Press, USA, 2005.

14. Brenda Waning; Michael Montagne; William W McCloskey, Pharmacoepidemiology : principles and practice, New York : McGraw-Hill, 2001
15. Various Guidelines like:
- ICH (International Conference on Harmonisation), GCP for registration of pharmaceuticals for human use. ICH Harmonised Tripartite
 - Guideline for Good Clinical Practice, E6, 1996.
 - ICMR Guideline – Ethical Guidelines for Biomedical Research on Human Subjects.
 - Indian GCP – Central Drugs Standard Control Organization. Good Clinical Practices
 - Guidelines for Clinical Trials on Pharmaceutical Products in India. New Delhi: Ministry of Health; 2001.
 - Pharmacovigilance Programme of India (PvPI)

SEMESTER II

BRANCH : MEDICINAL CHEMISTRY

Core 1

Advanced Pharmaceutical and Medicinal Chemistry

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Enzyme Inhibition	16
1.1	Coverage of basic aspects of enzyme kinetics, catalysis, transition-state theory.	2
1.2	Drug Resistance through alterations of drug uptake, overproduction of enzyme, alterations of the enzyme active site, overproduction of the substrate or new pathways for formation of the product	1
1.3	Drug synergism, concepts and mechanisms.	1
1.4	Reversible enzyme inhibitors – competitive inhibition, non-competitive inhibition, uncompetitive inhibition with suitable examples. Detection of type of inhibition by suitable plotting methods. Concepts of IC_{50} and K_i .	4
1.5	Slow-tight binding inhibitors, covalent enzyme inhibitors and mechanism-based inhibitors with suitable examples. Concept of K_{inact} and K_i for irreversible inhibitors	4
1.6	<i>Self study of specific examples of different types of inhibitors and their design (some examples like COX inhibitors, ACE inhibitors, RT inhibitors, HIV protease inhibitors, aromatase inhibitors, DHFR inhibitors, viral DNA polymerase inhibitors, thymidylate synthase inhibitors and others)</i>	4
2	QSAR	14
2.1	Historical Aspects	1
2.2	Electronic Effects- the Hammett equation, lipophilic effects, experimental measurement of lipophilicity, logP and logD, effect of ionization on logP, calculation of logP and logD, Steric effects- the Taft equation	3
2.3	Hansch Analysis, Free-Wilson method, Topliss operational scheme	2
2.4	Basics of regression analysis - linear and multilinear regression, introduction to PCA, PCR, PLS, ANN and GFA. Correlation coefficients (r^2 and r^2_{pred}), F-test, standard error, validation methods like cross-validation by calculation of q^2 , boot-strap analysis and randomization. Application domain for predictions using a QSAR model.	6

2.5	Design of training and test sets using factorial design	2
3	Peptides and Peptidomimetics	14
3.1	Coverage of peptide structure, biosynthesis of peptides and solid-phase/solution synthesis of peptides.	4
3.2	Design of peptidomimetics by manipulation of the amino acids, modification of the peptide backbone, incorporating conformational constraints locally or globally, α -helix, β -sheet, β - and γ -turn mimetics	4
3.3	<i>Self study of examples of peptidomimetics for some enzymes and receptors like ACE, CCK, bradykinin</i>	4
4	Antisense therapeutic agents	6
4.1	History and principles	2
4.2	Design of antisense oligonucleotides and small interfering RNAs (siRNAs) with some examples	4
5	Molecular Biology, Genetic engineering and Biotechnology in production of biologicals as drugs.	6
5.1	<i>Self study of biotechnology based drugs, vaccines and diagnostic agents with respect to their biological source, their design and the mechanism of their actions</i>	4
	Total	60

Books

1. The Organic Chemistry of Drug Design and Drug Action, Silverman R. B., Academic Press.
2. Textbook of Drug Design and Discovery, Eds. Krogsgaard-Larsen P., Liljefors T., Madsen U., Taylor & Francis.
3. Medicinal Chemistry: An Introduction, Thomas G, Wiley.
4. Peptide and Protein Design for Biopharmaceutical Applications, Ed Jensen K. J., Ch. 3 Aspects of Peptidomimetics by Maes V., Tourwé D., John Wiley & Sons, Ltd, Chichester, UK.
5. Comprehensive Medicinal Chemistry, Series Ed., Hansch C., Pergamon Press.
6. Burgers Medicinal Chemistry, Drug Discovery and Development, Wiley.

Core 2

Drug Metabolism

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Introduction to xenobiotic/drug metabolism	6
1.1	Introduction to xenobiotic/drug metabolism and its relation to other defence systems (Physical barriers, excretion, immune system).	2
1.2	Types of reactions (I and II), consequences of drug metabolism (DM) [inactivation, bioactivation, prodrugs], organs of DM, localization of drug metabolizing enzymes, factors affecting drug metabolism.	4

2	Cytochrome P450s: Introduction to the family of enzymes, their classification and nomenclature.	20
2.1	Introduction to the family of enzymes, their classification and nomenclature.	2
2.2	CYP450 catalytic cycle, different types of reactions catalyzed by CYP450s and the mechanisms of catalysis.	8
2.3	Human CYP450s involved in DM, their distribution and properties, typical substrates, specific probe substrates, specific inhibitors, induction of CYPs and specific inducers	7
2.4	Genetic polymorphism in CYP450 expression	3
3	NON P450 enzymes	20
3.1	Introduction to NON P450 enzymes involved in drug metabolism	05
3.2	<i>Self study of NON P450s - glucuronosyltransferases, sulfotransferases, glutathione S-transferases, N-acetyl transferases, xanthine oxidase, aldehyde oxidase, esterase, epoxide hydrolase, nitro/azo reductases and FMO [on lines similar to that specified for CYPs as listed above].</i>	15
4	Introduction to methods for studying DM. Discussion of in vitro and in vivo tools, along with their advantages and limitations {recombinant enzymes, subcellular fractions, hepatocytes, liver slices, perfused liver and whole animal studies}.	5
5	Discussion of types of DM studies – metabolic stability, cross species comparisons, metabolite profiling and identification, reaction phenotyping, CYP inhibition and CYP induction studies.	6
6	Introduction to <i>in silico</i> drug metabolite predictions and associated algorithms.	3
	Total	60

Books:

1. Comprehensive Medicinal Chemistry, Series Ed., Hansch C., Pergamon Press.
2. Wilson and Gisvold's, Textbook of Organic Medicinal and Pharmaceutical Chemistry, Lippincott-Raven
3. Foye's Principles of Medicinal Chemistry, Lippincott Williams and Wilkins.
4. Drug Metabolizing Enzymes-Cytochrome P450 and Other Drug Metabolizing Enzymes in Drug Discovery and Development, Lee JS, Obach SR and Fisher MB, Marcel Dekker, Fontis India, 2003
5. Pharmaceutical Profiling in Drug Discovery for Lead Selection, Borchardt RT, Kerns EH, Lipinski CA, Thakker DR and Wang B, AAPS Press, 2004
6. Drug Metabolism – Current Concepts, Ionescu C and Caira MR, Springer International Edition
7. Handbook of Drug Metabolism, Woolf TF, Marcel Dekker, 1999.

SEMESTER II

BRANCH : PHARMACEUTICAL BIOTECHNOLOGY

Core 1

Pharmaceutical Biotechnology

4 hrs/week

Unit	Course Content (Topics)	Hours
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1	Production and Control of Biotech derived products	25
1.1	Recombinant DNA products – insulin, growth hormone, erythropoietin, cytokines	5
1.2	Vaccines – attenuated virus, genetic alterations of live virus as a vector of other pathogens (recombinant virus or recombinant vaccinia virus)	7
1.3	Diagnostic proteins – protein A, protein G, antibodies	4
1.4	Quality control testing of biotech products – determining impurities, contamination -viral, bacterial endotoxin, rabbit pyrogen test, sterility, protein identification, finger prints by electrophoresis, isoelectric focussing, immunogenicity, partial sequence analysis.	9
2	Plant biotech products	10
2.1	Substances produced by plant cell culture	2
2.2	Transgenic plants and their application	4
2.3	Biotransformations with plant cell culture	4
3	Biotech products through fermentation	13
3.1	Fermentation – batch, continuous fermentation	2
3.2	Role of bioengineering in fermentation – geometry of fermentation tanks, design of impellers, agitation systems and environmental conditions of fermentation	3
3.3	Fermentative production of important secondary metabolites – penicillins, amino glycosides polyene macrolides, macrolides, anthracyclines	3
3.4	Principles of downstream processing of fermentation products	3
3.5	Unit operations and techniques employed inn downstream processing of fermentation products, microbial strain selection and preservation methods	3
3.6	Genotype and phenotype variation of characters of microbes	
4	Self Study - Biotransformation	12
4.1	<i>Biotransformation principles and industrial applications in the production of chemicals and drugs</i>	
4.2	<i>Immobilization of enzymes, proteins and their applications – biosensors, enzyme electrodes, immunosensors, optical sensors</i>	
	Total	60

Books:

1. Biotechnology, H. J. Rechm, G. Reed. Vols 1 – 12, A. Pulher, P. Stadler Eds, Weinheim, New York
2. A text book of Biotechnology, H. D. Kumar Affiliated, East – West Press Pvt. Ltd.
3. Genetic Engineering Fundamentals, Karl Kammer, Meyer Virginia, C. Clark.
4. Genes V, Benjamin Lewin, Oxford University Press.
5. Methods in Plant Molecular Biology and Biotechnology, Bernard R Glick, John E Thompson, CRC Press.
6. Genetic and Biochemistry of Antibiotics Production, Leo C Vining, Colin, Stuttard Butterworth, Heinemann.
7. Biotechnology – Applications and Research, Paul N Chermisinoff, Robert P Ouellett, Technomic Publishing Co. Inc.

8. Transgenic Plants: A production systems for industrial and pharmaceutical proteins, Meran R. L. Owen, Jan Pen, John Wiley and Sons.
9. Biotechnology of antibiotics, William R Strohl, Marcel Dekker.
10. Molecular Biochemistry – Therapeutic applications and strategies, Sunil Maulik and Salil D Patel, John Wiley and Sons, Inc.

Core 2

Basic Molecular Biology

4 hrs/week

Unit	Course Content (Topics)	Hours
1	The beginnings of molecular biology	1
2	DNA Structure and Role of DNA	16
2.1	Organization of the genome, building from nucleotides to chromatin	6
2.2	The genetic code and its relationship to protein structure	2
2.3	DNA replication, Telomere maintenance, mechanisms of DNA repair, DNA recombination	8
3	The versatility of RNA	23
3.1	Transcription and translation in prokaryotes; Transcription and translation in eukaryotes	8
3.2	Epigenetics and monoallelic gene expression	3
3.3	RNA processing and post-transcriptional gene regulation	8
3.4	Mechanisms of translation	4
4	Genetically modified organisms: Use in basic and applied research	14
4.1	Recombinant DNA technology, molecular cloning, & some tools for analyzing gene expression	8
4.2	Genome analysis: DNA typing; Genomics and beyond; Medical molecular biology: applications in Cancer and Gene therapy; Genes and behaviour	6
5	Plant tissue culture and animal cell culture	6
	Total	60

Books:

1. Genes IX, Ed Benjamin Lewin. Oxford University Press.
2. Molecular Cell Biology, Lodish H, Berk A, Zipursky S L, Matsudaira P., Baltimore D, Darnell J, Publisher W. H. Freeman.
3. Molecular Biology of the Cell, Alberts Publisher Garland Science.
4. Watson, J. D. Tania A. Baker, Stephen P. Bell, Alexander Gann, Michael Levine, Richard Losick, Molecular Biology of the Gene, Benjamin Cummings; 6th Edition, 2007.
5. Molecular Biology in Medicinal Chemistry, Dingemann Th, Steinhilber D and Folkers G, Wiley-VCH, Germany
6. Basic Principles of Gene Manipulation, Primrose SB, Twyman RM and Old RW, Blackwell.
7. Molecular Biology and biotechnology, Walker JM and Rapley R, Royal Society of Chemistry

SEMESTER II

BRANCH: DRUG DELIVERY TECHNOLOGY

Core 1

Advanced Pharmaceutics – I

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Solids – oral SR system	13
1.1	<i>Self study-Overview of Single oral unit SR systems .</i>	3
1.2	<i>Self study-Structure and physiology of GIT.</i>	1
1.3	Mechanism of Release & Release kinetic equations. Types – Diffusion controlled, Dissolution controlled, Reservoir, Matrix, Osmotic systems, Ion exchange systems Mucosal drug delivery systems- buccal, gingival, sublingual.	4
1.4	Multiparticulate systems-pelletization (emphasis on extrusion and spheronization). Orodispersible systems. Pulsatile Drug delivery systems.	5
2	Parenteral SR systems	12
2.1	Need and concept, routes employed	1
2.2	Approaches- aqueous systems (complexation, use of polymers), aqueous suspensions (depot injections, microspheres, magnetic microspheres), Oily solutions & suspensions, Emulsions (Microemulsions, multiple emulsions,), Implants (in detail-concept, properties desired, various approaches), prodrugs (chemical modifications), infusion pumps.	6
2.3	<i>Self study-Biopharmaceutical aspects, Sterilization & stability issues</i>	2
2.4	Characterization-special emphasis on release studies	1
2.5	Issues related to Safety, Toxicity & Tissue Injury	2
3	Specialized Emulsions	9
3.1	Microemulsions, Multiple emulsions, Self Emulsifying Drug Delivery systems & SMEDDS; Formulation and phase behaviour; Preparation & Characterization; Bioavailability Aspects; Applications.	6
3.2	<i>Self study-Theories of Emulsification, Factors influencing type of emulsion formed.</i>	3
4	Gastro-retentive Drug Delivery Systems	8
4.1	<i>Self study –Introduction; concept of absorption window; need for GRDDS, gastric motility; principles of Gastro-retention; Factors controlling performance of GRDDS.</i>	3
4.2	Different Approaches- High density systems, floating systems, muco-adhesive systems, Expandable systems, Magnetic systems, Superporous Hydrogels	4
4.3	Evaluation.	1
5	Ocular drug delivery systems.	7

5.1	<i>Self study-Structure and physiology of eye; Drug absorption and disposition in the eye.</i>	2
5.2	Methods to prolong ocular drug residence with emphasis on mucoadhesive systems.	1
5.3	Intraocular inserts; Nonerodible inserts / Erodible inserts. Novel ophthalmic drug delivery systems, Nanoparticles , liposomes and prodrugs. Ocular penetration enhancers.	4
6	Transdermal Drug Delivery Systems	7
6.1	<i>Self study-Structure and physiology of skin.</i>	1
6.2	Principles of skin permeation. Kinetics of skin permeation & penetration enhancers. Types (Gels, Patches/films) , Pressure sensitive adhesives.	3
6.3	Development & evaluation – <i>in vitro, in vivo.</i>	1
6.4	Iontophoresis.	1
6.5	Recent advances –use of microneedles in transdermal drug delivery.	1
7	Introduction to Pharmaceutical Processing Development. (As per ICH guidelines)	4
7.1	Elements in Pharmaceutical development <ul style="list-style-type: none"> • Target product profile • Critical Quality Attributes. • Linking Material Attributes & process parameters to CQA's Risk Assessment • Design space • Control Strategy • Product Lifecycle management & continual improvement. 	3
7.2	Submission of Pharmaceutical Development and related information in CTD format. <i>Relevant Examples.</i>	1
Total		60

Books:

1. Targeted and Controlled Drug Delivery: Novel Carrier Systems by Vyas SP, Khar RK, CBS Publishers and Distributors, 1stedn, 2002.
2. Controlled and Novel Drug Delivery by Jain NK, CBS Publishers and Distributors, 2008.
3. Controlled Drug Delivery: Fundamentals and Applications by Robinson JR, Lee VHL, Dekker, 2ndedn, Vol 29, 1987.
4. Novel Drug Delivery System by Chien YW, 2ndedn, Vol 50, Informa Healthcare, 2003.
5. Progress in Controlled and Novel Drug Delivery Systems by Jain NK, CBS Publishers and Distributors; 2004.

6. Ophthalmic Drug Delivery Systems, Mitra AK, 2nd edn., Drugs and Pharmaceutical Sciences Series, Vol. 130, Marcel Dekker, 2003.
7. Polymeric drug delivery system, Kwon GS, Marcel Dekker, Vol 148, 2005.
8. Nanoparticulate Drug Delivery System by Thassu D, Deleers M, Pathak Y, Marcel Dekker, Vol 166, 2007.
9. Controlled Drug Delivery- Challenges and Strategies by Park K, American Chemical Society, 1997.
10. Colloidal Drug Delivery System by Kreuter J, Marcel Dekker Vol 66, 1994.
12. www.ich.org
13. Pharmaceutical Dosage Forms: Disperse Systems by Lieberman HA, Rieger MM, Banker GS, Marcel Dekker, Vol 3, 2nd edn, 2005.
14. Pharmaceutical Emulsions and Suspensions by Nielloud F, Marti- Mestres G, Marcel Dekker, Vol 105, 2000.
14. Controlled Release Systems Fabrication Technology by Dean STH, CRC Press, Vol 1, 1988.
15. Bioadhesive Drug Delivery Systems by Mathiowitz.E , Chickering DE , Lehr CM, Marcel Dekker Vol 98 ,1999.
16. Pharmaceutical Skin Penetration Enhancement by Walters. K A, Hadgraft J, Marcel Dekker, Vol 59, 1993.
17. Percutaneous Absorption by Bronaugh RL, Maibach HI, Taylor and Francis, 3rd edn, Vol 97, 2005.
18. Transdermal Controlled Systemic Medication by Chien YW, Marcel Dekker, Vol. 31, 1987.
19. Oral Mucosal Drug Delivery by Rathbone MJ, Marcel Dekker, Vol 74, 1996.
20. Modified Release Drug Delivery Technology by Rathbone MJ, Hadgraft J, Roberts MS, Lane ME, Informa Healthcare, 2nd edn, Vol 183(1), 2008.
21. Pharmaceutical Pelletization Technology, Ghebre-sellassie. I, , Marcel Dekker, Vol. 37

Core 2

Models for DDS Evaluation

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Pharmacodynamic models for evaluation of DDS containing drugs of various categories e.g. cardiovascular agents, antidiabetic, anti-inflammatory, antiepileptic, anticancer, hepatoprotectives, analgesics, antistress, antiasthmatic and antitussives.	20
2	In vitro cell culture techniques for evaluation of drug permeation from DDS, including isolation, maintenance of cell lines, culturing monolayers, evaluation of drug transport	8
3	In vitro/ex vivo models for evaluation of drug absorption	5
4	In vitro cytotoxicity evaluation using cell cultures and techniques such as MTT assay, dye uptake etc	5
5	Toxicity testing – in vitro – In vitro toxicity testing and its application to safety evaluation, general perspectives, in vitro trends and issues, ocular and cutaneous irritation, validation of in vitro toxicity tests – acute, sub acute and chronic toxicity testing, biochemical basis of toxicity, design of toxicological studies, quality assurance in toxicological studies, toxicity by routes – parenteral, oral, percutaneous and inhalation, target organ toxicity exemplified by hepatotoxicity and cutaneous (dermal) toxicity.	10

	Total	48
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Books:

1. Bioassay Techniques for drug development, Atta Ur Rahman, M. Iqbal Choudhar, William J Thomsen.
2. In vitro Methods in Pharmaceutical Research, Eds. J. V. Castell, M. J. Gomer, Lechon, Academic Press
3. In vitro Toxicity Testing, John M Frazier.
4. General and Applied Toxicology, Bryan Ballantyne, T Marrs and P. Turner.

Elective Subjects

Elective 1

Rational Drug Design

4 hrs/week

Unit	Course Content (Topics)	Hours
1.0	Molecular Mechanics and the forcefield. General form of a generic force field, force field parametrization.	5
1.1	<i>Self Study – Comparison between the different forcefields in existence at present time</i>	1
2.0	Energy minimization	6
2.1	Steepest descents, conjugate gradients, Newton Raphson method, advantages and limitations of each method	
3.0	Conformational analysis	10
3.1	Systematic search, Monte Carlo simulations, Molecular dynamics simulations, distance geometry, strengths and limitations of each method	
4.0	Docking	10
4.1	Docking by energy minimization, superimposition, molecular dynamics, Metropolis Monte Carlo, genetic algorithms, build-up approach. Different types of scoring function, e.gs of successful application of docking.	8
4.2	<i>Self Study – Successful applications of docking</i>	2
5.0	de novo ligand design	10
5.1	Classes of de novo ligand design – active site analysis methods, whole-molecule methods, connection methods, random connection and disconnection methods, e.gs of successful application of <i>de novo</i> ligand design	
5.2	Fragment based drug design	2
5.3	<i>Self Study – Successful applications of de novo drug design</i>	2
6.0	Pharmacophore modelling	9
6.1	Techniques of developing a pharmacophore map covering both ligand based and receptor based approaches, incorporating additional geometric features into a 3D pharmacophore, use of a pharmacophore model in drug design,.	7

6.2	<i>Self study - Successful e.g. of pharmacophore maps in drug design</i>	2
7.0	Virtual Screening based on similarity, docking, pharmacophore maps and filters for drug-likeness and ADME	3
8.0	3D-QSAR	6
8.1	CoMFA and CoMSIA. Mention of other 3D-QSAR techniques and introduction to the 4 th , 5 th and 6 th dimension in QSAR.	4
8.2	Self Study – 3D-QSAR methods other than CoMFA and CoMSIA	2
	Total	60

Books

1. Molecular Modelling – Principles and Applications, Leach A. R., Prentice Hall.
2. Practical Application of Computer-Aided Drug Design, Ed. Charifson P., Marcel Dekker Inc.
3. 3D QSAR in Drug Design: Theory, Methods and Applications, Ed. Kubinyi H., Lediten ESCOM.
4. Molecular Modeling and Simulation -An Interdisciplinary Guide, Schlick T., Springer.

Elective 2

Spectroscopic Structural Elucidation

4 hrs / week

Problems of structural elucidation involving the following techniques:		
<ul style="list-style-type: none"> • UV spectroscopy. • IR spectroscopy. • ¹H-NMR spectroscopy. • ¹³C-NMR spectroscopy. • Mass Spectrometry. 		
The problems should cover the following aspects:		
Unit	Course Contents (Topics)	Hours
1	Calculation of λ max for dienes, α,β –unsaturated ketones by UV spectroscopy. <i>Self study-practice problems (1 hr)</i>	5
2	Prediction of characteristic IR bands , NMR spectra (¹ H NMR) –chemical shift, splitting pattern and ratio of proton intensity, (¹³ C NMR)-number of signals, chemical shift and splitting pattern, mass fragmentation patterns. <i>Self study-practice problems (2 hrs)</i>	10
3	Distinguishing compounds using UV / IR / ¹ H NMR / ¹³ C NMR and /or Mass spectrometry. <i>Self study-practice problems (2 hrs)</i>	10
4	Interpretation of mass spectra with explanation of fragmentation patterns. <i>Self study-practice problems (2 hrs)</i>	9
5	Problems involving structure elucidation by- UV / IR / ¹ H NMR / ¹³ C NMR and / or Mass spectrometry, <i>Self study- practice problems (8 hrs)</i>	26
	Total	60

Books:

- 1 Introduction to Spectroscopy by D.L.Pavia, G.M.Lampman & G.S.Kriz, Latest edition, Thomson Brooks/Cole, United States.
- 2 Spectrometric Identification of Organic compounds by Robert.M.Silverstein & Francis.X.Webster, D.J.Kiemle, Latest edition, John Wiley & Sons.
- 3 Organic Spectroscopy by William Kemp.
- 4 Applications of absorption spectroscopy of organic compounds by John Robert Dyer, Prentice Hall, London.

Elective 3**Title of the Course: Advanced Biochemistry****4 hrs/week**

Unit	Course Content (Topics)	Hours
1	Proteins	15
1.1	Structure – primary, secondary, tertiary, quaternary; motifs, structural and functional domains, protein families and macromolecular assemblies	5
1.2	Mechanisms for regulating protein function: Protein-protein interactions, interaction with ligands; Ca ²⁺ and GTP as modulators, cyclic phosphorylation and dephosphorylation, proteolytic cleavage.	2
1.3	Purification and characterisation of proteins: electrophoresis, ultracentrifugation and liquid chromatography, use of biological assays, use of radioisotopes; MS, X-ray crystallography, NMR and homology modelling to determine structures; amino acid analysis; cleavage of peptides; protein sequencing.	4
1.4	Protein biosynthesis: translation machinery in prokaryotic and eukaryotic systems; comparison of similarities and differences, drug affecting protein biosynthesis and protein function	4
2	DNA and nucleic acids	15
2.1	DNA, RNA structure, nomenclature, double helix, conformations, higher order packing and architecture of DNA, transcription and replication of DNA – mechanisms in prokaryotic and eukaryotic systems, DNA repair mechanisms, drug affecting nucleotide biosynthesis, RNA and DNA biosynthesis and RNA and DNA function	15
3	Carbohydrates	8
3.1	Mono, di and polysaccharides and their nomenclature, stereochemistry, types of linkages; conjugates of carbohydrates with other molecules – glycoproteins, glycolipids, proteoglycans, lipopolysaccharides and their biological roles	8
4	Lipids	7
4.1	Classification, nomenclature, stereochemistry, storage lipids, membrane lipids, lipids as secondary messengers and cofactors, biological role of lipids, drug affecting lipid metabolism.	7
5	<i>Self study of protein superfamilies, N and C terminal sequencing, DNA structures other than B-DNA, DNA sequencing, DNA pyrosequencing, cerebrosides, sphingolipids.</i>	15
	Total	60

Books:

1. Principles of Biochemistry, Lehninger, Nelson D.L., C.B.S Publishers, New Delhi.
2. Biochemistry, Stryer L, W. H. Freymont & Co., New York.
3. Molecular Cell Biology, Lodish H, Darneu J, Scientific American Books, N.Y.
4. Biochemistry- The chemical reactions of living cells, Vol 1 &2, Metzler DE, Elsevier Academic Press.
5. Biochemistry, Berg JM, Tymoczko JL and Stryer L, WH Freeman and Company and Sumanas Inc.
6. Biomacromolecules- Introduction to structure, function and informatics, Stan Tsai C, Wiley-Liss
7. Protein: Structure and Molecular properties, Thomas E Creighton, W. H. Freeman.
8. Physical Biochemistry- Principles and applications, Sheehan D, Wiley-Blackwell

Elective 4

Pharmacokinetics

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Introduction to pharmacokinetics and its utility in drug design and dosage regimen design. Definitions of absorption, distribution, metabolism, excretion, elimination. Different approaches for determination of pharmacokinetics of drugs – non-compartmental, physiological, and compartmental modeling. Assumptions involved in the evolution of single and multi-compartment models.	4
2	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following different dosing methods/protocols [blood/plasma/urine sampling]	40
2.1	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following intravenous bolus dosing [blood/plasma/urine sampling]	5
2.1	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following intravenous multiple bolus dosing [blood/plasma]	5
2.2	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following intravenous constant infusion dosing [blood/plasma].	5
2.3	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following extravascular bolus dosing [blood/plasma]. Discussion of the concepts of bioavailability (absolute and relative) and bioequivalence.	5
2.4	Discussion (including mathematical description and equations) of the pharmacokinetics of drugs showing one compartment pharmacokinetics following extravascular multiple bolus dosing [blood/plasma].	5
2.5	Discussion of approaches to solve problems related to the analysis of pharmacokinetic study data obtained after different types of dosing. Discussion of approaches to problem solving involving data from bioavailability and bioequivalence studies. Discussion of approaches to dosage regimen design	5
2.6	<i>Self study of problems and problem solving related to the theoretical concepts outlined above (blood ad urine data analysis)</i>	10
3	Discussion of the processes of absorption, distribution and elimination with respect to how these processes impact the values of rate constants for absorption/distribution/elimination and the values of bioavailability, volume of distribution and clearance.	10

4	Introduction to drug transporters and their impact on the pharmacokinetics of drugs and pharmacokinetic drug-drug interactions.	3
5	Brief introduction to the concept of dose- and time-dependent pharmacokinetics [non-linear pharmacokinetics] and their impact on drug development and clinical use.	3
	Total	60

Books:

1. Clinical Pharmacokinetics and Pharmacodynamics-Concepts and Applications, Rowland M and Tozer TN, Walters Kluwer – Lippincott Williams and Wilkins.
2. Applied Biopharmaceutics and Pharmacokinetics, Shargel L and Yu ABC, Appleton and Lange, International Edition
3. Handbook of Basic Pharmacokinetics including clinical applications, Ritschel WA and Kearns GL, APhA,
4. Basic Pharmacokinetics, Jambhekar SS and Breen PJ, Pharmaceutical Press.
5. Biopharmaceutics and Pharmacokinetics, Venkateshwarlu V, Pharma Book Syndicate
6. Drug Bioavailability- Estimation of solubility, permeability, absorption and bioavailability, van der Waterbeemd H, Lennernas H and Artursson P, Wiley VCH.
7. Modelling in Biopharmaceutics, Pharmacokinetics and Pharmacodynamics – Homogenous and Heterogenous approaches, Macheras P and Iliadis A, Springer

Elective 5

Molecular Biology

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Introduction to biological macromolecules	8
2	DNA replication, transcription, gene splicing, gene cloning, gene expression	10
3	Genotype and phenotype variation of characters and microbes	8
4	Genetic organization of prokaryotic and eukaryotic cells	8
5	Protein biosynthesis and regulation – induction, repression of enzymes, catabolic expression	8
6	Methods of alteration of genetic apparatus – genes- mutation, transition, recombination, protoplast fusion	8
7	Proteomics and genomics	10
	Total	60

Books:

1. Molecular Biotechnology, Principles and applications of recombinant DNA, Glick B. R. And Pasternak J. J. ASM Press, Washington DC

- Principles of Genome Analysis and Genomics, Primrose S. B. And Twyman R. M., Blackwell Publishing, U.K.
- Gene Biotechnology and Techniques, Gen Engg, Mutagenesis, Separation Technology, Chirirjian J. G, Jones and Bartlett Publishers
- Pharmaceutical Biotechnology – An introduction for Pharmacists and Pharmaceutical Scientists, Crommelin D. A and Sindelar R. D., Harwood Academic Publishers.

Elective 6

Green Chemistry

4 hrs/week

Unit	Course Content (Topics)	Hours
1	Introduction to the concepts of Green Chemistry – history, need, goals, limitations, obstacles and opportunities	5
1.1	Introduction to the principles of Green Chemistry – prevention of waste/by products, maximum incorporation of the materials used in the process into the final product (atom economy), green metrics, prevention/minimization of hazardous/toxic products, designing safer chemicals –basic approaches, selection of appropriate auxiliary substances (solvents, separation agents etc), energy requirements for reactions, selection of starting materials, renewable starting materials, avoidance of unnecessary derivatization – careful use of blocking/protecting groups	15
1.2	Microwave assisted organic synthesis; photochemical transformations; sonication; solid phase transformations; aqueous phase transformations; enzymatic transformations; etc	8
1.2	<i>Self Study - transformations using ionic liquids, PEG, polymer supported reagents</i>	4
2	Application of green synthetic reactions, green starting materials, green reagents, green solvents and reaction conditions, green catalysis and examples of green synthesis, green analytical methods	13
2.1	<i>Self Study – Examples of Green synthesis</i>	3
3	Future trends in green chemistry – oxidation reduction reagents and catalysts; biomimetics and multifunctional reagents; combinatorial green chemistry; solventless reactions; non-covalent derivatization; biomass conversion; emission control; biocatalysis	12
	Total	60

Books

- Green Chemistry: Theory and Practice, Anastas P T and Warner J C, Oxford University Press.
- Green Chemistry: Introductory Text, Lancaster M, RCS London
- Introduction to Green Chemistry, Ryan M. A., Tinnensand M., American Chemical Society (Washington).
- Handbook of Green Chemistry and Technology, Clarke J and Macquarrie D, Blackwell Publishing.
- Green Chemistry – Greener alternative to synthetic organic transformations, Ahluwalia V K, Narosa Publications, New Delhi.
- Organic Synthesis – Special Techniques, Ahluwalia V K and Aggarwal R, Narosa Publications.

Elective 7

Drug Regulatory Affairs

4 hrs/week

No.	Course Contents (Topics)	Hours
1.0	Need for Regulations	1
2.0	Indian Regulations	15
2.1	Introduction to Indian Regulations	1
2.2	Drugs & Cosmetic Act & Rules - Overview and recent amendments	5
2.2.1	<ul style="list-style-type: none"> Schedule DI and DII (Registration and Import) 	
2.2.2	<ul style="list-style-type: none"> Schedule M 	
2.2.3	<ul style="list-style-type: none"> Schedule Y 	
2.2.4	<ul style="list-style-type: none"> Central Drug Laboratories 	
2.3	ICMR guidelines for ethical considerations in biomedical research on human subjects	1
2.4	BA – BE studies	2
2.5	New drug application	1
2.6	Insurance, Compensation and Indemnification of trial subjects	1
2.7	Expert Referral	1
2.7.1	<ul style="list-style-type: none"> IBSC, RCGM 	
2.7.2	<ul style="list-style-type: none"> ICMR 	
2.7.3	<ul style="list-style-type: none"> NDAC 	
2.7.4	<ul style="list-style-type: none"> CBBTDEC 	
2.8	WHO GMP Certification, FSC and CoPP procedure	1
2.9	Procedures for obtaining Test license (Form 29 and Form 11); Export NOC	1
2.10	Loan license / Contract manufacturing	1
3.0	US Regulations	6
3.1	Introduction to US Regulations	1
3.2	Hatch Waxman Act and amendments, FDA Medicare Modernization Act	
3.3	Introduction to Orange Guide and 21-CFR	1
3.4	Investigational new drug (IND) filing	1
3.5	US Drug Master File (DMF) filing, amendments and annual reports	1
3.6	Abbreviated New Drug Application (ANDA) filing	1
3.7	New Drug Application (NDA) filing	

3.8	Post approval changes	1
4.0	European Regulations	6
4.1	Introduction to European Regulations	1
4.2	Active Substance Master File (ASMF) filing	1
4.3	CEP filing	
4.4	Marketing Authorization and filing procedures	2
4.4.1	<ul style="list-style-type: none"> National Procedure 	
4.4.3	<ul style="list-style-type: none"> Mutual Recognition Procedure (MRP) 	
4.4.4	<ul style="list-style-type: none"> Decentralized Procedure (DCP) 	
4.4.5	<ul style="list-style-type: none"> Centralized Procedure (CP) 	
4.5	Handling variations	
4.6	Clinical Trial Regulations in EU	2
5.0	Other applicable Regulations and Guidelines	10
5.1	Overview of ICH guidelines	1
5.2	CTD format of dossier	1
5.3	eCTD filing procedure	
5.4	21- CFR Part 11	1
5.5	Audits and Inspections, FDA 483's – Lessons learnt	1
5.6	Overview of registration process in other geographies	1
5.7	Biological license application (BLA)	1
5.8	Medical Device Registration process	1
5.9	Regulations governing Stem Cell therapeutics	1
5.10	Introduction to Pharmacovigilance and Drug Safety	1
5.11	Orphan Medicinal Products	1
6.0	Intellectual Property Rights (IPR)	4
6.1	Overview of patents from regulatory perspective	
6.2	PCT application & general rules	
6.3	WTO / GATT system	
6.4	TRIPS Agreement	
6.5	Compulsory licensing	

6.6	Patent search, drafting and filing procedure	
6.7	Patent infringement analysis	
6.8	Trademark/ copyright filing procedures	
	Total	42

Books

1. Good Drug Regulatory Practices: A Regulatory Affairs Quality Manual (Good Drug Development Series, Vol 1, [Helene I. Dumitriu](#))
2. <http://www.amazon.com/Good-Drug-Regulatory-Practices-Development/dp/1574910515>
3. Guide to Drug Regulatory Affairs / Buch, [Brigitte Friese](#).
4. Drugs and Cosmetics Act, 1940 and Rules, 1945.

Useful links:

5. <http://www.cdsc.nic.in/>
6. <http://clinicaltrials.gov/>
7. <http://dbtbiosafety.nic.in/>
8. <http://www.emea.europa.eu/>
9. <http://www.ich.org/>
10. <http://www.fda.gov/>

Elective 8

Quality Assurance Systems

4 hrs / week

Unit	Course Contents (Topics)	Hrs
1	Regulatory basis for validation: US FDA guidelines (cGMP guidelines, 21 CFR 210-211), EU guidelines, WHO guidelines.	5
2	Terminology and validation overview:	10
2.1	<i>Self Study: Validation versus verification, testing, calibration and qualification.</i>	3
2.2	Concepts of DQ, IQ, OQ and PQ.	3
2.3	Concepts of Prospective validation, retrospective validation, Concurrent and revalidation. Validation Master Plan.	4
3	Validation of Equipment	10
3.1	Dry Powder Mixers	1
3.2	Fluid Bed and Tray dryers	1
3.3	Tablet Compression Machine	2
3.4	<i>Self study :Dry Heat Sterilization/Tunnels</i>	1
3.5	Autoclaves	2
3.6	Capsule filling machines.	1
3.7	Validation of Integrated lines by media fill test.	2
4	Utilities Validation	7
4.1	Validation of Pharmaceutical Water System & pure steam,	2
4.2	Validation of HVAC system	3
4.3	Validation of Compressed air	2
5	Cleaning Validation: <i>Self study:Cleaning of Equipment, Cleaning of Facilities.</i>	4
6	Analytical Method Validation: General principles of analytical method validation, Validation of following analytical Instruments	6

6.1	HPLC	2
6.2	Dissolution test apparatus	2
6.3	U.V./Visible spectrophotometers	2
7	Process Validation:	13
7.1	<i>Self study : Prospective, concurrent, retrospective & revalidation Self-study</i>	1
	Process validation of following formulations	
7.2	Uncoated / Coated tablets	2
7.3	Hard gelatin Capsules	2
7.4	Ampoules & Vials	2
7.5	<i>Self study:Ointment/Creams</i>	2
7.6	<i>Self study :Liquid Orals</i>	2
7.7	Transdermal patches (Matrix systems)	2
8	<i>Self study-Computer system validation in controlling the manufacturing process.</i>	2
9	Process Analytical Technologies (PAT) and Quality by Design (QbD) (US FDA)	3
	Total	60

BOOKS:

1. Validation and Qualification in Analytical Laboratories by Ludwig Huber, Second edition (2007), Informa Health Care, New York, London.
2. Pharmaceutical Process Validation by R.Nash and Wachter, Volume 129, Latest edition, Marcel Dekker Inc, New York.
3. GMP for Pharmaceuticals by Sidney H. Willing, Fifth edition, Marcel Decker Series, New York.
4. United States Pharmacopoeia & Indian Pharmacopoeia.
5. Validation of Pharmaceutical process, F. J. Carleton and J. Agalloco, Marcel Dekker Inc.

INTERNET REFERENCES:

1. www.fda.gov (US FDA guidelines for PAT and QbD).
2. www.ich.org
3. WHO publications on related topics.
4. EMEA guidelines

Elective 9

Pharmaceutical Quality Management

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Concept of-	8
1.1	Total Quality Management (TQM)	2
1.2	Quality control and quality assurance	2
1.3	Quality control laboratory responsibilities	2
1.4	<i>Self study: Good laboratory practices</i>	2
2	GMP	8
2.1	Organization of pharmaceutical manufacturing unit, production management,	4

Unit	Course Contents (Topics)	Hours
2.2	<i>Self study : Revised schedule M.</i>	4
3	Personnel:	12
3.1	<i>Self study: Introduction, Human resource development, Qualification Experience and Training, Responsibilities, Personal Hygiene and Gowning.</i>	6
3.2	Location, Plant layout, Lighting, Sewage, Water handling-Sewage, Refuse and Disposal, Washing and toilet facility, Sanitation, Controls of contamination and Environmental controls.	6
4	Materials Management:	8
4.1	API's, raw materials & packaging materials, Purchase specifications, Selection of vendors, Intermediates & Finished products, Rejected and Recovered materials, Recalled products, Reagents & culture media, Reference standards, Waste materials.	6
4.2	Warehousing- Good Warehousing Practices, distribution and records.	2
5	Manufacturing Operations and Control:	8
5.1	<i>Self study: Sanitation of Manufacturing Premises, Line clearance, Mix-ups and Cross contamination, Processing and holding of Intermediates and Bulk Products</i>	3
5.2	Packaging, I.P.Q.C., Release and storage of Finished Product, Process Deviations and Incidents, Drug product inspection, Yield calculations	3
5.3	Expiry dating, Manufacturing record review and approval.	2
6	Documentation and Records: In-process and Product Release Specifications, Master production and control record, Batch production and control record, Standard Operating Procedures (SOP), Change Control, Site master file.	6
7	Post Operational Activities:	5
7.1	Distribution, Complaints and recalls, evaluation of complaints, Recall procedures, related records and documents.	2
7.2	Outsourcing: Facility audit, Manufacturing, Packaging, Analytical, Clinical and other services outsourcing.	3
8	Site and Plant security: Security personnel, Entry procedures to site & plant, Internal security, Vehicle parking, Fuel storage, Canteen & cooking, Garden & horticulture.	2
9	Audits: Principle of Quality audit, Plant level, Department wise documentation.	3
	Total	60

Books

1. Quality Assurance of Pharmaceuticals, Vol. 2, Updated Edition, World Health Organization, Geneva.
2. S.H. Willing, Good Manufacturing Practices for Pharmaceuticals; A plan for total Quality control, Latest Edition, Marcel Dekker.
3. Regulatory guidelines related to GMP by
 - a. 21 Code of Federal Regulation, Parts 210, 211&58 (USFDA guidelines)
 - b. EU, MHRA, UK Guidelines on GMP
 - c. Schedule M of Drug & Cosmetics Act.
4. Quality Planning & Analysis by J. M. Juran and F. M. Gryna, Tata Mcgraw Hill, India.
5. Quality Assurance Guide by Organization of Pharmaceutical Producers of India.

Unit	Course Contents (Topics)	Hours
1	General Anatomy and Physiology of skin, hair, nail and tooth:	8
1.1	<i>Self study :Anatomy and physiology of skin, hair, nail and tooth-emphasis on points with reference to cosmetics.</i>	4
1.2	Problems associated with normal functioning of skin, aged skin, dry skin, sensitive skin, acne, pigmentation disorders. Common hair problems - hair loss, manageability problems, split ends, shine and luster disorders; nail problems; tooth problems.	4
2	General raw materials in cosmetic formulations:	19
2.1	Overview of raw materials-Water, natural & synthetic oils, fats& waxes, inorganic solids, emulsifiers, thickeners, hydrocolloids, polymers, surfactants, antioxidants, humectants, polysiloxanes, preservatives.	2
2.2	Colouring agents used in cosmetics. Quality evaluation of colors, safety, toxicity and regulatory aspects of colors w.r.t. cosmetic products	3
2.3	Perfumes in cosmetics: raw materials in perfumery, developing a perfume composition, current trends including emulsified and solid perfumery, analytical and separation techniques of perfumes, sensory analysis, safety and toxicological evaluation of perfumes, manufacturing and packaging of perfumes, legislation and regulations for perfumes in cosmetics.	6
2.4	Therapeutic ingredients in various cosmetics like skin products, dentifrices, hair care and nail preparations, and performance evaluation of these activities.	3
2.3	<i>Self study: Details of general raw materials (oils, fats, waxes, surfactants, preservatives, polysiloxanes), Historical purview of perfumes, Approved colours as per Indian, European and US specifications</i>	5
3	Application of novel approaches in cosmetic formulations	4
3.1	Concepts of microemulsions, liposomes, niosomes, nanoparticles, iontophoresis, to enhance functional attributes & delivery of cosmeceuticals.	4
4	Herbal cosmetics	6
4.1	Current trends in use of herbal materials in cosmetics.	2
	<i>Self study: Discussion on aloe vera, henna, tea tree oil, neem in various cosmetic products</i>	4
5	Packaging and labelling of cosmetic products	5
5.1	Packaging materials, speciality packages for cosmetics, labelling requirements for cosmetics	5
6	Quality standards of cosmetic products:	18
6.1	BIS guidelines for quality of finished products for cosmetics, quality control, textural analysis, performance and psychometric evaluation of various cosmetic products such as creams, gels, powders, lipstick, nail lacquer, shampoo, sunscreen products, dentifrices.	8
6.2	Microbiological quality of cosmetic products.	2
6.3	Safety and toxicity evaluation of cosmetic products	4

6.4	Legal considerations and regulatory procedures of cosmetic products	2
6.5	<i>Self study- BIS, European and US specifications about quality standards of cosmetic products</i>	2
	Total	60

Books:

1. Harry's Cosmeticology Edited by J.B. Wilkinson and R. J. Moore, Longman Scientific & Technical Publishers
2. Cosmetics Science and Technology, Edited by M.S. Balsam, E. Sagarin, S.D. Gerhon, S.J.Strianse and M.M.Rieger, Volumes 1,2 and 3.Wiley-Interscience, Wiley India Pvt. Ltd.,2008
3. Poucher's Perfumes, cosmetics & Soaps, 10th Ed, Editor- Hilda Butler, Kluwer Academic Publishers, Netherlands, 2000
4. Cosmetic Technology, Ed. By S.Nanda, A. Nanda and R. Khar, Birla Publications Pvt. Ltd., New Delhi, 2007
5. Handbook of Cosmetic Science and Technology, edited by M. Paye, A.O.Barel, H. I. Maibach, Informa Healthcare USA,Inc. 2007.
6. Encyclopedia of Pharmaceutical Technology, Vol. 6, Eds. James Swarbrick, James C. Boylan, Marcel Dekker Inc., 1992
BIS Guidelines for different cosmetic products
7. Drugs & Cosmetics Act & Rules, 1940 (with latest amendments).

Elective 11

Polymers in Pharmacy

4 hrs/week

Unit	Course Contents (Topics)	Hrs.
1.0	Historical Background, Basic definitions, Applications	1
2.0	Classification of Polymers	7
2.1	Classification based on reaction to temperature and structure/arrangement/architecture - linear, branched, crosslinked.	2
2.2	Polymerization mechanisms- Addition & step-growth polymerization-Free radical, cationic, anionic and Ziegler Natta mechanisms	5
3.0	Copolymerization	5
3.1	Theoretical aspects of copolymerization	3
3.2	<i>Self study- Case studies of any two copolymers</i>	2
4.0	Properties & Characterization of polymers	12
4.1	Factors affecting and Overview	1
4.2	Molecular weight and determination of molecular weight,	2
4.3	Solid state characterization- glass transition temperature, Crystallinity,	3
4.4	Solubility of polymers & Swelling pr, Mechanical properties	3
4.5	<i>Self study- Case study-any 3 polymers – characteristics & comparison</i>	3
5.0	Methods of Preparation of Polymers	11

5.1	Bulk polymerization, Solution polymerization,	3
5.2	Suspension polymerization, Emulsion polymerization.	3
5.3	Additives in polymers, Fabrication of polymeric devices/systems- casting, extrusion, moulding etc	3
5.4	<i>Self study- One example polymer for each method</i>	2
6.0	Biocompatibility of Polymers	10
6.1	Safety & Biocompatibility issues- Overview	1
6.2	Reaction of polymer to tissues, effect of body/host systems to polymers	2
6.3	Mechanisms of tissue reactions/injury,	2
6.4	Evaluation of biocompatibility of polymers	3
6.5	<i>Self study- Pharmacopoeial & other tests for toxicity evaluation of polymers</i>	2
7.0	Biocompatible Polymers	10
7.1	General features of biocompatible polymers, enzymatically degradable bonds in polymers	2
7.2	Design of biocompatible polymers & evaluation,	4
7.3	<i>Self study- some examples-PLGA, cellulose, acrylates, hydrogels.</i>	4
8.0	Applications of polymers in pharmacy.	4
8.1	Overview of applications as thickeners, binders, coating agents, adhesives, as release modifying agents, including smart polymers, elastomers	2
8.2	<i>Self study : One example each of – adhesive polymer, coating agent, drug release modifier, smart polymer</i>	2
	Total	60

Books:

1. Fundamental Principles of Polymeric Materials by Rosen SL, Wiley- Interscience Publication, 2nd edn, 1993.
2. Martin's Physical Pharmacy and Pharmaceutical Sciences by Sinko PJ, Ed Lea & Feiger, Lippincott Williams & Wilkins, 6th edn, 2010.
3. Controlled Drug Delivery: Fundamentals and Applications, Robinson JR, Lee VHL, Dekker, 2nd edn, Vol 29, 1987.
4. Biodegradable Polymers as Drug Delivery Systems, Chasin M, Langer R, Marcel Dekker, Vol 45, 1990.
5. Controlled and Novel Drug Delivery by Jain NK, CBS Publishers and Distributors, 2008.
6. Controlled Drug Delivery: Clinical Applications, by Bruk SD, CRC Press Inc., Vol 2, 1983.
7. Polymeric Drug Delivery System by Kwon GS, Marcel Dekker, Vol 148, 2005.
8. Aqueous polymeric coating for pharmaceutical dosage forms by McGinity J W , Marcel Dekker, Vol 79.

Unit	Course Contents (Topics)	Hours
1		19
1.1	Basic principles of drug discovery and biological screening <ul style="list-style-type: none"> • Correlation between various animal models and human situations. • Correlation between <i>in vitro</i> and <i>in vivo</i> screens. • Care, handling, breeding techniques of lab animals. • CPCSEA, OECD, ICH guidelines in brief. 	6
1.2	High throughput screening in drug discovery. Techniques for high throughput screening. <ul style="list-style-type: none"> • Cell based assays • Biochemical assays. • Radio ligand binding assays. 	6
1.3	Detection methods <ul style="list-style-type: none"> • Fluorescence based assay techniques • Chemiluminescence based assay techniques. 	3
1.4	Self study Use of alternative methods of screening: <ul style="list-style-type: none"> • <i>Zebrafish model</i> • <i>Drosophila</i> <i>Types of drugs for which these models can be used.</i>	4
	Target based drug discovery and in vitro screening techniques for	26
2.1	Anti-platelet activity- Turbidimetric, GP IIB – IIIA assays using platelet agregometer.	2
2.2	CNS: <ul style="list-style-type: none"> • Alzheimer's disease: <i>in vivo</i> which includes aluminum induced, scopolamine induced memory loss. <i>In vitro</i> includes acetylcholinesterase activity. • Parkinson's disease: <i>in vivo</i> includes Haloperidol, reserpine, rotenone, MPTP induced models. • Anti depressant and anti-convulsants. 	6
2.3	Anti-diabetic: Alloxan, STZ, genetically diabetic animals and various in vitro methods	3
2.4	<ul style="list-style-type: none"> • Anti-tubercular: BACTEC • Anticancer: Few <i>in vitro</i> cell lines, models for metastasis. • Anti-HIV: Various targets involved • Anti malarial. 	6
2.5	Immunomodulatory: <i>in vivo</i> and <i>in vitro</i> methods.	2
2.6	Anti-inflammatory: Acute, subacute and chronic models.	2
2.7	<i>Self study-Antioxidant activity</i>	5
3	<ul style="list-style-type: none"> • Estimation of drugs from complex media like biological fluids Eg. blood, tissues, CSF etc. 	5

	<ul style="list-style-type: none"> • <i>Self study-US FDA guidelines for bio analysis methods including validation.</i> 	2
4	<ul style="list-style-type: none"> • <i>-In vitro skin irritation and eye irritation tests</i> • <i>-In vitro tests for pyrogenicity</i> • <i>Self study-Alternative methods for toxicity testing(in vitro)</i> 	5
	Total	60

Books

1. H.G. Vogel, Drug discovery and evaluation- Pharmacological Assays-Springer Verlag.
2. R.A.Turner, Screening methods in pharmacology- Academic Press.
3. D.R.Laurence and A.L.Bacharach- Evaluation of drug activities: Pharmacometrics. Academic Press.
4. A. Schwartz, Methods in Pharmacology- Plenum Publishing Corporation.
5. Website--Altox.org/ttrc/validation-va

Elective 13

Analytical Method Development and Validation Techniques

4 hrs/week

Unit	Course Contents (Topics)	Hours
1	Calibration & Validation of analytical instruments: <ol style="list-style-type: none"> HPLC. UV-VIS spectrophotometer. FTIR. Dissolution test apparatus. 	4
2	HPLC assay method development for API and drug products:	20
2.1	Preliminary investigations- Nature of sample, its composition and properties. (It should also include significance of pK_a , partition coefficient and current methods to determine the same), separation goals, sample pretreatment and detection, developing separation.	5
2.2	Basics of separation-Resolution, Resolution as a function of- solvent strength, selectivity and column plate number; and sample size effect.	1

2.3	<i>Self study-Detection-Comparison of sensitivity, selectivity, advantages, disadvantages and applications with respect to detectors such as U.V, Fluorescence, PDA, Refractive Index, Evaporative light scattering detector and electrochemical detectors.</i>	2
2.4	Sample preparation and pretreatment for solid, liquid, semisolid samples; column switching and pre and post column derivatisation.	2
2.5	<i>Self study-Columns-characteristics of column and column packing and column specifications.</i>	1
2.6	Method development for Reverse-phase, Ion pair and ion exchange chromatography, Gradient elution-principle and development of gradient separation. <i>Self study- pharmaceutical examples for these methods-(1 hr).</i>	3
2.7	Quantitation analysis- measurement of signals, quantitation methods, sources of errors, procurement, storage and use of reference standards and working standards.	2
2.8	ICH guidelines for analytical method validation (Q2 with latest revision) .System suitability testing as per USP, IP.	3
2.9	<i>Self study-One detailed HPLC analysis of any API by USP or IP (1hr)</i>	1
3	HPTLC : Method development and validation for fixed dose combination drugs and herbal analysis.	3
4	Impurity profiling:	9
4.1	a. <i>Self study-Sources of impurities and ICH terminologies-Organic impurities, Inorganic impurities, Residual solvents, Isolation and characterisation methods for impurities (3 hrs).</i> b. Analytical method development and quantitation of impurities.	5
4.2	ICH guidelines- Q3A, Q3B, Q3C with latest revisions for all.	4
5	Bioanalytical method development and validation:	13
5.1	Steps followed-sample preparation, liquid-liquid extraction, precipitation, solid-phase extraction, sintered column solid phase extraction.	3
5.2	Bioanalytical method validation including full, partial, cross validation, selectivity, accuracy, calibration curve, stability (freeze-thaw and mobile phase), recovery.	4
5.3	CDER and ICH guidelines for bioanalytical method validation.	4
5.4	<i>Self study- Examples of bioanalytical method development and validation for a specified drug estimated in urine/ plasma/serum samples.</i>	2
6	Stability testing:	11
6.1	Drug development cycle and stability testing.	2
6.2	Stress testing of drug substances.	1
6.3	Stability indicating assays (specific and selective), Role of kinetic studies.	3
6.4	Stability testing protocols.	1
6.5	Retest period / Shelf-life determination of drug substances / phytopharmaceuticals / biotechnological products and equipments.	2

6.6	ICH guidelines-Q1A and Q1B with latest revisions.	2
	Total	60

Books

1. Practical HPLC Method Development by L.R.Snyder, 2nd edition, John Wiley & Sons.
2. Analytical Method Validation and Instrument Performance Verification by Chung Chow Chan, Herman Lam, Y.C. Lee, Xue-Ming Zhang, Wiley Interscience, John Wiley & Sons, Incorp Publications.
3. United States Pharmacopoeia and Indian Pharmacopoeia.
4. Handbook of Isolation and Characterisation of Impurities in Pharmaceuticals by Satinder Ahuja & Karen Mills Alsante, Volume 5, Academic Press, USA.
5. Handbook of Bioanalysis & Drug metabolism by Gary Evans, CRC Press (2004),
6. HPLC method development by Satinder Ahuja
7. Sethi's Quantitative Analysis of Pharmaceutical Formulations by P.D.Sethi, fourth edition, volume-1, CBS Publishers and Distributors, New Delhi.
8. Remington-The Science and Practice of Pharmacy, 20th edition Remington-The Science and Practice of Pharmacy, 20th edition.
9. Validation and Qualification in Analytical Laboratories, 2nd edition, Ludwig Huber; Informa Healthcare.
10. Handbook of stability testing in pharmaceutical development - Regulations, Methodologies and Best practices; Editor Kim Huynh-Ba, Springer.
11. J.T. Carstensen, C.T. Rhodes, "Drug stability: principles & Practices". Latest Edition. Marcel Dekker Inc., New York

INTERNET REFERENCES:

1. US FDA (CDER) and ICH guidelines for Bioanalytical method validation.
2. ICH guidelines- Q1A(R), Q3A(R), Q3B, Q3C, Q6A.
3. ICH guidelines for analytical method validation.

Experimental Techniques in Pharmaceutical Sciences 8 hrs/week

Syllabus for the course Experimental Techniques in Pharmaceutical Sciences for the main branches of pharmacy namely Pharmaceutical Chemistry/Medicinal Chemistry, Pharmaceutics, Pharmacology, Pharmaceutical Analysis/Quality Assurance and Pharmacognosy is given below. Students of other branches of specialization like Clinical Pharmacy, Medicinal Natural Products, Biopharmaceutics and Pharmacokinetics, etc. can choose to do the exercises prescribed for any one of the above mentioned branches, preferably selecting something closest to their branch of specialization.

An attempt should be made to cover a **minimum** of 8 exercises in the syllabus of a particular branch

1. Pharmaceutical Chemistry (and Medicinal Chemistry)

Unit	Experiments
1.0	Measurement of logP of a poorly water soluble and a highly water soluble drug
2.0	Determination of the pK_a of a drug (weak acid and weak base) by potentiometric titration and/or by UV/visible spectroscopy
3.0	Measurement of V_{max} and K_m of an hydrolase enzyme (can use esterase, 70hosphatise, or lipase type enzymes Student should learn how to plot data by Linweaver Burke and Eadie Hofstee methods)
4.0	Estimation of two drugs by simultaneous equation method and by absorbance ratio method. (preferably use combinations of drugs that are used as fixed drug combination)
5.0	Synthesis of some drugs (for e.g. thiazide and hydrothiazide derivatives) involving multistep (at least three steps) reactions. (Students should learn to monitor the reaction by TLC, separate the main product from impurities by column chromatography and learn use of IR and 1H and ^{13}C NMR to check the structures of the intermediates and the final compounds and estimate overall yield of the reaction).
6.0	Resolution of racemic mixtures of acidic and basic compounds by formation of diastereomers
7.0	Synthesis of prodrugs of any one of the common drugs and study of their decomposition (kinetics) in plasma or serum to the parent drug (suggest use of DCC based coupling to obtain ester prodrugs)
8.0	Establish a RPHPLC method for the separation of a mixture of two or more compounds (e.g fixed dose combination drugs, or prodrugs synthesized above or apply to reaction monitoring)
9.0	Working with physical models- ball and stick or space-filling models. Students should learn to construct physical models for glucose, vitamin C, propranolol, chloramphenicol. This will enable students to identify stereocenters and assign correct stereochemistry to them.
10	Demonstration of some molecular modelling exercises like energy minimization, molecular dynamics simulations, docking, 2D/3D-QSAR, structure based drug, pharmacophore mapping etc. using either commercially available programs or freeware.

2. Pharmaceutics

1. Study of dissolution profile of IR and ER products. Mathematical treatment of data for release Kinetics and f_1 and f_2 analysis.
2. Simple Optimization design (formulation study/pH-stability study)
3. Design and evaluation of Orally Disintegrating Drug Delivery System
4. Preparation and evaluation of microspheres for inhalation system
5. Preparation and evaluation of transdermal/mucoadhesive/gastroretentive system
6. Constructing phase diagram for one system of oil, surfactant- cosurf, water
7. Design of one vesicular system - niosomes/liposomes systems can be made
8. Design of lipid particulate system (nanosystems with wax can be tried)

3. Pharmacognosy

- 1) Extraction and evaluation of Mucilage from suitable sources such as aloes or Leaves of *Annona squamosa* or any other suitable source.
- 2) Extraction and analysis of alkaloids such as purine alkaloids and Tropane or indole alkaloids from suitable natural sources.
- 3) Preparation and evaluation of extract of anthraquinone glycosides from *Cassia angustifolia/Cassia fistula* or rhubarb or other suitable source
- 4) Extraction and analysis of Volatile oil from suitable sources such as clove or eucalyptus or any other.
- 5) Extraction and detection of inulin from suitable sources such as chicory or Kuth.
- 6) Activity guided fractionation of any herb for its antimicrobial and/or antioxidant activity.
- 7) Preparation and evaluation of any herbal 'churna, dosage form (Situpaladia or Triphala).
- 8) Extraction and analysis of carotenoid derivatives or flavonoid derivatives from suitable natural sources for each.

4. Pharmaceutical Analysis and Quality Assurance.

1. Determination of pK_a by U.V. spectroscopy.
2. Sample preparation for I.R. spectroscopy (solid/liquids) and interpretation of IR bands for important functional groups.
3. Assays for drugs in combination by UV derivative spectroscopy.
4. Structural elucidation workshop: Interpretation of 1H NMR, ^{13}C NMR, IR and Mass spectrometry of simple compounds (maximum 12 carbon atoms).
5. Standard calibration curve by UV spectroscopy at –
 - a. λ max
 - b. λ max + 10 nm
 - c. λ max – 10 nm
6. Determination of response factor by HPLC.
7. Qualitative and Quantitative HPTLC analysis (minimum mixture of 3 compounds).
8. Assay determination by Simultaneous equation, Absorbance ratio and Difference spectroscopy.
9. Determination of Response factor by HPLC analysis of drugs.
10. Preparative TLC analysis.
11. Bioanalysis by HPLC.
12. pH stability evaluation of Aspirin by TLC.
13. Failure investigation/Investigations of 'Out of Specification' report for products and analytical methodology.
14. Qualifications of Instruments/Equipments.
15. Validation of analytical method/procedure/process.
16. Separation of components by column chromatography.
17. Calibration of UV spectrophotometer / HPLC column

5. Pharmacology

- I) Experiments to be performed by students
 - 1) Assay of antagonist using a suitable isolated tissue preparation
 - 2) Determination of pA_2 values of antagonists using a suitable tissue preparation
 - 3) In vitro antioxidant screening assays (any two) e.g. DPPH, NO, superoxide.
- II) Demonstrations
 - a. Experiments on intact animals to evaluate:
 - 1) Anti cataleptic activity
 - 2) Anti anxiety activity
 - 3) Anti inflammatory/analgesic activity
 - 4) Muscle relaxant activity
 - b. Techniques of drug administration and blood withdrawal
 - c. Non invasive methods of measuring blood pressure, pulse, ECG etc.

III) Tutorials

- 1) Care and handling of animals
- 2) CPCSEA, OECD, ICH guidelines in brief
- 3) Use of alternative methods of screening (Types of drugs for which these models can be used)
 - Zebra fish
 - Drosophila
- 4) Techniques for high throughput screening
 - Cell based assays
 - Biochemical assays
 - Radioligand binding assays

References:

Latest editions of the following books/CDs can be referred.

1. Expharm Pro-Simulated animal experiments in Pharmacology, Elseviers
2. Expharm-Stimulated animal experiments, Ravindran
3. H. G. Vogel, Drug discovery and evaluation-Pharmacological Assays-Springer Verlag
4. M. N. Ghosh, Fundamentals of Experimental Pharmacology, Scientific Book Agency
5. S. K. Kulkarni, Practical Pharmacology and Clinical Pharmacy, Vallabh Publications.
6. CPCSEA, OECD, ICH Guidelines.

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