Semester: M. Pharmacy 1st Semester Branch: Pharmacy

Subject: Advance Research Methods

Total Theory Periods: 50

Subject Code: 565111(41)

Total Tutorial Periods: 12

Total Marks in the End Semester: **100**Minimum of Class Test to be Conducted: **2**

Unit - 1:

Spectroscopic Method – Introduction, application structure elucidation using UV, IR, NMR, Mass spectrometry with examples.

<u>Unit - 2:</u>

Separation Techniques – Theory, Instrumentation and application of GLC, HPLC, HPTLC, Chiral chromatography, Ion Pair Chromatography.

<u>Unit - 3:</u>

Thermal Analysis – Theory, Instrumentation and application of thermo-gravimentric analysis, differential thermal analysis.

Unit - 4:

Calorimetric analysis – theory, instrumentation, chemical application and structural elucidation, differential scanning calorimetric (DSC), Isothermal titration.

<u>Unit - 5 :</u>

Immunochemical techniques – Immunelectrophoresis, immunoprecipation, ELISA, radioimmunoassay.

Books Recommended:

- 1. Practical Pharmaceutical Chemistry, Backett, and Stenlake.
- 2. Spectrophotometric identification of organic compound, Silverstein.
- 3. Vogel's Text book of Quality analysis, 5th and 6th edition, Svehla.
- 4. Textbook of Pharmaceutical chemistry, L. G. Chatten.
- 5. Instrumental Method of Chemical Analysis.

Semester: M. Pharmacy 1st Semester

Subject: Pharmacology and Biostatistics

Total Theory Periods: 50

Total Marks in the End Semester: **100**Minimum of Class Test to be Conducted: **2**

Branch: **Pharmacy**

Subject Code: **565112(41)** Total Tutorial Periods: **12**

Unit - 1:

Drug dependence, tolerance, abuse drug allergy and resistance.

<u>Unit - 2:</u>

Genetics, gene cloning, gene delivery and recombinant DNA.

Unit - 3:

Molecular pharmacology, receptor theories, receptor isolation radio- ligand binding studies, Signal transduction mechanism of the cell.

Unit - 4:

Therapeutics regimens – therapeutics response and toxicity, dosage regimens, clinical trial studies, ADME – Pharmacokinetics, Drug – drug interaction and bioassay.

<u>Unit - 5:</u>

Biological screening of new compounds and New drug discovery.

Unit - 6:

Bio-statistics – Student "t" test, chi-square test, correlation probit analysis, analysis of variances.

Books Recommended:

- 1. The Pharmacological basis of therapeutics-Goodman and Gill man's
- 2. Pharmacology-Rang & Dale.
- 3. Pharmacology-Katzung.
- 4. Fundamentals of experimental Pharmacology-By M.N.Ghosh
- 5. Hand book of Experimental Pharmacology-S.K.Kulakarni
- 6. Text book of in vitro practical Pharmacology by Ian Kitchen
- 7. Pharmacological Experiments on intact preparations by Churchill Living stone.
- 8. Hand book of Clinical Pharmacokinetics Gibaldi and Prescott.
- 9. Indian Pharmacopoeia and other Pharmacopeias.
- 10. Screening methods in Pharmacology by Robert Turner.A
- 11. Clinical trials and tribulations by Allien E.Cato
- 12. Drug discovery and Evaluation by Vogel H.G.

Semester: M. Pharmacy 1st Semester Branch: **Pharmacy**

Subject: Drug Regulatory Affairs and Quality Assurance Subject Code: **565113(41)** Total Tutorial Periods: 12

Total Theory Periods: 50

Total Marks in the End Semester: 100 Minimum of Class Test to be Conducted: 2

<u>Unit - 1:</u>

Requirement of GMP, CGMP, GLP, USFDA, WHO guidelines and ISO 9000 series.

Drug and cosmetics acts and rules. Drug regulatory affairs.

Unit - 2:

Documentation - Protocols, forms and maintenance of record in Pharmaceuticals industry.

Preparation of documentation of new drug approval and export registration, processing and its application intellectual property rights (patent, copyright and trade marks) Sewage disposal and pollution control.

Unit - 4:

Concept in validation of manufacturing, analytical and process, validation and its application.

<u>Unit - 5 :</u>

Basic concept of quality control and quality assurance system, source and control of quality variation of raw material, containers, closures personnel, environmental etc.

Unit - 6:

In process quality control test, IPQC problem in pharmaceutical industries, ICH guidelines.

<u>Unit - 7:</u>

Sampling plans, Sampling and characteristics curves, Master formula generation and maintenance, standard operating procedure (SOP) for different dosage forms.

Book Recommended:

- 1. Theory and Practice of Industrial Pharmacy By Lachmann and Libermann
- 2. Pharmaceutical dosage forms: Tablets Vol. 1-3 by Leon Lachmann.
- 3. Pharmaceutical Dosage forms: Disperse systems, Vol, 1-2; By Leon Lachmann.
- 4. Pharmaceutical Dosage forms: Parenteral medications Vol. 1-2; By Leon Lachmann.
- 5. Modern Pharmaceutics; By Gillbert and S. Banker.
- 6. Remington's Pharmaceutical Sciences.
- 7. Advances in Pharmaceutical Sciences Vol. 1-5; By H.S. Bean & A.H. Beckett.
- 8. Physical Pharmacy; By Alfred martin
- 9. Bentley's Textbook of Pharmaceutics Rawbins.
- 10. Good manufacturing practices for Pharmaceuticals: A plan for total quality control, Second edition; By Sidney H. Willig.
- 11. Quality Assurance Guide; By Organization of Pharmaceutical producers of India.
- 12. Drug formulation manual; By D.P.S. Kohli and D.H.Shah. Eastern publishers, New Delhi.
- 13. How to practice GMPs; By P.P.Sharma. Vandhana Publications, Agra.
- 14. Pharmaceutical Process Validation; By Fra. R. Berry and Robert A. Nash.
- 15. Pharmaceutical Preformulations; By I.J. Wells.
- 16. Applied production and operations management; By Evans, Anderson, Sweeney and Williams.

Semester: M. Pharmacy 1st Semester Subject: Formulation Development

Total Theory Periods: 50

Total Marks in the End Semester: **100** Minimum of Class Test to be Conducted: **2**

Branch: **Pharmacy**Subject Code: **565114(41)**Total Tutorial Periods: **12**

Unit - 1:

Stability, solubility, Pka, Dissolution rate, Partition Coefficient. In Vitro and In Vivo evaluation techniques, product formulation and CGMP.

<u>Unit -2:</u>

Designing of Pharmaceuticals - Tablets formulation, special tablets and preparation of components for compression. Characterization of granulation, Coating of tablets, evaluation of tablets. Equipment and processing problem in tablets.

<u>Unit - 3:</u>

Topical and rectal absorption of drug, formulations and evaluations.

Unit - 4:

Formulation consideration of oral liquids, suspension, emulsion, development of various products.

<u>Unit - 5:</u>

Formulation consideration of parenteral ophthalmic, depot products, large volume and small volume parenteral, environmental control and quality assurance in parenteral drug manufacturing.

Unit - 6:

Stability in pharmaceuticals and study of stability kinetics.

Unit - 7:

Introduction to controlled and novel drug delivery system, Sustained release dosage form, prodrug concept, Nanoparticals, Liposomes, Resealed erythrocytes, Transdermal and other Novel drug delivery systems.

<u>Unit - 8:</u>

Types of container and closures, packaging and stability assessment.

Optimization techniques in pharmaceutical formulations and processing.

Pilot plant and scale up techniques.

Book Recommended:

- 1. Controlled Drug Delivery System, J.R. Robinson and V.H.S.L. Lee.
- 2. Physical Pharmacy, 4th edition, A. Martin, J.C. Swarbrick.
- 3. Pharmaceutical analysis, 'Ramington' A. R. Gennaro.
- 4. The theory and practice of Industrial pharmacy, IIIrd edition, L. Lachman, H. A. Liberman.
- 5. Modern Pharmaceutics, IInd edition, G. S. Banker, C.T. Rhodes.

Semester: M. Pharmacy 1st Semester Branch: Pharmacy

Subject: Advance Research Methods (Lab)

Subject Code: 565121(41)

Total Practical Periods: 72

Total Marks in the End Semester: 100

List of Experiment:

- 1. Determination of α max and Linearity of methylene blue by spectroscopic method.
- 2. To determine the absorption curve of aromatic hydrocarbons and the analysis of binary mixture
- 3. Estimation of Aspirin by colorimetry.
- 4. Assay of Paracetamol tablet by UV spectroscopy.
- 5. Determination of the active constituents in a medicinal preparation by derivative spectroscopy.
- 6. Estimation of Paracetamol by HPLC.
- 7. Identification of given sample by paper chromatography.
- 8. Identification of drug's by TLC.
- 9. To determine the purity of commercial benzoic acid using compressed discs (IR).
- 10. Interpretation of given sample by IR spectra.

Books Recommended:

- 1. Practical Pharmaceutical Chemistry, Backett, and Stenlake.
- 2. Spectrophotometric identification of organic compound, Silverstein.
- 3. Vogel's Text book of Quality analysis, 5th and 6th edition, Svehla.

Semester: M. Pharmacy 1st Semester Branch: Pharmacy

Subject: Pharmacology and Biostatics (Lab)

Subject Code: 565122(41)

Total Practical Periods: 72

Total Marks in the End Semester: 100

List of Practicals:

1. To Study the maintenance of common laboratory animals.

- 2. Bioassay of the more important biogenic agents by various methods.
- 3. Pharmacological Screening methods used for CNS, Local anesthetics, Endocrine and In-vitro microbial screening.
- 4. Protocol design of Clinical Trials.
- 5. Biostatical study of given data.

Books Recommended:

- 1. The Pharmacological basis of therapeutics-Goodman and Gill man's
- 2. Pharmacology- Rang & Dale.
- 3. Pharmacology-Katzung.
- 4. Fundamentals of experimental Pharmacology-By M.N.Ghosh
- 5. Hand book of Experimental Pharmacology-S.K.Kulakarni
- 6. Text book of in vitro practical Pharmacology by Ian Kitchen
- 7. Pharmacological Experiments on intact preparations by Churchill Living stone.
- 8. Hand book of Clinical Pharmacokinetics Gibaldi and Prescott.
- 9. Indian Pharmacopoeia and other Pharmacopeias.
- 10. Screening methods in Pharmacology by Robert Turner.A
- 11. Clinical trials and tribulations by Allien E.Cato
- 12. Drug discovery and Evaluation by Vogel H.G.

JOURNALS

- 1. Indian Journal of Pharmacology.
- 2. Indian Journal of Physiology and Pharmacology.
- 3. Indian Journal of Experimental Biology.
- 4. Pharmacological research.

Semester: M. Pharmacy 1st Semester Branch: Pharmacy

Subject: Formulation Development (Lab) Subject Code: 565123(41)

Total Practical Periods: 72

Total Marks in the End Semester: 100

- 1. To prepare and evaluate aspirin tablets by wet granulation method.
- 2. To evaluate and compare at least three marketed Paracetamol tablets.
- 3. To study the effect of various binders on the hardness and dissolution rate of ascorbic acid tablets, at different concentration.
- 4. To prepare 10gm of sustained release granules of ascorbic acid by Microencapsulation method.
- 5. To perform the pre-formulation studies of the given sample of ascorbic acid.
- 6. To study the dissolution profile of marketed sustained release products of aspirin.
- 7. To prepare and evaluate partially flocculated suspension of Paracetamol by using electrolyte.
- 8. To prepare and evaluate suspension of aspirin.
- 9. To study the effect of various suspending agents on sedimentation rate at different concentration.

Book Recommended:

- 1. Controlled Drug Delivery System, J.R. Robinson and V.H.S.L. Lee.
- 2. Physical Pharmacy, 4th edition, A. Martin, J.C. Swarbrick.
- 3. Pharmaceutical analysis, 'Ramington' A. R. Gennaro.
- 4. The theory and practice of Industrial pharmacy, IIIrd edition, L. Lachman, H. A. Liberman.
- 5. Modern Pharmaceutics, IInd edition, G. S. Banker, C.T. Rhodes.

Semester: M. Pharmacy 2nd Semester

Subject: Pharmaceutical Quality Assurance-I

Total Theory Periods: 40

Total Marks in the End Semester: 100 Minimum of Class Test to be conducted: 2 Branch: Pharmacy

Subject Code: 5119211(041) Total Tutorial Periods: 12

Unit-1:

A detailed study of the principles, instrumentations and applications in drug analysis of: GC-MS, LC-MS & FTIR with reference to drug metabolism, toxicologic and forensic studies, diagnosis of disease state, quantification of drugs in biological samples, Super critical fluid chromatography and size exclusion chromatography

Unit-2:

A detailed study of the principles, instrumentation and applications of the following Instrumental analysis: X-ray fluorescence spectrometry, atomic absorption spectroscopy, Inductively coupled plasma- mass spectroscopy

Unit-3:

Microscopy: General aspects, hot stage microscopy, scanning electron microscopy (SEM), transmission electron microscopy (TEM): principle, instrumentation and applications.

Particles size analysis: Zetameter, Photon correlation spectroscopy, counter-counter apparatus, atomic force microscopy and confocal.

Unit-4:

A detailed study of the various principles and procedure involved in the quantitative analysis of pharmaceutical preparations and dosage forms containing the following groups of drugs included in I.P. (Biological and microbiological methods excluded)

- (a) Analgesics and Antipyretics (b) Sedatives & Tranquillizers
- (c) Antihypertensives (d) Antibiotics (e) Cardiovascular drugs (f) Vitamins (g) Antihistaminics

Unit – 5:

Impurity Profile: Sources of impurities, their effect on drug stability and therapeutic action. Determination of impurities in bulk drugs - Isolation, characterization, and analytical methods. Formulation related impurities - Isolation, characterization, and analytical methods. ICH and WHO guidelines for impurity and related substances in the drugs.

RECOMMENDED BOOKS:

- 1. Pharmaceutical Analysis by Ohannason
- 2. Chemical Analysis by Settle
- 3. Pharmaceutical Analysis Modern Methods by Munson
- 4. Chemical Analysis Modern Instrumentation methods and techniques by Wiley.
- 5. Instrumental methods of analysis by Willard Dean & Merrit.
- 6. Hand book of Instrumental techniques for analytical chemistry edited by Frank settle pub. by Prentice Hall Inc.
- 7. A text book of Pharmaceutical analysis by K.A.Conners (John Wiley)
- 6. I.P. 1996, Vol.-I & II

Semester: M. Pharmacy 2nd Semester

Subject: Pharmaceutical Quality Assurance-II

Total Theory Periods: 40

Total Marks in the End Semester: 100
Minimum of Class Test to be Conducted: 2

Branch: **Pharmacy**Subject Code: **5119212(041)**

Total Tutorial Periods: 12

Unit-1: Regulatory requirements for import, export, manufacture and sale of drug and formulation. Types of manufacturing licenses. Master Formula and Batch Manufacturing Records.

Unit -2: Air-handling systems required for pharmaceutical manufacturing. Regulatory requirements and monitoring. Validation of air-handling system.

Finished products release, Quality review, Quality audits, Batch release documents, Complains and recalls, Evaluation of complaints recall procedures, relevant records and documents.

Unit -3: Guidelines for Industrial Safety & Health. Materials Safety Data Sheet (MSDS). Procedures for obtaining license for manufacture of narcotic and psychotropic drugs. Records to be maintained for manufacturing alcoholic preparations.

Unit -4: The Drug Price Control Order, 1985. The Medicinal & Toiletry Preparations Act, (Excise Duties Act, 1955 & Rules, 1976). Disposal of wastes and scraps, procedures and records.

Unit -5: Legislation to regulate import, manufacture, sale and distribution of cosmetics. Quality Control and Quality Assurance of cosmetics.

Legislation to regulate import, manufacture, sale and distribution of herbal drugs and their preparations. Quality Control and Quality Assurance of herbal drugs.

RECOMMENDED BOOKS:

- 1. The internal quality audit by Monica Girmaldi and Janet Gough Davis Harwood International Publishing.
- 2. Validation Master plan by Terveeks or Deeks, Davis Harwood International Publishing.
- 3. Validation of Asceptic Pharmaceutical Processes, 2nd Edition, by Carleton & Agalloco, (Marcel Dekker).
- 4. Statistical Design and Analysis in Pharmaceutical Science, by Chow, (Marcel Dekker).
- 5. Automation & Validation of Information in Pharmaceutical Processing, by deSPAUTZ, (Marcel Dekker).
- 6. Guidelines for Laboratory Quality Auditing, by Singer, (Marcel Dekker).
- 7. Pharmaceutical Experimental Design, by Lewis, (Marcel Dekker).
- 8. New Drug approval process, 2nd edition, Vol. 56, by Guarino, Marcel Dekker., New York.
- 9. Hosting a compliance Audit by Janet Gough Davis Harwood International Publishing.

Semester: M. Pharmacy 2nd Semester Branch: Pharmacy

Subject: Pharmaceutical Quality Assurance–III

Total Theory Periods: 40

Subject Code: 5119213(041)

Total Tutorial Periods: 12

Total Marks in the End Semester: 100
Minimum of Class Test to be Conducted: 2

Unit – 1:

Basis for process validation. Process validation and Quality Assurance, Process validation as a Quality Introduction to Pharmaceutical Validation:

Definition, Manufacturing Process Model, Government regulation, scope of Validation Advantage of Validation, Organization for Validation, Validation Master plan, URS, D.Q., IQ, OQ & P.Q. of facilities. Regulatory assurance tool, Prospective, retrospective and concurrent process validation.

Unit – 2:

Validation of Equipment

Concept of URS, DQ, IQ, OQ & PQ, Validation of following equipment

- Dry Powder Mixers
- Fluid Bed and Tray dryers.
- Tablet Compression M/c.
- Dry Heat Sterilization/Tunnels
- Autoclaves
- Capsule filling machines.
- Validation of Integrated lines by media fill test.
- Validation of existing equipment.
- Sterile product validation,
- Validation of water systems for sterile and non-sterile products.

Unit -3:

Validation of Utilities

Validation of Pharmaceutical Water System & pure steam, Validation of HAVC system. Validation of Compressed air. Cleaning of Equipment, Cleaning of Facilities. Computer System Validation.

Unit – 4:

Equipment and Analytical Method Validation

General principles of analytical method validation, Validation of following analytical Instruments include UV-visible spectrophotometer, FT-IR spectrometer, HPLC and GC-MS.

Unit – 5:

Process Validation

Prospective, concurrent, retrospective & revalidation, Process validation of following formulations

- Coated tablets
- Capsules
- Ampoules & Vials
- Ointment/Creams
- Liquid Orals

RECOMMENDED BOOKS

- 1. How to Practice GMPs, P.P.Sharma, Vandana Publication, 5th Ed.
- 3. Q.A. Mannual, D.H.Shah, Business Horizon, 1st Ed.
- 4. SOP Guidelines, D.H.Shah, Business Horizon, 2nd Ed.
- 5. The International Pharmacopoeia Vol 1,2,3,4, General Methods of Aalysis & Quality Specifications for Pharmaceutical Substances, Excipients, Dosage forms., CBS (WHO), 3rd Ed.
- 6. Quality Assurance of Pharmaceuticals A compendium of guidelines and related materials Vol.1 & Vol.2, WHO, 1999
- 7. Pharmaceutical Process Validation, Wachter & Mash, M.Dekker, 3rd Ed.
- 8. Pharmaceutical Process Validation, Berry & Mash, M.Dekker, 3rd & 4thEd.

Semester: M. Pharmacy 2nd Semester

Subject: Pharmaceutical Quality Assurance-IV

Total Theory Periods: 40

Total Marks in the End Semester: 100 Minimum of Class Test to be conducted: 2 Branch: **Pharmacy**

Subject Code: 5119214(041)

Total Tutorial Periods: 12

Unit – 1:

1. Concept of total quality management, philosophy of GMP, CGMP and GLP.

Introduction to GMPs-MHRA, GMPs-HPFBI, GMPs-MCC, GMPs-EDQM etc

2. Audits: GMP compliance audit, Audit policy, Internal, external, second party and third party audits. Preparation for audit, conducting audit, audit analysis, audit report and follow up.

Unit-2:

- 3. Organization and personnel, responsibilities, training hygiene.
- 4. Premises: Location, design, plan layout, construction, maintenance and sanitations, environmental control, sterile areas, control of contamination.
- 5. Equipments: Selection, purchase specifications, maintenance, clean in place, sterilize in place.
- 6. Raw materials: Purchase specifications, maintenance of stores, selection of vendors, controls and raw materials.
- 7. Manufacture of and controls on dosage forms: Manufacturing documents master formula, batch formula records, standard operating procedures, quality audits of manufacturing processes and facilities.

Unit-3:

- 8. In process quality control on various dosage forms sterile, biological products and non-sterile, standard operating procedures for various operations like cleaning, filling, drying, compression, coating, disinfection, sterilization, membrane filtration etc. Guidelines for Quality assurance of Human Blood products and Large volume parenterals.
- 9. Packaging and labeling controls, line clearance and other packaging materials.
- 10. Quality control laboratory: Responsibilities, good laboratory practices, routine controls, instruments, protocols, non-clinical testing, controls on animal house, data generation and storage, quality control documents, retention samples, records, audits of quality control facilities.

Unit-4:

- 11. Finished products release: Quality review, quality audits, batch release document.
- 12. Distribution and distribution-records: Handling of returned goods recovered materials and reprocessing.
- 13. Complaints and recalls, evaluation of complaints recall procedures, related records and documents.

Unit-5:

14. Good Laboratory Practices: Scope, Organization, personnel- technical competence, desirable qualities of analyst, analyst validation, QBD (quality by design) responsibilities of key personnel in the QC laboratories. Routine controls, instruments, protocols, non-clinical testing, controls on animal house, data generation and storage, quality control documents, retention samples, records, audits of quality control facilities, raw data maintenance.

RECOMMENDED BOOKS:

- 1. The International Pharmacopoeia Vol 1,2,3,4, 3rd Edition General methods of analysis and quality specifications for pharmaceutical substances, excipients, dosage forms.
- 2. Quality Assurance of Pharmaceuticals A compendium of guidelines and related materials Vol.1 and Vol.2, WHO, (1999)
- 3. Basic tests for pharmaceutical substances WHO (1988)
- 4. Basic tests for pharmaceutical dosage forms WHO (1991)
- 5. GMP-Mehra
- 6. How to Practice GMPs P.P.Sharma
- 7. The Drugs and Cosmetic Act 1940 Vijay Malik
- 8. Pharmaceutical Process Validation by Berry and Nash.
- 9. Q.A. Mannual by D.H.Shah
- 10. SOP Guidelines by D.H.Shah
- 11. Quality Assurance Guide by OPPI

Semester: M. Pharmacy 2nd Semester

Subject: Pharmaceutical Quality Assurance-I (Lab)

Total Practical Periods: 72

Total Marks in the End Semester: 100 Minimum of Class Test to be conducted: 2 Branch: **Pharmacy**

Subject Code: 5119221(041)

Total Tutorial Periods: 12

List of experiments:

1. Estimation of multicomponent formulation by UV- Spectrophotometer in formulations. (2 experiments).

- 2. Experiments based on HPLC (Isocratic and Gradient elution) techniques. (2 experiments).
- 3. Interpretation of drugs by IR spectra.
- 4. Workshop of spectroscopy: (UV, IR, NMR, MASS) structural elucidation of at least 5 compounds (4experiments).
- 5. Separation of protein drug substances by electrophoresis.
- 6. Analysis of active pharmaceutical ingredients (API) (2 experiments).
- 7. Identification of impurities and related substances in API's (Albendazole, metronidazole, diclofenac, paracetamol, aspririn, ibuprofen) (2 experiments).
- 8. Use of Super critical fluid chromatography for the analysis of Pharmacopoeial compounds.

Semester: M. Pharmacy 2nd Semester

Subject: Pharmaceutical Quality Assurance-II (Lab)

Total Practical Periods: 72

Total Marks in the End Semester: 100 Minimum of Class Test to be conducted: 2 Total Tutorial Periods: 12

Subject Code: 5119222(041)

Branch: Pharmacy

List of Experiments

1. Complete analysis of formulations, one of each type, as per protocol and comment there on.

- 2. Validation of processes manufacturing & analytical (at least 3 exercises on each)
- 3. Calibration of at least 3 equipments and 3 instruments.
- 4. Cleaning validation of at least 2 equipments.
- 5. Exercises on statistical analysis of data using different models (at least 6 exercises).
- 6. Exercises on determination of (1) order of degradation, (2)shelf-life prediction and(3) interpretation of results, based on given data(arbitrary)(at least 3 exercises).
- 7. Exercises on pharmacological cases (at least 3).

Semester: M. Pharmacy 2nd Semester Branch: Pharmacy

Subject: Pharmaceutical Quality Assurance–III (Lab)

Total Practical Periods: 72

Subject Code: 5119223(041)

Total Tutorial Periods: 12

Total Marks in the End Semester: 100 Minimum of Class Test to be conducted: 2

List of experiments:

- 1. Validation of (analytical) instruments. (IQ,OQ & PQ) (UV, IR, HPLC).
- 2. Validation of analytical methods.
- 3. Standard operating procedure (SOP) for analytical instrumentation.
- 4. Standard operating procedure (SOP) for cleaning validation.
- 5. Standard test procedure (STP) for monograph analysis including COA (certificate of analysis).
- 6. Comparison of methods available in the official methods mentioned in IP, BP, USP etc for various dosage forms.
- 7. Analytical method validation for evaluation of drugs from biological samples.
- 8. Analysis of drugs in biological fluids.
- 9. Cleaning validation method, swab and rinse sample, maximum allowable concentration calculations.



Scheme of Teaching and Examination

Courses of Study and Scheme of Examination of Master of Pharmacy (Pharmaceutical Quality Assurance)

Semester - III

S. No	Board of Study	Subject Code	Subject	Period per Week			Scheme of Examination Theory/Practical			Total Marks	Credit L+(T+P)/ 2
				L	Т	P	ESE	CT	TA		
1.	Pharmacy	5119321 (041)	Minor Dissertation (Synopsis Submission Seminar & Viva)	-	3	36	300	-	100	400	-
	Total				03	36	300	-	100	400	-

L – Lecture,

T – Tutorial,

P - Practical,

ESE – End Semester Examination,

CT – Class Test,

TA- Teacher Assessment



Scheme of Teaching and Examination

Courses of Study and Scheme of Examination of Master of Pharmacy (Pharmaceutical Quality Assurance)

Semester - IV

S. No.	Board of Study	Subject Code	Subject	Period per Week			Scheme of Examination Theory/Practical			Total Marks	Credit L+(T+P)/ 2
	-			L	T	P	ESE	CT	TA		
1	Pharmacy	5119421 (041)	Major Dissertation (Seminar & Viva)	-	03	36	400	-	200	600	-
Total				-	03	36	400	-	200	600	-

L – Lecture,

T – Tutorial,

P – Practical,

ESE – End Semester Examination,

CT – Class Test,

TA- Teacher Assessment