ADVANCED INSTRUMENTAL ANALYSIS (MPA 201T)

Scope

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.

Objectives

After completion of course student is able to know,

- interpretation of the NMR, Mass and IR spectra of various organic compounds
- ☐ theoretical and practical skills of the hyphenated instruments
- identification of organic compounds

THEORY 60 Hrs

- 1. HPLC: Principle, instrumentation, pharmaceutical applications, 12 peak shapes, capacity factor, selectivity, plate number, plate Hrs 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.
- 2 Biochromatography: Size exclusion chromatography, ion 12 exchange chromatography, ion pair chromatography, affinity Hrs 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.

3 Super critical fluid chromatography: Principles, 12 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.
- 4 Mass spectrometry: Principle, theory, instrumentation of mass 12 spectrometry, different types of ionization like electron impact, Hrs 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.
- 5 NMR spectroscopy: Quantum numbers and their role in NMR, 12 Principle, Instrumentation, Solvent requirement in NMR, Hrs 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 with reference to 13CNMR: Spin spin and spin lattice relaxation phenomenon. 13C NMR, 1-D and 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.

MODERN BIO-ANALYTICAL TECHNIQUES (MPA 202T)

Scope

This subject is designed to provide detailed knowledge about the importance of analysis of drugs in biological matrices.

Objectives

Upon completion of the course, the student shall be able to understand

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

THEORY 60 Hrs

- 1. Extraction of drugs and metabolites from biological matrices: 12
 General need, principle and procedure involved in the Hrs
 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.
- 2 Biopharmaceutical Consideration:

12 Hrs

- 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.
- 3 Pharmacokinetics and Toxicokinetics:

12 Hrs

- 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.
- 4 Cell culture techniques

12 Hrs

Basic equipments 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.

5 Metabolite identification:

12 Hrs

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.

- 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.
- Pharmaceutical Analysis Modern methods Part B J W Munson, Volume 11, Marcel Dekker Series
- 5. Practical HPLC method Development Snyder, Kirkland, Glaich, 2nd Edition, John Wiley & Sons, New Jercy. USA.
- 6. Chromatographic Analysis of Pharmaceuticals John A Adamovics, 2nd Edition, Marcel Dekker, Newyork, USA. 1997.
- 7. Chromatographic methods in clinical chemistry & Toxicology Roger L Bertholf, 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

QUALITY CONTROL AND QUALITY ASSURANCE (MPA 203T)

Scope

This course deals with the various aspects of quality control and quality assurance aspects of pharmaceutical industries. It covers the important aspects like cGMP, QC tests, documentation, quality certifications, GLP and regulatory affairs.

Objectives

At the completion of this subject it is expected that the student shall be able to know

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

THEORY

60 hrs

Concept and Evolution of Quality Control and Quality

Assurance

Hrs

Good Laboratory Practice, GMP, Overview of ICH Guidelines – QSEM, with special emphasis on Q-series guidelines.

Good Laboratory Practices: Scope of GLP. Definitions. Quality

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.

- 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.
- 3. Analysis of raw materials, finished products, packaging 12 materials, in process quality control (IPQC), Developing Hrs 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.

12

Hrs

- 4. 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.
- 5. 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.

- 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, 4th 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.

HERBAL AND COSMETIC ANALYSIS (MPA 204T)

Scope

This course is designed to impart knowledge on analysis of herbal products. Regulatory requirements, herbal drug interaction with monographs. Performance evaluation of cosmetic products is included for the better understanding of the equipments used in cosmetic industries for the purpose.

Objectives

At compl	etion of this course student shall be able to understand
	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.

THEORY 60 Hrs

- Herbal remedies- Toxicity and Regulations: Herbals vs 12
 Conventional drugs, Efficacy of herbal medicine products, Hrs
 Validation of Herbal Therapies, Pharmacodynamic and
 Pharmacokinetic issues. Herbal drug standardization: WHO and
 AYUSH guidelines.
- 2 12 Adulteration Deterioration: Introduction. and types adulteration/substitution of herbal drugs, Causes and Measure of Hrs 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.
- 3 Testing of natural products and drugs: Effect of herbal 12 medicine on clinical laboratory testing, Adulterant Screening using Hrs 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 quidelines in quality assessment of herbal drugs.

4 Herbal drug-drug interaction: WHO and AYUSH guidelines for 12 safety monitoring of natural medicine, Spontaneous reporting Hrs schemes for bio drug adverse reactions, bio drug-drug and bio drug-food interactions with suitable examples. Challenges in monitoring the safety of herbal medicines.

12

Hrs

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 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. Pharmacognosy by Trease and Evans
- 2. Pharmacognosy by Kokate, Purohit and Gokhale
- 3. Quality Control Methods for Medicinal Plant, WHO, Geneva
- 4. Pharmacognosy & Pharmacobiotechnology by Ashutosh Kar
- 5. Essential of Pharmacognosy by Dr.S.H.Ansari
- 6. Cosmetics Formulation, Manufacturing and Quality Control, P.P. Sharma, 4th edition, Vandana Publications Pvt. Ltd., Delhi
- 7. Indian Standard specification, for raw materials, BIS, New Delhi.
- 8. Indian Standard specification for 28 finished cosmetics BIS. New Delhi
- 9. Harry's Cosmeticology 8th edition
- 10. Suppliers catalogue on specialized cosmetic excipients
- 11. Wilkinson, Moore, seventh edition, George Godwin. Poucher's Perfumes, Cosmetics and Soaps
- 12. Hilda Butler, 10th Edition, Kluwer Academic Publishers. Handbook of Cosmetic Science and Technology, 3rd Edition,

PHARMACEUTICAL ANALYSIS PRACTICALS - I (MPA 205P)

- 1. Comparison of absorption spectra by UV and Wood ward Fiesure rule
- 2. Interpretation of organic compounds by FT-IR
- 3. Interpretation of organic compounds by NMR
- 4. Interpretation of organic compounds by MS
- 5. Determination of purity by DSC in pharmaceuticals
- 6. Identification of organic compounds using FT-IR, NMR, CNMR and Mass spectra
- 7. Bio molecules separation utilizing various sample preparation techniques and Quantitative analysis of components by gel electrophoresis.
- 8. Bio molecules separation utilizing various sample preparation techniques and Quantitative analysis of components by HPLC techniques.
- 9. Isolation of analgesics from biological fluids (Blood serum and urine).
- 10. Protocol preparation and performance of analytical/Bioanalytical method validation.
- 11. Protocol preparation for the conduct of BA/BE studies according to guidelines.
- 12. In process and finished product quality control tests for tablets, capsules, parenterals and creams
- 13. Quality control tests for Primary and secondary packing materials
- 14. Assay of raw materials as per official monographs
- 15. Testing of related and foreign substances in drugs and raw materials
- 16. Preparation of Master Formula Record.
- 17. Preparation of Batch Manufacturing Record.
- 18. Quantitative analysis of rancidity in lipsticks and hair oil
- 19. Determination of aryl amine content and Developer in hair dye
- 20. Determination of foam height and SLS content of Shampoo.
- 21. Determination of total fatty matter in creams (Soap, skin and hair creams)
- 22. Determination of acid value and saponification value.
- 23. Determination of calcium thioglycolate in depilatories