

#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

#### SEMESTER - I

S.	Course	Course Name	Hour	s per we	eek	Credits
No.	codes		L	T	P	
1.	21S01101	Modern Pharmaceutical Analytical Techniques	4	-	-	4
2.	21S07101	Advanced Pharmaceutical Analysis	4	-	-	4
3.	21S07102	Pharmaceutical and Food Analysis	4	-	_	4
4.	21S07103	Quality Control And Quality Assurance	4	-	-	4
5.	21S01105	Modern Pharmaceutical Analytical Techniques Lab	-	-	6	3
6.	21S07104	Pharmaceutical and Food Analysis Lab	-	-	6	3
7.	21DAC101a 21DAC101b 21DAC101c	Audit Course – I English for Research paper writing Disaster Management Sanskrit for Technical Knowledge	2	-	-	0
8.		Seminar/Assignment	-	1	6	4
		Total	18	1	18	26

#### SEMESTER – II

S.No.	Course	Course Name	Hou	rs per	week	Credits
	codes		L	T	P	
1.	21S07201	Advanced Instrumental Analysis	4	-	-	4
2.	21S07202	Modern Bio-Analytical Techniques	4	-	-	4
3.	21SOE301a	Pharmaceutical Validation	4	-	-	4
4.	21S07203	Herbal and Cosmetic Analysis	4	-	-	4
5.	21S07204	Advanced Instrumental Analysis Lab	-	-	6	3
6.	21S07205	Modern Bio-Analytical Techniques Lab	-	-	6	3
7.	21DAC201a 21DAC201b	Audit Course – II Pedagogy Studies Stress Management for Yoga Personality Development through Life Enlightenment Skills	2	1	ı	0
8.	21S07206	Seminar/Assignment	-	1	6	4
		Total	18	1	18	26



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### COURSE STRUCTURE & SYLLABI SEMSTER - III

S.No.	Course	Course Name	Hours	Hours per week		Credits
	codes		L	T	P	
1.	21DRM101	Research Methodology and Intellectual Property Right	4	-	-	4
2.	21SOE301f	Open Electives Biological Screening methods Stability of Drugs and Dosage forms Pharmacoepidemiology and Pharmacoeconomics	3	1	1	3
3.	21S07301	Teaching Practice/Assignment	-	-	4	2
4.	21S07302	Comprehensive viva voce	-	-	4	2
5.	21S07303	Research Work - I	-		24	12
		Total	7	-	32	23

#### **SEMESTER - IV**

S.No.	Course	Course Name	Hours	Hours per week		Credits
	codes		L	T	P	
1.	21S07401	Co-Curricular Activities	2			2
2.	21S07402	Research Work - II	3		30	18
		Total	5		30	20



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Course Code	MODERN PHARMACEUTICAL ANALYTICAL	L	T	P	C
<b>21S01101</b>	TECHNIQUES	4	0	0	4
	Semester	ster		[	

#### **Course Objectives:**

The course is designed to impart the knowledge in the field of Pharmaceutical Analysis. The various modern analytical techniques like UV-Visible, IR, NMR, Mass, GC, HPLC, different chromatographic methods and other important topics are taught to enable the students to understand and apply the principles involved in the determination of different bulk drugs and their formulation. In addition to the theoretical aspects, the basic practical knowledge relevant to the analysis is also imparted.

#### Course Outcomes (CO): Student will be able to

- Modern Analytical Techniques and can apply the theories in analysis of various drugs in single and combination dosage forms
- Theoretical and practical skills of the instruments
- Apply their knowledge in developing the new methods for the determination and validate the procedures.

#### UNIT - I

#### UV-Visible spectroscopy

Introduction, Theory, Laws, and Instrumentation associated with UV-Visible spectroscopy, Choice of solvents and solvent effect and Applications of UV-Visible spectroscopy, Difference/ Derivative spectroscopy.

#### UNIT - II

#### IR spectroscopy

Theory, Modes of Molecular vibrations, Sample handling, Instrumentation of Dispersive and Fourier -Transform IR Spectrometer, Factors affecting vibrational frequencies and Applications of IR spectroscopy, Data Interpretation.

#### UNIT - III

#### NMR spectroscopy

Quantum numbers and their role in NMR, Principle, Instrumentation, Solvent requirement in NMR, Relaxation process, NMR signals in various compounds, Chemical shift, Factors influencing chemical shift, Spin-Spin coupling, Coupling constant, Nuclear magnetic double resonance, Brief outline of principles of FT-NMR and 13C NMR. Applications of NMR spectroscopy

#### UNIT - IV

#### **Mass Spectroscopy**

Principle, Theory, Instrumentation of Mass Spectroscopy, Different types of ionization like electron impact, chemical, field, FAB and MALDI, APCI, ESI, APPI Analyzers of Quadrupole and Time of Flight, Mass fragmentation and its rules, Meta stable ions, Isotopic peaks and Applications of Mass spectroscopy.

#### UNIT - V

#### Chromatography

Introduction to chromatography and classification of chromatographic methods based on the mechanism of separation, Principle, instrumentation, selection of solvents; chromatographic parameters, factors affecting resolution, applications of the following:

- a) Thin Layer chromatography;
- b) High Performance Thin Layer Chromatography
- c) Paper Chromatography;
- d) Column chromatography

e) Gas chromatography;

f) High Performance Liquid chromatography



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g) Affinity chromatography;

h) Gel Chromatography

i)Hyphenated techniques:

- Ultra High Performance Liquid chromatography- Mass spectroscopy
- Gas Chromatography-Mass Spectroscopy

#### **Textbooks:**

- 1. Instrumental Methods of Chemical Analysis by B.K Sharma
- 2. Vogel's Text book of Quantitative Chemical Analysis by A.I. Vogel
- 3. Textbook of Pharmaceutical Analysis, KA.Connors, 3rd Edition, John Wiley& Sons, 1982.

- 4. Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John Wiley & Sons, 2004.
- 5. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, T imothy A. Nieman, 5th edition, Eastern press, Bangalore, 1998.
- 6. Instrumental methods of analysis Willards, 7th edition, CBS publishers.
- 7. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4<sup>th</sup>edition, CBS Publishers, New Delhi, 1997.
- 8. Organic Spectroscopy William Kemp, 3rd edition, ELBS, 1991.
- 9. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi,3rd Edition, CBS Publishers, New Delhi, 1997.
- 10. Pharmaceutical Analysis Modern Methods Part B J W Munson, Vol11, Marcel. Dekker Series
- 11. Spectroscopy of Organic Compounds, 2nd edn., P.S/Kalsi, Wiley esternLtd., Delhi.
- 12. Organic Chemistry by I. L. Finar
- 13. Quantitative Analysis of Drugs by D. C. Garrett
- 14. HPTLC by P.D. Seth
- 15. Indian Pharmacopoeia 2007
- 16. High Performance thin layer chromatography for the analysis of medicinal plants by Eike
- 17. Reich, Anne Schibli
- 18. Introduction to instrumental analysis by Robert. D. Braun



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Course Code	ADVANCED PHARMACEUTICAL ANALYSIS	L	T	P	C
21S07101		4	0	0	4
	Semester	I			
			<u> </u>		

#### **Course Objectives:**

This subject deals with the various aspects of Impurity, Impurities in new drug products, in residual solvents, Elemental impurities, Impurity profiling and characterization of degradents, Stability testing of phytopharmaceuticals and their protocol preparation. It also covers the biological testing of various vaccines and their principle and procedure.

#### **Course Outcomes (CO):** Student will be able to

- Appropriate analytical skills required for the analytical method development.
- Principles of various reagents used in functional group analysis that renders necessary support in research methodology and demonstrates its application in the practical related problems.
- Analysis of impurities in drugs, residual solvents and stability studies of drugs and biological products

## UNIT - I Impurity and stability studies

Definition, classification of impurities in drug Substance or Active Pharmaceutical Ingredients and quantification of impurities as per ICH guidelines Impurities in new drug products: Rationale for the reporting and control of degradation products, reporting degradation products content of batches, listing of degradation products in specifications, qualification of degradation products

Impurities in residual solvents: General principles, classification of residual solvents, Analytical procedures, limits of residual solvents, reporting levels of residual solvents

#### UNIT - II

#### **Elemental impurities**

Element classification, control of elemental impurities, Potential Sources of elemental Impurities, Identification of Potential Elemental Impurities, analytical procedures, instrumentation & C,H, N and S analysis

#### Stability testing protocols

Selection of batches, container orientation, test parameters, sampling frequency, specification, storage conditions, recording of results, concept of stability, commitment etc. Important mechanistic and stability related information provided by results of study of factors like temperature, pH, buffering species ionic strength and dielectric constant etc. on the reaction rates. With practical considerations.

#### UNIT – III

#### Impurity profiling and degradent characterization

Method development, Stability studies and concepts of validation accelerated stability testing & shelf life calculation, WHO and ICH stability testing guidelines, Stability zones, steps in development, practical considerations. Basics of impurity profiling and degradent characterization with special emphasis. Photo stability testing guidelines, ICH stability guidelines for biological products

#### UNIT – IV

#### Stability testing of phytopharmaceuticals

Regulatory requirements, protocols, HPTLC/HPLC finger printing, interactions and complexity.

#### Biological tests and assays of the following

Adsorbed Tetanus vaccine b. Adsorbed Diphtheria vaccine c. Human anti haemophilic vaccine d. Rabies vaccine e. Tetanus Anti toxin f. Tetanus Anti serum g. Oxytocin h. Heparin sodium IP i. Antivenom. PCR, PCR studies for gene regulation, instrumentation (Principle and Procedures)



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#### **COURSE STRUCTURE & SYLLABI**

UNIT – '	V
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#### Immunoassays (IA)

Basic principles, Production of antibodies, Separation of bound and unbound drug, Radioimmunoassay, Optical IA, Enzyme IA, Fluoro IA, Luminiscence IA, Quantification and applications of IA.

- 1. Vogel's textbook of quantitative chemical analysis Jeffery J Bassett, J.Mendham, R. C. Denney, 5th edition, ELBS, 1991.
- 2. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4<sup>th</sup>Edition, CBS publishers, New Delhi, 1997.
- 3. Textbook of Pharmaceutical Analysis K A Connors, 3rd Edition, John Wiley& Sons, 1982.102.
- 4. Pharmaceutical Analysis Higuchi, Brochmman and Hassen, 2nd Edition, Wiley Inter science Publication, 1961.
- 5. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi,3rd Edition, CBS Publishers New Delhi, 1997.
- 6. Pharmaceutical Analysis- Modern methods J W Munson Part B, Volume 11, Marcel Dekker Series.
- 7. The Quantitative analysis of Drugs D C Carratt, 3rd edition, CBS Publishers, New Delhi, 1964.
- 8. Indian Pharmacopoeia VolI, II & III 2007, 2010, 2014.
- 9. Methods of sampling and microbiological examination of water, first revision, BIS
- 10. Practical HPLC method development Snyder, Kirkland, Glajch, 2<sup>nd</sup>edition, John Wiley & Sons.
- 11. Analytical Profiles of drug substances Klaus Florey, Volume 1 20, Elsevier, 2005
- 12. Analytical Profiles of drug substances and Excipients Harry G Brittan, Volume 21 30, Elsevier, 2005.
- 13. The analysis of drugs in biological fluids Joseph Chamberlain, 2<sup>nd</sup>edition, CRC press, London.
- 14. ICH Guidelines for impurity profiles and stability studies.



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Course Code	PHARMACEUTICAL AND FOOD ANALYSIS	L	T	P	C
21S07102		4	0	0	4
·	Semester			Ī	
Course Objectives:					
This course is designe	ed to impart knowledge on analysis of food constituents a	nd f	inish	ed fo	ood
products. The course in	cludes application of instrumental analysis in the determinati	on			
of pesticides in variety	11				

#### **Course Outcomes (CO):** Student will be able to

various analytical techniques in the determination of

- Food constituents
- Food additives
- Finished food products
- Pesticides in food
- Pharmaceuticals (API & Dosage forms)
- And also student shall have the knowledge on food regulations and legislations

#### UNIT - I

#### Carbohydrates

Classification and properties of food carbohydrates, General methods of analysis of food carbohydrates.

#### **Proteins**

Chemistry and classification of amino acids and proteins, Physico-Chemical properties of protein and their structure, general methods of analysis of proteins and amino acids

#### UNIT - II

#### Lipids

Classification, general methods of analysis, refining of fats and oils; hydrogenation of vegetable oils, Determination of adulteration in fats and oils.

#### **Vitamins**

Classification of vitamins, methods of analysis of vitamins, Principles of microbial assay of vitamins of B-series

#### UNIT – III

#### **Probiotics**

Definition, history, importance, mode of action, identification advantages and disadvantages of probiotics. Applications of Probiotics

#### UNIT - IV

#### Food additives

Introduction, analysis of Preservatives, antioxidants, artificial sweeteners, flavors, flavor enhancers, stabilizers, thickening and jelling agents.

#### Pigments and synthetic dyes

Natural pigments, their occurrence and characteristic properties, permitted synthetic dyes, Non-permitted synthetic dyes used by industries, Method of detection of natural, permitted and non-permitted dyes.

#### UNIT – V

#### Milk (constituents and milk products)

General Analytical methods for milk, milk constituents and milk products like ice cream, milk powder, butter, margarine, cheese including adulterants and contaminants of milk.

• Analysis of fermentation products like wine, spirits, beer and vinegar.



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- Pesticides Analysis in food like organophosphorus and organochlorine
- And also student shall have knowledge in food regulations and legislations

#### **Textbooks:**

- 1. The chemical analysis of foods David Pearson, Seventh edition, Churchill Livingstone, Edinburgh London, 1976
- 2. Introduction to the Chemical analysis of foods S. Nielsen, Jones & Bartlett publishers, Boston London, 1994.
- 3. Official methods of analysis of AOAC International, sixth edition, Volume I & II, 1997.
- 4. Analysis of Food constituents Multon, Wiley VCH.
- 5. Dr. William Horwitz, Official methods of analysis of AOAC International
- 6. 18th edition, 2005. Theory and Practice of Industrial Pharmacy by Lieberman and Lachman

- 1. Indian Pharmacopoeia 2012
- 2. Remington's Pharmaceutical Sciences by Alfonso and Gennaro



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#### COURSE STRUCTURE & SYLLABI

<b>Course Code</b>	QUALITY CONTROL AND QUALITY	L	T	P	C				
21S07103	ASSURANCE	4	0	0	4				
	Semester I								
<b>Course Objectives:</b>									
	with the various aspects of quality control and quality assu stries. It covers the important aspects like cGMP, QC tests								

quality certifications, GLP and regulatory affairs. Course Outcomes (CO): Student will be able to

- The cGMP aspects in a pharmaceutical industry
- To appreciate the importance of documentation
- To understand the scope of quality certifications applicable to Pharmaceutical industries
- To understand the responsibilities of QA & QC departments

#### UNIT - I

#### **Quality Control and Quality Assurance**

Concept and Evolution of Quality Control and Quality Assurance Good Laboratory Practice, GMP, Overview of ICH Guidelines -QSEM, with special emphasis on Q-series guidelines.

#### **Good Laboratory Practices**

Scope of GLP, Definitions, Quality assurance unit, protocol for conduct of non clinical testing, control on animal house, report preparation and documentation.

#### UNIT - II

#### cGMP

cGMP guidelines according to schedule M, USFDA (inclusive of CDER and CBER) Pharmaceutical Inspection Convention(PIC), WHO and EMEA covering: Organization and personnel responsibilities, training, hygiene and personal records, drug industry location, design, construction and plant lay out, maintenance, sanitation, environmental control, utilities and maintenance of sterile areas, control of contamination and Good Warehousing Practice. CPCSEA guidelines.

#### UNIT - III

Analysis of raw materials, finished products, packaging materials, in process quality control (IPQC), Developing specification (ICH Q6 and Q3) Purchase specifications and maintenance of stores for raw materials. In process quality control and finished products quality control for following formulation in Pharma industry according to Indian, US and British pharmacopoeias: tablets, capsules, ointments, suppositories, creams, parenterals, ophthalmic and surgical products (How to refer pharmacopoeias), Quality control test for containers, closures and secondary packing materials.

### UNIT – IV Documentation in pharmaceutical industry

Three tier documentation, Policy, Procedures and Work instructions, and records (Formats), Basic principles- How to maintain, retention and retrieval etc. Standard operating procedures (How to write), Master Formula Record, Batch Formula Record, Quality audit plan and reports. Specification and test procedures, Protocols and reports. Distribution records. Electronic data.

#### UNIT - V

#### **Manufacturing operations and controls:**

Sanitation of manufacturing premises, mix-ups and cross contamination, processing of intermediates and bulk products, packaging operations, IPQC, release of finished product, process deviations, charge-in of components, time limitations on production, drug product inspection, expiry date calculation, calculation of yields, production record review, change control, sterile products, aseptic process control, packaging.



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#### **COURSE STRUCTURE & SYLLABI**

- 1. Quality Assurance Guide by organization of Pharmaceutical Procedures of India, 3rd revised edition, Volume I & II, Mumbai, 1996.
- 2. Good Laboratory Practice Regulations, 2nd Edition, Sandy Weinberg Vol.69, Marcel Dekker Series, 1995.
- 3. Quality Assurance of Pharmaceuticals- A compedium of Guide lines and Related materials Vol I & II, 2nd edition, WHO Publications, 1999.
- 4. How to Practice GMP's P P Sharma, Vandana Publications, Agra, 1991.
- 5. The International Pharmacopoeia vol I, II, III, IV & V General Methods of Analysis and Quality specification for Pharmaceutical Substances, Excepients and Dosage forms, 3rd edition, WHO, Geneva, 2005.
- 6. Good laboratory Practice Regulations Allen F. Hirsch, Volume 38, Marcel Dekker Series, 1989.
- 7. ICH guidelines
- 8. ISO 9000 and total quality management
- 9. The drugs and cosmetics act 1940 Deshpande, Nilesh Gandhi, 4<sup>th</sup>edition, Susmit Publishers, 2006.
- 10. QA Manual D.H. Shah, 1st edition, Business Horizons, 2000.
- 11. Good Manufacturing Practices for Pharmaceuticals a plan for total quality control Sidney H. Willig, Vol. 52, 3rd edition, Marcel Dekker Series.
- 12. Steinborn L. GMP/ISO Quality Audit Manual for Healthcare Manufacturers and Their Suppliers, Sixth Edition, (Volume 1 With Checklists and Software Package). Taylor & Francis; 2003.
- 13. Sarker DK. Quality Systems and Controls for Pharmaceuticals. John Wiley& Sons; 2008.



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21801105	TECHNIQUES LAB	0	0	6	3
	Semester	Ī			

#### List of Experiments

- 1. Analysis of Pharmacopoeial compounds and their formulations by UV Vis Spectrophotometer.
- 2. Simultaneous estimation of multi component containing formulations by UV Spectrophotometry
- 3. Effect of pH and solvent on UV –Spectrum
- 4. Determination of Molar absorption coefficient
- 5. Estimation of riboflavin/ quinine sulphate by fluorimetry
- 6. Study of quenching effect by fluorimetry
- 7. Estimation of sodium or potassium by flame photometry
- 8. Colorimetric determination of drugs by using different reagents
- 9. Qunatitative determination of functional groups
- 10. Experiments based on Column chromatography
- 11. Experiments based on HPLC
- 12. Experiments based on Gas Chromatography
- 13. Preparation of Master Formula Record.
- 14. Preparation of Batch Manufacturing Record.



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Course Code	PHARMACEUTICAL AND FOOD ANALYSIS LAB	L	T	P	C
21S07104		0	0	6	3
	Semester		]	[	

#### **List of Experiments**

- 1. Determination of total reducing sugar
- 2. Determination of proteins
- 3. Determination of saponification value, Iodine value, Peroxide value, Acid value in food products
- 4. Determination of fat content and rancidity in food products
- 5. Analysis of natural and synthetic colors in food
- 6. Determination of preservatives in food
- 7. Determination of pesticide residue in food products
- 8. Analysis of vitamin content in food products
- 9. Determination of density and specific gravity of foods
- 10. Determination of benzoic acid by titrimetric analysis in beverages/ sauces/ ketchup/ jam
- 11. Assay of any two Analgesic & Antipyretic drugs (API & dosage forms) official in IP
- 12. Assay of any two Antihistamines (API & dosage forms) official in IP
- 13. Assay of any two Diuretics (API & dosage forms) official in IP



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Course Code	ADVANCED INSTRUMENTAL ANALYSIS		L	T	P	C
21S07201			4	0	0	4
Pre-requisite		Semester		I	I	

#### **Course Objectives:**

This subject deals with various hyphenated analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are LC-MS, GC-MS, and hyphenated techniques.

#### Course Outcomes (CO): Student will be able to

- Interpretation of the NMR, Mass and IR spectra of various organic compounds
- Theoretical and practical skills of the hyphenated instruments
- · Identification of organic compounds

#### UNIT - I

#### **HPLC**

Principle, instrumentation, pharmaceutical applications, peak shapes, capacity factor, selectivity, plate number, plate height, resolution, band broadening, pumps, injector, detectors, columns, column problems, gradient HPLC, HPLC solvents, trouble shooting, sample preparation, method development, New developments in HPLC-role and principles of ultra, nano liquid chromatography in pharmaceutical analysis. Immobilized polysaccharide CSP's: Advancement in enantiomeric separations, revised phase Chiral method development and HILIC approaches. HPLC in Chiral analysis of pharmaceuticals. Preparative HPLC, practical aspects of preparative HPLC.

#### UNIT - II

#### **Biochromatography**

Size exclusion chromatography, ion exchange chromatography, ion pair chromatography, affinity chromatography general principles, stationary phases and mobile phases.

Gas chromatography: Principles, instrumentation, derivatization, head space sampling, columns for GC, detectors, quantification.

#### High performance Thin Layer chromatography

Principles, instrumentation, pharmaceutical applications.

#### UNIT – III

### **Super critical fluid chromatography:** Principles, instrumentation, pharmaceutical applications **Capillary electrophoresis:**

Overview of CE in pharmaceutical analysis, basic configuration, CE characteristics, principles of CE, methods and modes of CE. General considerations and method development in CE, Crown ethers as buffer additives in capillary electrophoresis. CE-MS hyphenation.

#### UNIT - IV

#### Mass spectrometry

Principle, theory, instrumentation of mass spectrometry, different types of ionization like electron impact, chemical, field, FAB and MALD, APCI, ESI, APPI mass fragmentation and its rules, meta stable ions, isotopic peaks and applications of mass spectrometry. LC-MS hyphenation and DART MS analysis. Mass analysers (Quadrpole, Time of flight, FT-ICR, ion trap and Orbitrap) instruments. MS/MS systems (Tandem: QqQ, TOF-TOF; Q-IT, Q-TOF, LTQ-FT, LTQ-Orbitrap.

#### UNIT – V

#### **NMR** spectroscopy

Quantum numbers and their role in NMR, Principle, Instrumentation, Solvent requirement in NMR, Relaxation process, NMR signals in various compounds, Chemical shift, Factors influencing chemical shift, Spin-Spin coupling, Coupling constant, Nuclear magnetic double resonance, Brief



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outline of principles of FT-NMR with reference to 13CNMR:Spin spin and spin lattice relaxation phenomenon. 13C NMR, 1-Dand 2-D NMR, NOESY and COSY techniques, Interpretation and Applications of NMR spectroscopy. LC-NMR hyphenations.

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



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Course Code	MODERN BIO-ANALYTICAL TECHNIQUES	L	T	P	C			
21S07202		4	0	0	4			
<u>.</u>	Semester							
biological matrices.	ned to provide detailed knowledge about the importance of an	alysi	s of	drugs	s in			
Course Outcomes (C	<b>CO):</b> Student will be able to							
• Extraction of drug	gs from biological samples							

- Separation of drugs from biological samples using different techniques
- Guidelines for BA/BE studies.

#### UNIT – I

#### Extraction of drugs and metabolites from biological matrices

General need, principle and procedure involved in the Bioanalytical methods such as Protein precipitation, Liquid -Liquid extraction and Solid phase extraction and other novel sample preparation approach.

Bioanalytical method validation: USFDA and EMEA guidelines

#### UNIT – II

#### **Biopharmaceutical Consideration**

Introduction, Biopharmaceutical Factors Affecting Drug Bioavailability, In Vitro: Dissolution and Drug Release Testing, Alternative Methods of Dissolution Testing Transport models, Biopharmaceutics Classification System. Solubility: Experimental methods. Permeability: In-vitro, in-situ and In-vivo methods.

#### UNIT – III

#### Pharmacokinetics and Toxicokinetics:

Basic consideration, Drug interaction (PK-PD interactions), The effect of protein-binding interactions, The effect of tissue-binding interactions, Cytochrome P450-based drug interactions, Drug interactions linked to transporters. Microsomal assays Toxicokinetics-Toxicokinetic evaluation in preclinical studies, Importance and applications of toxicokinetic studies. LC-MS in bioactivity screening and proteomics

#### UNIT - IV

#### Cell culture techniques

Cell culture techniques Basic equipment's used in cell culture lab. Cell culture media, various types of cell culture, general procedure for cell cultures; isolation of cells, subculture, cryopreservation, characterization of cells and their applications. Principles and applications of cell viability assays (MTT assays), Principles and applications of flow cytometry.

#### UNIT - V

#### Metabolite identification

In-vitro / in-vivo approaches, protocols and sample preparation. Microsomal approaches (Rat liver microsomes (RLM) and Human liver microsomes (HLM) in Met –ID. Regulatory perspectives. In-vitro assay of drug metabolites & drug metabolizing enzymes.

Drug Product Performance, In Vivo: Bioavailability and Bioequivalence: Drug Product Performance, Purpose of Bioavailability Studies, Relative and Absolute Availability. Methods for Assessing Bioavailability, Bioequivalence Studies, Design and Evaluation of Bioequivalence Studies, Study Designs, Crossover Study Designs, Generic Biologics (Biosimilar Drug Products), Clinical Significance of Bioequivalence Studies.



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- 1. Analysis of drugs in Biological fluids Joseph Chamberlain, 2nd Edition.CRC Press, Newyork. 1995.
- 2. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5th edition, Eastern press, Bangalore, 1998.
- 3. Pharmaceutical Analysis Higuchi, Brochmman and Hassen, 2nd Edition, Wiley Interscience Publications, 1961.
- 4. Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11, Marcel Dekker Series
- 5. Practical HPLC method Development Snyder, Kirkland, Glaich, 2<sup>nd</sup>Edition, John Wiley & Sons, New Jercy. USA.
- 6. Chromatographic Analysis of Pharmaceuticals John A Adamovics, 2<sup>nd</sup>Edition, Marcel Dekker, Newyork, USA. 1997.
- 7. Chromatographic methods in clinical chemistry & Toxicology Roger LBertholf, Ruth E Winecker, John Wiley & Sons, New Jercy, USA. 2007.
- 8. Good Laboratory Practice Regulations, 2nd Edition, Sandy Weinberg Vol.69, Marcel Dekker Series, 1995.
- 9. Good laboratory Practice Regulations Allen F. Hirsch, Volume 38, Marcel Dekker Series, 1989.
- 10. ICH, USFDA & CDSCO Guidelines.
- 11. Palmer



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#### **COURSE STRUCTURE & SYLLABI**

Course Code	PHARMACEUTICAL VALIDATION	L	T	P	C
21SOE301a		4	0	0	4
	Semester		I	I	
<b>Course Objectives:</b>					
* *	of the subject is to understand about validation and how it can				
•	mprove the quality of the products. The subject covers the comp	olete	info	rmat	ion
	es, methodology and application				
Course Outcomes (	CO): Student will be able to				
• Explain the aspec	ct of validation				
• Carryout validati	on of manufacturing processes				
<ul> <li>Apply the knowl</li> </ul>	edge of validation to instruments and equipments				
* * *	ufacturing facilities				
UNIT – I	diactaring racinities				
	' CO 1'C' ' 1X71'1' A1 ' CX71'1'	G.	1		- (
	ion of Qualification and Validation, Advantage of Validation,	Str	eami	ınıng	OI
	lation process and Validation Master Plan.			-	
	Requirement Specification, Design Qualification, Factory				
	otance Test (SAT), Installation Qualification, Operationa				
	fication, Re- Qualification (Maintaining status-Calibrat				
	e management), Qualification of Manufacturing Equipments,	Qua	lifica	ation	of
	ts and Laboratory equipments.				
UNIT – II					
Qualification of ana					
	pH meter, UV-Visible spectrophotometer, FTIR, GC,				LC
Qualification of Glas	XI - 1 4 Cl - 1 1 1 1	- d h	urott.	_	
	sware: Volumetric flask, pipette, Measuring cylinder, beakers a	na b	uren	ᡛ.	
UNIT – III		na o	uren	е.	
		na o	uren	<del>с.</del>	
UNIT – III Validation of Utility					ing
UNIT – III Validation of Utility Pharmaceutical Water	y systems	trog	en. C	Clean	
Validation of Utility Pharmaceutical Water Validation: Cleaning	v systems er System &pure steam, HVAC system, Compressed air and ni	trog	en. C	Clean ation	of
Validation of Utility Pharmaceutical Water Validation: Cleaning	systems er System &pure steam, HVAC system, Compressed air and ni g Validation - Cleaning Method development, Validation a	trog	en. C	Clean ation	of
UNIT – III  Validation of Utility Pharmaceutical Wate Validation: Cleaning analytical method us	systems er System &pure steam, HVAC system, Compressed air and ni g Validation - Cleaning Method development, Validation a	trog	en. C	Clean ation	of
Validation of Utility Pharmaceutical Wate Validation: Cleaning analytical method us (CIP).	r systems er System &pure steam, HVAC system, Compressed air and ni g Validation - Cleaning Method development, Validation and ed in cleaning. Cleaning of Equipment, Cleaning of Facilities.	trog	en. C	Clean ation	of
Validation of Utility Pharmaceutical Wate Validation: Cleaning analytical method us (CIP). UNIT – IV Analytical method v	r systems er System &pure steam, HVAC system, Compressed air and ni g Validation - Cleaning Method development, Validation and ed in cleaning. Cleaning of Equipment, Cleaning of Facilities.	trog	en. C	Clean ation	of

#### **General Principles of Intellectual Property**

GAMP.
UNIT – V

Concepts of Intellectual Property (IP), Intellectual Property Protection (IPP), Intellectual Property Rights (IPR); Economic importance, mechanism for protection of Intellectual Property –patents, Copyright, Trademark; Factors affecting choice of IP protection; Penalties for violation; Role of IP in pharmaceutical industry; Global ramification and financial implications. Filing a patent applications; patent application forms and guidelines. Types patent applications-provisional and non-provisional, PCT and convention patent applications; International patenting requirement procedures and costs; Rights and responsibilities of a patentee; Practical aspects regarding maintaining of a Patent file; Patent infringement meaning and scope. Significance of transfer technology (TOT), IP and ethics-



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

#### COURSE STRUCTURE & SYLLABI

positive and negative aspects of IPP; Societal responsibility, avoiding unethical practices.

- 1. B. T. Loftus & R. A. Nash, "Pharmaceutical Process Validation", Drugs and Pharm Sci. Series, Vol. 129, 3rd Ed., Marcel Dekker Inc., N.Y.
- 2. The Theory & Practice of Industrial Pharmacy, 3rd edition, Leon Lachman, Herbert A. Lieberman, Joseph. L. Karig, Varghese Publishing House, Bombay.
- 3. Validation Master Plan by Terveeks or Deeks, Davis Harwood International publishing.
- 4. Validation of Aseptic Pharmaceutical Processes, 2nd Edition, by Carleton & Agalloco, (Marcel Dekker).
- 5. Michael Levin, Pharmaceutical Process Scale-Upl, Drugs and Pharm. Sci.Series, Vol. 157,2nd Ed., Marcel Dekker Inc., N.Y.
- 6. Validation Standard Operating Procedures: A Step by Step Guide for Achieving Compliance in the Pharmaceutical, Medical Device, and Biotech Industries, Syed Imtiaz Haider
- 7. Pharmaceutical Equipment Validation: The Ultimate Qualification Handbook, Phillip A. Cloud, Interpharm Press
- 8. Validation of Pharmaceutical Processes: Sterile Products, Frederick J.Carlton (Ed.) and James Agalloco (Ed.), Marcel Dekker, 2nd Ed.
- 9. Analytical Method validation and Instrument Performance Verification by Churg Chan, Heiman Lam, Y.C. Lee, Yue. Zhang, Wiley Inter Science.



#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

Course Code	HERBAL AND COSMETIC ANALYSIS	L	T	P	C					
21S07203		4	0	0	4					
	Semester									
Course Objectives	:									
requirements, herba	esigned to impart knowledge on analysis of herbal producted all drug interaction with monographs. Performance evaluation of ce better understanding of the equipments used in cosmetic in	osm	etic j	produ	acts					
<b>Course Outcomes</b>	(CO): Student will be able to									
• Determination	of herbal remedies and regulations									

- Analysis of natural products and monographs
- Determination of Herbal drug-drug interaction
- Principles of performance evaluation of cosmetic products.

#### UNIT – I

#### Herbal remedies- Toxicity and Regulations

Herbals vs Conventional drugs, Efficacy of herbal medicine products, Validation of Herbal Therapies, Pharmacodynamic and Pharmacokinetic issues. Herbal drug standardization: WHO and AYUSH guidelines

#### UNIT – II

#### **Adulteration and Deterioration:**

Introduction, types of adulteration/substitution of herbal drugs, Causes and Measure of adulteration, Sampling Procedures, Determination of Foreign Matter, DNA Finger printing techniques in identification of drugs of natural origin, heavy metals, pesticide residues, phototoxin and microbial contamination in herbal formulations. Regulatory requirements for setting herbal drug industry: Global marketing management, Indian and international patent law as applicable herbal drugs and natural products and its protocol.

#### UNIT – III

#### Testing of natural products and drugs

Effect of herbal medicine on clinical laboratory testing, Adulterant Screening using modern analytical instruments, Regulation and dispensing of herbal drugs, Stability testing of natural products, protocol. Monographs of Herbal drugs: Study of monographs of herbal drugs and comparative study in IP, USP, Ayurvedic Pharmacopoeia, American herbal Pharmacopoeia, British herbal Pharmacopoeia, Siddha and Unani Pharmacopoeia, WHO guidelines in quality assessment of herbal drugs.

#### UNIT – IV

#### Herbal drug-drug interaction

General principles, Validation of analytical method as per ICH guidelines and USP.

Computerized system validation: Electronic records and digitalsignificance-21 CFR part 11 and GAMP.

#### UNIT – V

#### **Evaluation of cosmetic products:**

Determination of acid value, ester value, saponification value, iodine value, peroxide value, rancidity, moisture, ash, volatile matter, heavy metals, fineness of powder, density, viscosity of cosmetic raw materials and finished products. Study of quality of raw materials and general methods of analysis of raw material used in cosmetic manufacture as per BIS.

Indian Standard specification laid down for sampling and testing of various cosmetics in finished



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

forms such as baby care products, skin care products, dental products, personal hygiene preparations, lips sticks. Hair products and skin creams by the Bureau Indian Standards

- 1. B. T. Loftus & R. A. Nash, "Pharmaceutical Process Validation", Drugs and Pharm Sci. Series, Vol. 129, 3rd Ed., Marcel Dekker Inc., N.Y.
- 2. The Theory & Practice of Industrial Pharmacy, 3rd edition, Leon Lachman, Herbert A. Lieberman, Joseph. L. Karig, Varghese Publishing House, Bombay.
- 3. Validation Master Plan by Terveeks or Deeks, Davis Harwood International publishing.
- 4. Validation of Aseptic Pharmaceutical Processes, 2nd Edition, by Carleton & Agalloco, (Marcel Dekker).
- 5. Michael Levin, Pharmaceutical Process Scale-Upl, Drugs and Pharm. Sci.Series, Vol. 157,2nd Ed., Marcel Dekker Inc., N.Y.
- 6. Validation Standard Operating Procedures: A Step by Step Guide for Achieving Compliance in the Pharmaceutical, Medical Device, and Biotech Industries, Syed Imtiaz Haider
- 7. Pharmaceutical Equipment Validation: The Ultimate Qualification Handbook, Phillip A. Cloud, Interpharm Press
- 8. Validation of Pharmaceutical Processes: Sterile Products, Frederick J.Carlton (Ed.) and James Agalloco (Ed.), Marcel Dekker, 2nd Ed.
- 9. Analytical Method validation and Instrument Performance Verification by Churg Chan, Heiman Lam, Y.C. Lee, Yue. Zhang, Wiley Inter Science.



#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

C	ourse Code	ADVANCED INSTRUMENTAL ANALYSIS LAB	$\mathbf{L}$	T	P	C
	21S07204		0 0		6	3
		Semester		I	I	
List	of Experiment	S				
1.	Comparison of	absorption spectra by UV and Wood ward – Fiesure rule				
2.	Interpretation of	of organic compounds by FT-IR				
3.	Interpretation of	of organic compounds by NMR				
4.	Interpretation of	of organic compounds by MS				
5.	Determination	of purity by DSC in pharmaceuticals				
6.	Identification of	f organic compounds using FT-IR, NMR, CNMR and Mass spe	ectra			
7.	Testing of relat	ed and foreign substances in drugs and raw materials				
8.	-	naterials as per official monographs				

9. Calibration of UV – Visible Spectrophtometer/ HPLC/ GC/ FTIR

10. Cleaning validation of any one analytical equipment



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#### **COURSE STRUCTURE & SYLLABI**

Course Code	MODERN BIO-ANALYTICAL TECHNIQUES LAB	L	T	P	C
21S07205		0	0	6	3
	Semester		I	I	

#### List of Experiments

- 1. Protocol preparation and performance of bioanalytical method validation
- 2. Protocol preparation for the conduct of BA/BE studies according to guidelines
- 3. Bio molecules separation utilizing various sample preparation techniques and Quantitative analysis of components by HPLC techniques
- 4. Isolation of analgesics from biological fluids (blood serum and urine)
- 5. Identification of anti-histaminics drug in urine by TLC
- 6. Extraction of drugs and metabolites from biological matrices by SPE/LLE
- 7. HPLC separation of modern drug from plasma and its formulations (Diclofenac)
- 8. Stability indicating method development by HPLC of any API
- 9. Bio molecules separation utilizing various sample preparation techniques and Quantitative analysis of components by gel electrophoresis
- 10. Quality control methods for herbal materials/ Medicinal plant materials



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#### **COURSE STRUCTURE & SYLLABI**

Course Code	RESEARCH METHODOLOGY AND	L	T	P	C
21DRM101	INTELLECTUAL PROPERTY RIGHTS	4	0	0	4
	Semester		I	II	
Course Objectiv	es:				
To under	stand the research problem				
	the literature studies, plagiarism and ethics				
<ul> <li>To get the</li> </ul>	e knowledge about technical writing				
<ul> <li>To analyz</li> </ul>	ze the nature of intellectual property rights and new developments				
<ul> <li>To know</li> </ul>	the patent rights				
Course Outcome	es (CO): Student will be able to				
Understa	nd research problem formulation.				
Analyze	research related information				
<ul> <li>Follow re</li> </ul>	esearch ethics				
Understa	nd that today's world is controlled by Computer, Information	Tec	hnol	ogy,	but
tomorrow	world will be ruled by ideas, concept, and creativity.				
	nding that when IPR would take such important place in growth				
	is needless to emphasis the need of information about Intellectual	Prop	erty	Righ	nt to
	ted among students in general & engineering in particular.				
	nd that IPR protection provides an incentive to inventors for furth				
	stment in R & D, which leads to creation of new and better proc	lucts	, and	l in	turn
	out, economic growth and social benefits.				
UNIT - I					
Research Proble					
	arch problem, Sources of research problem, Criteria Character				
	, Errors in selecting a research problem, Scope and objectives of r				
	investigation of solutions for research problem, data coll	ectic	on,	analy	/S1S,
	cessary instrumentations				
UNIT – II					
Literature review					
	re studies approaches, analysis, Plagiarism, Research ethics.				
UNIT – III					
Report writing					
	al writing, how to write report, Paper Developing a Research Propo	sal,	Forn	nat o	f
	, a presentation and assessment by a review committee	1			
UNIT – IV					
Nature of Intelle	<u> </u>				
_	, Trade and Copyright. Process of Patenting and Developme			_	
research, innovat	ion, patenting, development. International Scenario: Internationa	l co	oper	ation	on

research, innovation, patenting, development. International Scenario: International cooperation on Intellectual Property. Procedure for grants of patents, Patenting under PCT.

#### UNIT – V

#### **Patent Rights:**

Scope of Patent Rights. Licensing and transfer of technology. Patent information and databases. Geographical Indications. New Developments in IPR: Administration of Patent System. New developments in IPR; IPR of Biological Systems, Computer Software etc. Traditional knowledge Case Studies, IPR and IITs.



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

#### **Textbooks:**

- 1. Stuart Melville and Wayne Goddard, "Research methodology: an introduction for science & engineering students"
- 2. Wayne Goddard and Stuart Melville, "Research Methodology: An Introduction"

- 1. Ranjit Kumar, 2nd Edition, "Research Methodology: A Step by Step Guide for beginners"
- 2. Halbert, "Resisting Intellectual Property", Taylor & Francis Ltd ,2007.
- 3. Mayall, "Industrial Design", McGraw Hill, 1992.
- 4. Niebel, "Product Design", McGraw Hill, 1974.
- 5. Asimov, "Introduction to Design", Prentice Hall, 1962.
- 6. Robert P. Merges, Peter S. Menell, Mark A. Lemley, "Intellectual Property in New Technological Age", 2016.
- 7. T. Ramappa, "Intellectual Property Rights Under WTO", S. Chand, 2008



#### M.PHARM. PHARMACEUTICAL ANALYSIS

**COURSE STRUCTURE & SYLLABI** 

# AUDIT COURSE-I



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

Course Code			T	P	C
21DAC101a		2	0	0	0
	Semester			I	
G 01: 4:	TDI: '11 11 . 1 .				
Course Objectiv	es: This course will enable students:				
<ul> <li>Understa</li> </ul>	nd the essentials of writing skills and their level of readability				
<ul> <li>Learn ab</li> </ul>	out what to write in each section				
	ualitative presentation with linguistic accuracy				
Course Outcome	es (CO): Student will be able to				
<ul> <li>Understa</li> </ul>	nd the significance of writing skills and the level of readability				
<ul> <li>Analyze</li> </ul>	and write title, abstract, different sections in research paper				
<ul> <li>Develop</li> </ul>	the skills needed while writing a research paper				
UNIT - I			e Hr		
	Research Paper- Planning and Preparation- Word Order- Useful I				
up Long Sentenc	es-Structuring Paragraphs and Sentences-Being Concise and Remo	oving	g Red	unda	ncy
-Avoiding Ambig	guity				
UNIT - II			e Hr		
	nents of a Research Paper- Abstracts- Building Hypothesis-Re			roblei	n -
Highlight Finding	gs- Hedging and Criticizing, Paraphrasing and Plagiarism, Cauteri	zatio	n		
UNIT - III	L	ectur	e Hrs	s:10	
	ew of the Literature - Methodology - Analysis of the Data-Find	ings	- Dis	cussi	on-
Conclusions-Rec	ommendations.				
UNIT - IV	1	Ιa	ctura	Hrs:	<u> </u>
	l for writing a Title, Abstract, and Introduction	LC	Cture	1115.	,
UNIT - V	Tot writing a Title, Abstract, and introduction	Lo	oturo	Hrs:	)
	luage to formulate Methodology, incorporate Results, put forth Ar				
Conclusions	uage to formulate Methodology, incorporate Results, put form Al	gume	21118 0	iiiu ui	aw
Suggested Read	ing .				
	R (2006) Writing for Science, Yale University Press (available or	God	oda I	Rooks	
	urriculum of Engineering & Technology PG Courses [Volume-I]	OUC	igic i	JOOKS	)
	006) How to Write and Publish a Scientific Paper, Cambridge Uni	verci	tv Pr	229	
•	N (1998), Handbook of Writing for the Mathematical Sciences, S		•	<b>C</b> C C C C C C C C C C C C C C C C C C	
J. Highman Highman		** #TA1	•		
_	Vallwork, English for Writing Research Papers, Springer New Yo	rk Do	ordre	cht	
	rg London, 2011	(			
	<u>,</u>				



#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### COURSE STRUCTURE & SYLLABI

Course Code				L	T	P	C
21DAC101b		DISASTER MANAGEMENT		2	0	0	0
		Semest	er		]	Ī	
Course Objecti	ves: This cour	se will enable students:					
• Learn to	demonstrate	e critical understanding of key concept	s in	disast	ter risk	reducti	ion
and hun	nanitarian resp	onse.					
<ul> <li>Criticall</li> </ul>	y evaluatedisa	sterriskreduction and humanitarian response	pol	licy and	l practic	e from	
^	e perspectives.						
		ngofstandardsofhumanitarianresponseandpr	acti	calrelev	anceins	specific	types
	ters and conflic						_
		estrengthsandweaknessesofdisastermanager					
program UNIT - I	iming in differ	ent countries, particularly their home countr	y or	the co	untries t	they wo	rk in
Introduction:	_						
	tion Easterson	dCianifiaanaa Diffamanaa Datuu an Hazandana	Dia	o atom. NI	otumo10 m	.d	
		dSignificance;DifferenceBetweenHazardance	DIS	aster, IN	aturaian	ıu	
		ce, Nature, Types and Magnitude.					
Disaster Prone				1 4 1	1		D
•		as Prone to Floods and Droughts, Landslide					
-	nd Coastal Ha	zards with Special Reference to Tsunam	ı; P	ost- Di	saster	Disease	s and
Epidemics		T					
UNIT - II							
Repercussions							
	-	Human and Animal Life, Destruction of		-			
_	· ·	ones, Tsunamis, Floods, Droughts and Famines					
		Reactor Meltdown, Industrial Accidents, Oil	Sli	cks and	Spills,	Outbre	aks of
Disease and Ep	idemics, War	and Conflicts.					
UNIT - III							
Disaster Prepa	aredness and I	Management:					
Preparedness:	Monitoring of	of Phenomena Triggering ADisasteror	Haz	ard; E	valuatio	on of	Risk:
Application of	Remote Sens	sing, Data from Meteorological and Oth	er 1	Agencie	es, Med	lia Re	ports:
Governmental	and Communit	y Preparedness.					
UNIT - IV							
Risk Assessme	ent Disaster R	isk:					_
Concept and	Elements, Di	saster Risk Reduction, Global and Nati	onal	l Disas	ster Ris	sk Situ	ation.
^		,GlobalCo-OperationinRiskAssessmentand					
1				6, -	Ι	1	

## UNIT - V Disaster Mitigation:

in Risk Assessment. Strategies for Survival.

Meaning, Conceptand Strategies of Disaster Mitigation, Emerging Trends In Mitigation. Structural Mitigation and Non-Structural Mitigation, Programs of Disaster Mitigation in India.

#### **Suggested Reading**



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

- 1. R.Nishith, Singh AK, "Disaster Management in India: Perspectives, issues and strategies
- 2. "'New Royal book Company..Sahni,PardeepEt.Al.(Eds.),"DisasterMitigationExperiencesAndReflections",PrenticeHa ll OfIndia, New Delhi.
- 3. GoelS.L.,DisasterAdministrationAndManagementTextAndCaseStudies",Deep&Deep Publication Pvt. Ltd., New Delhi



#### M.PHARM. PHARMACEUTICAL ANALYSIS

<b>Course Code</b>	SANSKRIT	FOR TECHNICAL KNOWLEDGE	L	T	P	C
21DAC101c			2	0	0	0
		Semeste	r	I	I	
Course Objecti	ves: This course	will enable students:				
To get a	working knowle	dge in illustrious Sanskrit, the scientific la	nguage in	the wo	orld	
<ul> <li>Learnin</li> </ul>	g of Sanskrit to in	nprove brain functioning				
<ul> <li>Learnin</li> </ul>	gofSanskrittodeve	elopthelogicinmathematics, science & others	ubjects e	nhancin	g the	
memory	power					
• The eng	ineering scholars	equipped with Sanskrit will be able to exp	lore the	huge		
	dge from ancient					
	nes (CO): Studen					
	anding basic Sans					
		e about science &technology can be under	stood			
	logical language	will help to develop logic in students				
UNIT - I						
Alphabets in S	anskrit,					
UNIT - II						
	ure Tense, Simple	e Sentences				
UNIT - III						
Order, Introduct	ion of roots					
UNIT - IV						
Technical info	mation about San	skrit Literature				
UNIT - V						
Technical conc	epts of Engineering	ng-Electrical, Mechanical, Architecture, M	athematic	es		
Suggested Read	ling					
1."Abhyaspust	akam" –Dr.Vish	was, Sanskrit-Bharti Publication, New	Delhi			
2."Teach You	rself Sanskrit	" Prathama Deeksha- VempatiKutu	nbshastı	ri, Rash	triyaSa	nskrit
,	ew Delhi Public					
3."India's Gloa	ious ScientificT	radition" Suresh Soni, Ocean books (P	) Ltd.,N	ew Del	hi	



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

**COURSE STRUCTURE & SYLLABI** 

# AUDIT COURSE-II



#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

				1		<del></del>
Course Code		PEDAGOGY STUDIES	L	T	P	C
21DAC201a			2	0	0	0
		Semester		]	I	•
Course Objective	es: This cour	se will enable students:				
		ceonthereviewtopictoinformprogrammedesigna	ndpolic	y makii	ng	
	•	O, other agencies and researchers.				
• Identify c	ritical eviden	ce gaps to guide the development.				
Course Outcome	s (CO): Stud	lent will be able to				
Students will be a	ble to unders	tand:				
<ul> <li>Whatpeda</li> </ul>	igogicalpract	icesarebeingusedbyteachersinformalandinforma	alclassr	ooms in	develo	ping
countries'	?					
<ul> <li>What is th</li> </ul>	ne evidence o	n the effectiveness of these pedagogical practic	es, in v	vhat		
		hat population of learners?				
		on(curriculumandpracticum)andtheschoolcurric	culuma	nd guid	ance	
	best support	effective pedagogy?				
UNIT - I						
		ogy: Aims and rationale, Policy back ground,				
terminology	Theories	oflearning, Curriculum, Teachereducation. Con	ceptua	lframew	ork,Res	search
questions. Overv	iew of metho	odology and Searching.				
UNIT - II						
	viow. Podog	ogical practices are being used by teachers	in fo	rmol or	nd inf	ormal
		ogical practices are being used by teachers ntries. Curriculum, Teacher education.	111 10	illiai ai	ia iiii	Offilal
ciassioonis in de	veloping cou	nures. Curriculum, reacher education.				
UNIT - III						
	effectiveness	ofpedagogicalpractices,Methodologyfortheinder	ntheta or	anality	1 200ACC	men f
		teacher education (curriculumandpracticum)				
		ort effective pedagogy? Theory of change. Stren				
		ogical practices. Pedagogic theory and pedago				
attitudes and beli			<b>5</b> 1001 0	ppromen.		
		6.6				
UNIT - IV						
Professional dev	velopment: a	lignment with classroom practices and follow-u	p suppo	ort, Peer	suppor	t.
Support from the		1	. 11	,	11	
		riculumandassessment,Barrierstolearning:limite	dresour	cesand	large cl	ass
sizes						
TINITED TI						

#### **Suggested Reading**

UNIT - V

1. AckersJ, HardmanF(2001)ClassroominteractioninKenyanprimaryschools, Compare, 31 (2): 245-261.

Researchgapsandfuturedirections: Researchdesign, Contexts, Pedagogy, Teachereducation,

Curriculum and assessment, Dissemination and research impact.

 $2. \quad A grawal M(2004) Curricular reformins chools: The importance of evaluation, Journal of the control of th$ 



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

- 3. Curriculum Studies, 36 (3): 361-379.
- 4. AkyeampongK(2003) Teacher training in Ghana does it count? Multi-site teachereducation research project (MUSTER) country report 1. London: DFID.
- 5. Akyeampong K, LussierK, PryorJ, Westbrook J (2013)Improving teaching and learning of basic maths and reading in Africa: Does teacherpreparation count?International Journal Educational Development, 33 (3): 272–282.
- 6. Alexander RJ(2001) Culture and pedagogy: International comparisons in primary education. Oxford and Boston: Blackwell.
  - Chavan M (2003)ReadIndia: A mass scale, rapid, 'learning to read'campaign.
- 7. www.pratham.org/images/resource%20working%20paper%202.pdf.



#### M.PHARM. PHARMACEUTICAL ANALYSIS

<b>Course Code</b>	CEDI			L	T	P	C
21DAC201b	SIRI	ESSMANAGEMENT BY YOGA		2	0	0	0
		Semest	er		I	I	
Course Objecti	ves: This course	e will enable students:					
To achie	eve overall healt	h of body and mind					
• To over	come stres						
<b>Course Outcon</b>	es (CO): Studer	nt will be able to					
Develop	healthy mind ir	n a healthy body thus improving social hea	lth a	also			
• Improve	efficiency						
UNIT - I							
Definitions of I	Eight parts of yo	g.(Ashtanga)					
UNIT - II							
Yam and Niyar	n.						
UNIT - III							
Do`sand Don't	sin life.						
i) Ahinsa,satya	astheya,bramhad	charyaand aparigrahaii)					
	h,tapa,swadhyay	,ishwarpranidhan					
UNIT - IV							
Asan and Prana	yam						
UNIT - V							
i)Variousyogpo	sesand theirben	efitsformind &body					
ii)Regularizatio	onofbreathingtecl	hniques and its effects-Types of pranayam					
Suggested Read							
		ing-Part-I": Janardan SwamiYogabhyasiM					
		e Internal Nature" by Swami Vivekan	anda	a, Adv	aita		
Ashrama (Public	cation Department	nt), Kolkata					



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

Course Code	PERSONALIT	Y DEVELOPMENT THE	ROUGHLIFE	L	T	P	C
21DAC201c	EN	LIGHTENMENTSKILL		2	0	0	0
			Semester		]	<u> </u>	
G 011 (1	mi :						
Course Objecti	ves: This course v	vill enable students:					
<ul> <li>To learn</li> </ul>	to achieve the hi	ghest goal happily					
		stable mind, pleasing persor	nality and detern	nination	ı		
	ken wisdom in stu						
	nes (CO): Student						
•	•	d-Geetawillhelpthestudentin	developinghispe	rsonali	tyand a	chieve	
•	est goal in life	1.0					
		ed Geetawilllead the nation		•	•	perity	
	Neetishatakam v	vill help in developing versa	tile personality of	of stude	nts		
UNIT - I	T 1 1 1 1						
	•	ent of personality					
	20,21,22(wisdom)						
•	31,32(pride &hero	oism)					
	28,63,65(virtue)						
UNIT - II							
	•	ent of personality					
	53,59(dont's)						
	73,75,78(do's)		1				
UNIT - III							
* *	y to day work and						
		oter2-Verses41,47,48,					
•		Chapter6-Verses5,13,17,23,	35,				
	Verses45,46,48.						
UNIT - IV							
	asic knowledge.						
		oter2-Verses 56,62,68					
*	-Verses 13, 14, 15, 1						
<b>.</b>	of Rolemodel. Sh	rimad Bhagwad Geeta:					
UNIT - V							
•	Verses 17, Chapter?	3-Verses 36, 37, 42,					
•	Verses 18,38,39						
_	- Verses37,38,63						
Suggested Read							
•	wadGita"bySwam	iiSwarupanandaAdvaitaAsh	ram(Publication)	Departi	nent),		
Kolkata	1 C-4.1 OT	di anima an anto N. 1 - P. C.	Santards D. S.	· 0	-154		
		ti-sringar-vairagya) by P.C	opinatn, Kashti	iyaSan	skrit		
Sansthanam,	New Delli.						



#### M.PHARM. PHARMACEUTICAL ANALYSIS

**COURSE STRUCTURE & SYLLABI** 

# OPEN ELECTIVE



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

Course Code	BIOLOGICAL SCREENING METHODS	L	T	P	C
21SOE301d	( Elective)	3	0	0	3
	Semester	III			

#### **Course Objectives:**

The students are going to study about various techniques for screening of drugs for various pharmacological activities and guide lines for handling animals and human and animal ethics for screening of drugs.

#### Course Outcomes (CO): Student will be able to know

- How to handle animals
- About various techniques for screening of drugs for different pharmacological activities
- Guidelines and regulations for screening new drug molecules on animals.

#### UNIT – I

#### **Drug discovery process:**

Principles, techniques and strategies used in new drug discovery. High throughput screening, human genomics, robotics and economics of drug discovery, Regulations. Alternatives to animal screening procedures, cell-line, patch—clamp technique, In-vitro models, molecular biology techniques.

#### UNIT - II

#### **Bioassays:**

Basic principles of bioassays, official bioassays, experimental models and statistical designs employed in biological standardization.

#### UNIT – III

#### **Toxicity Evaluations**

Principles of toxicity evaluations, ED50, LD50 and TD values, International guidelines (ICH recommendations).

Preclinical studies: General principles and procedures involved in acute, sub-acute, chronic, teratogenicity, mutagenicity and carcinogenicity.

#### UNIT - IV

#### **Screening of drugs**

Screening of different classes of drugs using micro-organisms. Vitamin and antibiotic assays.

Screening methods involved in toxins and pathogens.

#### UNIT – V

#### **Enzymatic screening methods**

α-glucosidase, α- amylase, DNA polymerase, nucleases, L-asparginase, lipases and peptidases.

- 1. Basic and clinical pharmacology by Bertram G. Katzung (International edition) lange medical book / Mc Graw Hill, USA 2001 8th edition
- 2. Pharmacology by Rang H.P, Dale MM and Ritter JM., Churchill Livingston, London, 4/e
- 3. Goodman and Gilman's The pharmacological basis of therapeutics (International edition) Mc Graw Hill, USA 2001 10th edition.
- 4. General and applid toxicology by B.Ballantyne, T.Marrs, P.Turner (Eds) The Mc Millan press Ltd, London.
- 5. Drug Discovery by Vogel's
- 6. Drug Discovery and evaluation Pharmacological assays by H.Gerhard. Vogel, 2nd edition, Springer verlag, Berlin, Heidelberg.
- 7. Tutorial Pharmacy (Vol I and II) by Cooper and Gunns.



#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### COURSE STRUCTURE & SYLLABI

Course Code	STABILITY OF DRUGS AND DOSAGE FORMS	L	T	P	C
21SOE301f	( Elective)	3	0	0	3
	Semester	III			

#### **Course Objectives:**

These topics are designed impart a specialized knowledge to preserve the properties of drugs and dosage forms during manufacture storage and shelf life. The understanding of properties and evaluation of stability during storage, by solution and solid state against several factors of degradation.

#### Course Outcomes (CO): Student will be able to

- Evaluation of stability of solutions, solids and formulations against adverse conditions.
- Suggest the measures to retain stability and storage conditions for retaining the efficacy of the products.

#### UNIT – I

#### **Drug decomposition mechanisms**

- 1. Hydrolysis and acyl transfers: Nature of reaction, structure and utility, stabilization of Pharmaceutical examples.
- 2. Oxidation: Nature of oxidation, kinetics of oxidation, oxidation pathways of pharmaceutical, Interest Inhibition of oxidation
- 3. Photolysis: Energetics of photolysis, kinetics photolysis, photolytic reactions of pharmaceutical interest, prevention of photolytic reactions.

#### UNIT – II

#### Solid state chemical decomposition

Kinetic of solids state decomposition, Pharmaceutical examples of solid-state decomposition, Pure drugs, drug excipient and drug-drug interaction in solid state, methods of stabilization.

Physical stability testing of dosage forms:

- 1. Solids tablets, capsules, powder and granules
- 2. Disperse systems
- 3. Microbial decomposition
- 4. Over-view, physical stability of novel drug carriers, liposomes, niosomes, nano-particles.

#### UNIT – III

Identification and quantitative determination of preservatives, Antioxidants, colouring materials, emulsifiers and stabilizers in Pharmaceutical formulation.

Analysis of drugs from biological samples including, selection of biological sample, extraction of drugs by various methods as LLE, SPE and Membrane filtration. Factors affecting extraction of drugs.

#### $\overline{UNIT-IV}$

General method of analysis to determine the quality of raw materials used in cosmetic industry. Indian Standard Specifications (ISI) laid down for sampling and testing of various cosmetics in finished form by the Bureau of Indian Standards

#### UNIT - V

Methods of analysis to determine the quality of cosmetics in the finished forms such as Hair care products, Skin care products, Baby care products, Dental products, Personal hygiene products, Colour cosmetics, Ethnic products, Colour makeup preparation, Lipsticks, Hair setting lotions and Eye shadows. Toxicity testing in cosmetics and Safety and Legislation of Cosmetic products.

Stability studies: Concept of stability studies.

a) cGMP& ICH guidelines for Accelerated stability Testing.



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#### **COURSE STRUCTURE & SYLLABI**

b) Interaction of containers & closure Compatibility Testing.

- 1. Comprehensive Pharmacy Review 5th Edition by Leon Shargel, Alan H. Mutnick, Paul F. Souney, Larry N. Sawnson 2004.
- 2. A.H. Beckett and J. B. Stenlake Practical Pharmaceutical Chemistry, Part I and Part II, 4<sup>th</sup> Edition.
- 3. G. H. Jeffery, J. Basset, J. Mendham, R. C. Denny (Rev. by) Vogels Text Book of Quantitative Chemical Analysis, 5th Edition 1989, ELBS.
- 4. The Controller of Publications; New Delhi, Govt. of India, Indian Pharmacopoeia, Vol. I and Vol. II 2010.
- 5. J. B. Wilkinson and R. J. Moore, Herry's Cosmeticology; Longman Scientific and Technical Publishers, Singapore.
- 6. P.D. Sethi; Quantitative Analysis of Drugs in Pharmaceutical Formulations, 3rd Edition 1997,
- 7. Classification of cosmetics raw materials and adjuncts IS 3958 of Indian Standards Institution (BIS).
- 8. Cosmetic and toilet goods methods of sampling IS 3958 of Indian Standards Institution (BIS).
- 9. Methods of sampling and test for various cosmetics as laid down by Bureau of Indian Standards.
- 10. Drug stability: Principles and practices by Jens T. Carstensen
- 11. Stability Testing of Drug Products by W. Grimm. 12. Stability of Drugs and Dosage Forms by Yoshioka and Stella.



#### M.PHARM. PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

Course Code	PHARMACOEPIDEMIOLOGY &	L	T	P	C
21SOE301e	PHARMACOECONOMICS (Elective-I)	3	0	0	3
_	Semester	III			

#### **Course Objectives:**

This course enables students to understand various pharmacoepidemiological methods and their clinical applications. Also, it aims to impart knowledge on basic concepts, assumptions, terminology, and methods associated with Pharmacoeconomics and health related outcomes, and when should be appropriate Pharmacoeconomic model should be applied for a health care regimen.

#### Course Outcomes (CO): Student will be able to

- Understand the various epidemiological methods and their applications
- Understand the fundamental principles of Pharmacoeconomics.
- Identify and determine relevant cost and consequences associated with pharmacy products and services
- Perform the key Pharmacoeconomics analysis methods
- Understand the Pharmacoeconomic decision analysis methods and its applications.
- Describe current Pharmacoeconomic methods and issues.
- Understand the applications of Pharmacoeconomics to various pharmacy settings.

#### $UNIT - \overline{I}$

#### Introduction to Pharmacoepidemiology

Definition, Scope, Need, Aims & Applications; Outcome measurement: Outcome measures, Drug use measures: Monetary units, Number of prescriptions, units of drug dispensed, defined daily doses, prescribed daily doses, Diagnosis and Therapy surveys, Prevalence, Incidence rate, Monetary units, number of prescriptions, unit of drugs dispensed, defined daily doses and prescribed daily doses, medications adherence measurements.

Concept of risk:

Measurement of risk, Attributable risk and relative risk, Time- risk relationship and odds ratio

#### UNIT - II

#### Pharmacoepidemiological Methods

Qualitative models: Drug Utilization Review; Quantitative models: case reports, case series, Cross sectional studies, Cohort and case control studies, Calculation of Odds' ratio, Meta-analysis models, Drug effects study in populations: Spontaneous reporting, Prescription event monitoring, Post marketing surveillance, Record linkage systems, Applications of Pharmacoepidemiology

#### UNIT – III

#### **Introduction to Pharmacoeconomics**

Definition, history of Pharmacoeconomics, Need of Pharmacoeconomic studies in Indian healthcare system. Cost categorization and resources for cost estimation: Direct costs. Indirect costs. Intangible costs. Outcomes and Measurements of Pharmacoeconomics: Types of outcomes: Clinical outcome, Economic outcomes, Humanistic outcomes; Quality Adjusted Life Years, Disability Adjusted Life Years Incremental Cost-Effective Ratio, Average Cost-Effective Ratio. Person Time, Willingness to Pay, Time Trade Off and Discounting.

#### UNIT - IV

#### Pharmacoeconomic evaluations

Definition, Steps involved, Applications, Advantages and disadvantages of the following Pharmacoeconomic models: Cost Minimization Analysis (CMA), Cost Benefit Analysis (CBA), Cost Effective Analysis (CEA), Cost Utility Analysis (CUA), Cost of Illness (COI), Cost Consequences Analysis (COA).



#### M.PHARM. IN PHARMACEUTICAL ANALYSIS

#### **COURSE STRUCTURE & SYLLABI**

UNIT -	– V
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#### Health related quality of life (HRQOL)

Definition, Need for measurement of HRQOL, Common HRQOL measures. Definition, Steps involved, Applications of the following: Decision Analysis and Decision tree, Sensitivity analysis, Markov Modeling, Software used in Pharmacoeconomic analysis, Applications of Pharmacoeconomics

- 1. Rascati K L. Essentials of Pharmacoeconomics, Woulters Kluwe rLippincott Williams & Wilkins, Philadelphia.
- 2. Thomas E Getzen. Health economics. Fundamentals and Flow of Funds. John Wiley & Sons, USA.
- 3. Andrew Briggs, Karl Claxton, Mark Sculpher. Decision Modeling for Health Economic Evaluation, Oxford University Press, London.
- 4. K G Revikumar, Pharmacoepidemiology and Pharmacoeconomics Concepts and Practices.
- 5. Michael Drummond, Mark Sculpher, George Torrence, Bernie O'Brien and Greg Stoddart. Methods for the Economic Evaluation of Health Care Programs Oxford University Press, London.
- 6. George E Mackinnon III. Understanding health outcomes and Pharmacoeconomics.
- 7. Graker, Dennis. Pharmacoeconomics and outcomes.
- 8. Walley, Pharmacoeconomics.
- 9. Pharmacoeconomic ed. by Nowakowska University of Medical Sciences, Poznan.
- 10. Relevant review articles from recent medical and pharmaceutical literature
- 11. Guru Prasad Mohanta and P K Manna, Textbook of Pharmacovigilance Concepts and Practice