Table - 7: Course of study for M. Pharm. (Regulatory Affairs)

Course Code	Course	Credit Hours	Credit Points	Hrs./ wk	Marks	
	Semester I					
MRA 101T	Good Regulatory Practices	4	4	4	100	
MRA 102T	Documentation and Regulatory Writing	4	4	4	100	
MRA 103T	Clinical Research Regulations	4	4	4	100	
MRA 104T	Regulations and Legislation for Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals In India and Intellectual Property Rights	4	4	a	100	
MRA 105P	Regulatory Affairs Practical I	12	6	12	150	
	Seminar/Assignment	7	4	7	100	
	Total	35	2.6	35	650	
		ester IÍ				
MRA 201T	Regulatory Aspects of Drugs & Cosmetics	(1)	4	4	100	
MRA 202T	Regulatory Aspects of Herbal & Biologicals	4	4	4	100	
MRA 203T	Regulatory Aspects of Medical Devices	4	4	4	100	
MRA 204T	Regulatory Aspects of Food & Nutraceuticals	4	4	4	100	
MRA 205P	Regulatory Affairs Practical II	12	6	12	150	
	Semina./Assignment	7	4	7	100	
,	Total	35	26	35	650	

Table - 12: Course of study for M. Pharm. III Semester (Common for All Specializations)

Course Code	Course	Credit Hours	Credit Points
MRM 301T	Research Methodology and Biostatistics*	4	4
-	Journal club	1	1
-	Discussion / Presentation (Proposal Presentation)	2	2
-	Research Work	28	14
Total		35	21

^{*} Non University Exam

Table - 13: Course of study for M. Pharm. IV Semester (Common for All Specializations)

Course Code	Course		C. edit hours	Credit Points
-	Journal Club		1	1
-	Research Work	6	31	16
-	Discussion/Final Presentation	1019	3	3
	Total		35	20

Table - 14. Semester vise credits distribution

Table - 14. Semester vise credits distribution				
Semester	Credit Points			
I	26			
II	26			
III	21			
IV	20			
Co-curricular Activities (Attending Conference, scientific Presentations and Other Scholarly Activities)	Minimum=02 Maximum=07*			
Total Credit Points	Minimum=95			
	Maximum=100*			
*Credit Points for Co-curricular Activities				
Okalica				

PHARMACEUTICALREGULATORY AFFAIRS(MRA)

GOOD REGULATORY PRACTICES (MRA 101T)

Scope

This course is designed to impart fundamental knowledge on various Good Regulatory Practices viz., cGMP, GLP, GALP and GDP for Pharmaceuticals, Cosmetics, Food & Nutraceuticals, Medical devices, In-vitro Diagnostic Medical Devices (IVDs) and biological products and understand the rationale behind these requirements and will propose ways and means of complying with them.

Objectives

At completion of this course it is expected that students will be able to understand,

- The key regulatory and compliance elements with respect to Good Manufacturing Practices, Good Laboratory Practices, Good Automated Laboratory Practices and Good Documentation Practices.
- Prepare and implement the check lists and SOPs for various Good Regulatory Practices
- Implement Good Regulatory Practices in the Healthcare and related Industries
- Prepare for the readiness and conduct of audits and inspections.

THEORY 60 Hrs

- Current Good Manufacturing Practices: Introduction, US cGMP 12
 Part 210 and Part 21.1.EC Principles of GMP (Directive Hrs 91/356/EEC) Article to Article 14 and WHO cGMP guidelines GAMP-5; Medica device and IVDs Global Harmonization Task Force(GHTF) Guidance docs.
- 2 Good Laboratory Practices: Introduction, USFDA GLP 12
 Regulations (Subpart A to Subpart K), Controlling the GLP Hrs
 inspection process, Documentation, Audit, goals of Laboratory
 Quality Audit, Audit tools, Future of GLP regulations, relevant ISO
 and Quality Council of India(QCI) Standards
- Good Automated Laboratory Practices: Introduction to GALP, 12 Principles of GALP, GALP Requirements, SOPs of GALP, Hrs Training Documentation,21 CFR Part 11, General check list of 21CFR Part 11, Software Evaluation checklist, relevant ISO and OCI Standards.

- 4 Good Distribution Practices: Introduction to GDP, Legal GDP 12 requirements put worldwide, Principles, Personnel, Hrs Documentation, Premises and Equipment, Deliveries to Customers, Returns, Self-Inspection, Provision of information, Stability testing principles, WHO GDP, USP GDP (Supply chain integrity), relevant CDSCO guidance and ISO standards
- Quality management systems: Concept of Quality, Total Quality Management, Quality by design, Six Sigma concept, Out of Specifications (OOS), Change control. Validation: Types of Validation, Types of Qualification, Validation master plan (VMP), Analytical Method Validation. Validation of utilities, [Compressed air, steam, water systems, Heat Ventilation and Air conditioning (HVAC)]and Cleaning Validation. The International Conference on Harmonization (ICH) process, ICH guidelines to establish quality, safety and efficacy of drug substances and products. ISO 13485, Sch MIII and other relevant CDSCO regulatory guidance documents.

REFERENCES

- 1. Good Laboratory Practice Regulations, by Sandy Weinberg, Fourth Edition Drugs and the Pharmaceutical Sciences, Vol.168
- 2. Good Pharmaceutical Manufacturing practice, Rational and compliance by John Sharp, CRC Press
- 3. Establishing a cGMP Laboratory Audit System, A practical Guide by David M.Bleisner, Wiley Publication.
- 4. How to practice GLP by PP Sharma, Vandana Publications.
- 5. Laboratory Auditing for Quality and Regulatory compliance bu Donald C.Singer, Drugs and the Pharmaceutical Sciences, Vol.150.
- 6. Drugs & Cosmetics Act, Rules & Amendments

DOCUMENTATION AND REGULATORY WRITING (MRA 102T)

Scope

This course is designed to impart fundamental knowledge on documentation and general principles involved in regulatory writing and submission to agencies.

Objectives

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

- Know the various documents pertaining to drugs in pharmaceutical industry
- Understand the basics of regulatory compilation
- Create and assemble the regulation submission as per the requirements of agencies
- Follow up the submissions and post approval document requirements

THEORY 60 Hrs

- 1. Documentation in pharmaceutical industry: Exploratory 12 Product Development Brief (EPDB) for Drug substance and Drug Hrs product, Product Development Plan (PD?), Product Development Report (PDR), Master Formula Record, Batch Manufacturing Record and its calculations. Batch Reconciliation, Batch Packaging Records, Print Pack specifications, Distribution records, Certificate of Analysis (CoA), Site Master File and Drug Master Files (DMF).
- 2 Dossier preparation and submission: Introduction 12 overview of dossiers, contents and organization of dossier, Hrs binders and sections, compilation and review of dossier. Paper submissions, overview and modules of CTD, electronic CTD Electronic submission: Planning submission, requirements for submission, regulatory bindings and requirements. Tool and Technologies, electronic submission process and validating the submission, Electronic Submission Gateway (ESG). Non eCTD electronic submissions (NeeS), Asian CTD formats (ACTD) submission. Organizing, process and validation of submission. Submission in Sugam system of CDSCO.

- Audits: Introduction, Definition, Summary, Types of audits, GMP 12 compliance audit, Audit policy, Internal and External Audits, Hrs Second Party Audits, External third party audits, Auditing strategies, Preparation and conducting audit, Auditing strategies, audit analysis, audit report, audit follow up. Auditing/inspection of manufacturing facilities by regulatory agencies. Timelines for audits/inspection. GHTF study group 4 guidance document. ISO 13485.
- 4 Inspections: Pre-approval inspections, Inspection of 1.3 pharmaceutical manufacturers, Inspection of drug distribution in the channels, Quality systems requirements for national good manufacturing practice inspectorates, inspection report, model certificate of good manufacturing practices, Root cause analysis, Corrective and Preventive action (CAPA).
- Product life cycle management: Prior Approval Supplement 12 (PAS), Post Approval Changes [SUPAC], Changes Being Hrs Effected in 30 Days (CBE-30), Annual Report, Post marketing Reporting Requirements, Post approval Labeling Changes, Lifecycle Management, FDA inspection and Enforcement, Establishment Inspection Report (CIR), Warning Letters, Recalls, Seizure and Injunctions. ISO Pisk Management Standard

REFERENCES

- 1. Compliance auditing for Pharmaceutical Manufacturers. Karen Ginsbury and Gi. Bismuth, Interpharm/CRC, Boca Raton, London New York, Washington D.C.
- 2. Pharmaceutical Manufacturing Handbook, Regulations and Quality by Shayne Cox Gad. Wiley-Interscience, A John Wiley and sons, Inc., Publications.
- 3. Handbook of microbiological Quality control. Rosamund M. Baird, Norman A. Hodges, Stephen P. Denyar. CRC Press. 2000.
- 4. Laboratory auditing for quality and regulatory compliance. Donald C. Singer, Raluca-loana Stefan, Jacobus F. Van Staden. Taylor and Francis (2005).
- 5. Implementing Juran's Road Map for Quality Leadership: Benchmarks and Results, By Al Endres, Wiley, 2000
- 6. Understanding, Managing and Implementing Quality: Frameworks, Techniques and Cases, By Jiju Antony; David Preece, Routledge, 2002

- 7. Organizing for High Performance: Employee Involvement, TQM, Reengineering, and Knowledge Management in the Fortune 1000: The CEO Report By Edward E. Lawler; Susan Albers Mohrman; George Benson, Jossey-Bass, 2001
- 8. Corporate Culture and the Quality Organization By James W. Fairfield-Sonn, Quorum Books, 2001
- The Quality Management Sourcebook: An International Guide to Materials and Resources By Christine Avery; Diane Zabel, Routledge, 1997
- 10. The Quality Toolbox, Second Edition, Nancy R. Tague, A.Q. Publications
- 11. Juran's Quality Handbook, Sixth Edition, Joseph M. Juran and Joseph A. De Feo, ASQ Publications
- 12. Root Cause Analysis, The Core of Problem Solving and Corrective Action, Duke Okes, 2009, ASQ Publications
- 13. International Medical Device Regulators Forum (IMDRF) Medical Device Single Audit Program (MDSAP)

CLINICAL RESEARCH REGULATIONS (MRA 103T)

Scope

This course is designed to impart the fundamental knowledge on the clinical development process of drugs, pharmaceuticals and Medical Devices, phases and conduct of clinical trials and research, regulations and guidance governing the conduct of clinical research in India, USA and EU. It prepares the students to learn in detail on various laws, legislations and guidance related to safety, efficacy, ethical conduct and regulatory approval of clinical research.

Objectives

Upon completion of the course, the student shall be able to (know, do and appreciate)

- History, origin and ethics of clinical and biomedical research and evaluation
- Clinical drug, medical device development process and different types and phases of clinical trials
- Regulatory requirements and guidance for conduct of clinical trials and research

Theory 60 Hrs

1. Clinical Drug Development Process 12

• Different types of Clinical Studies Hrs

- Different types of Clinical Studies
- Phases of clinical trials, Chaical Trial protocol
- Phase 0 studies
 Phase 1 and subtype studies (single ascending, multiple ascending, dose escalation, methods, food effect studies, drug drug interaction, PK end points
- Phase I studies (proof of concept or principle studies to establish efficacy)
- Phase III studies (Multi ethnicity, global clinical trial, registration studies)
- Phase IV studies (Post Marketing Studies; PSUR)

Clinical Investigation and Evaluation of Medical Devices & IVDs

Different Types of Studies Key Concepts of Medical Device Clinical Evaluation Key concepts of Clinical Investigation 2 Ethics in Clinical Research:

- 12 Hrs
- Historical Perspectives: Nuremberg Code, Thalidomide study
 , Nazis Trials, Tuskegee Syphilis Study, The Belmont Report,
 The declaration of Helsinki
- Origin of International Conference on Harmonization Good Clinical Practice (ICH-GCP) guidelines.
- The ethics of randomized clinical trials
- The role of placebo in clinical trials
- Ethics of clinical research in special population
- Institutional Review Board/Independent Ethics Committee/Ethics Committee - composition, roles, responsibilities, review and approval process and organic monitoring of safety data
- Data safety monitoring boards.
- Responsibilities of sponsor, CRO, and investigator in ethical conduct of clinical research
 - Ethical principles governing informed consent process
 - Patient Information Sheet and Informed Consent Form
 - The informed consent process and accumentation
- Regulations governing Clinical Trials
 India: Clinical Research regulations in India Schedule Y & Hrs
 Medical Device Guidance

USA: Regulations to conduct drug studies in USA (FDA)

- NDA 505(b)(1) of the FD&C Act (Application for approval of a new drug)
- NDA 505(b)(2) of the FD&C Act (Application for approval of a new drug that relies, at least in part, on data not developed by the applicant)
- AND: 505(j) of the FD&C Act (Application for approval of a generic drug product)
- FDA Guidance for Industry Acceptance of Foreign Clinical
 Studies
- FDA Clinical Trials Guidance Document: Good Clinical Practice

EU: Clinical Research regulations in European Union (EMA)

Good Clinical Practice Guidelines (ICH GCP E6) Hrs Indian GCP Guidelines ICMR Ethical Guidelines for Biomedical Research CDSCO quidelines GHTF study group 5 guidance documents Regulatory Guidance on Efficacy and Safety ICH Guidance's E4 - Dose Response Information to support Drug Registration E7 - Studies in support of General Population: Geriatrics E8 - General Considerations of Clinical Trials E10 - Choice of Control Groups and Related Issues Clinical Trials. E 11 - Clinical Investigation of Medicinal Products in the **Pediatric Population** General biostatics principle applied in clinical research 5 USA & EU Guidance 12 USA: FDA Guidance Hrs CFR 21Part 50: Protection of Human Subjects CFR 21Part 54: Financial Disclosure by Clinical Investigators CFR 21 Part 312: IND Application CFR 21Part 314: Application for FDA Approval to Market a New Drug 320: Bioavailability and bioequivalence CFR 21 Part requirements CFR 21Part 812: Investigational Device Exemptions CFR 21 Part 822. Post-market surveillance FDA Safety Reporting Requirements for INDs and BA/BE Studies FDA Med Watch

12

Clinical Research Related Guidelines

- EudraLex (EMEA) Volume 3 Scientific guidelines for medicinal products for human use

Guidance for Industry: Good Pharmacovigilance Practices

EU Annual Safety Report (ASR)

and Pharmacoepidemiologic Assessment European Union: EMA Guidance FU Directives 2001

- Volume 9A Pharmacovigilance for Medicinal Products for Human Use
- EU MDD with respect to clinical research
- ISO 14155

4

REFERENCES

- 1. Clinical Trials and Human Research: A Practical Guide to Regulatory Compliance By Fay A. Rozovsky and Rodney K. Adams
- 2. HIPAA and Human Subjects Research: A Question and Answer Reference Guide By Mark Barnes, JD, LLM and Jennifer Kulynych, JD, PhD
- 3. Principles and Practices of Clinical Research, Second Edition Edited by John I. Gallin and Frederick P. Ognibene
- Reviewing Clinical Trials: A Guide for the Ethics Committee; Johan PE Karlberg and Marjorie A Speers; Karlberg, Johan Petter Einar, Hong Kong.
- 5. International Pharmaceutical Product Registration: Aspects of Quality, Safety and Efficacy; Anthony C. Cartwright; Taylor & Francis Inc. USA.
- 6. New Drug Approval Process: The Global Challenge; Guarino Richard A; Marcel Dekker Inc., NY.
- 7. FDA regulatory affairs: a guide for prescription drugs, medical devices, and biologics; Douglas J. Pisano, David Mantus; CRC Press, USA
- 8. Country Specific Guidelines from official websites.
- 9. Drugs & Cosmetics Act & Rules and Amendments

RECOMMENDED WEBSITES:

- 1. EU Clinical Research Directive 2007: http://www.eortc.be/services/doc/clinical-eudirective-04-april-01.pdf
- 2. Code of Federal Regulations, FDA: http://www.accessdata.fda.gov/_cripts/cdrh/cfdocs/cfcfr/cfrsearch.cfm
- 3. Guidelines of International Conference on Harmonization: http://www.ich.org/products/guidelines.html
- 4. Eudralex Guidelines: http://www.gmpcompliance.info/euguide.htm
- 5. FDA New Drug Application:
- http://www.fda.gov/regulatoryinformation/legislation/FederalFoodDruga ndCosmetic ActFDCAct/FDCActChapterVDrugsandDevices/ucm108125.htm
- 7. Medicines and Healthcare products Regulatory Agency: http://www.mhra.go.guk
- 8. Central Drugs Standard Control Organization Guidance for Industry: http://cdsco.nic.in/CDSCO-GuidanceForIndustry.pdf
- 9. ICMR Ethical Guidelines for Biomedical Research: http://icmr.nic.in
 http://icmr.nic.in
 http://icmr.nic.in

REGULATIONS AND LEGISLATION FOR DRUGS & COSMETICS, MEDICAL DEVICES, BIOLOGICALS & HERBALS, AND FOOD & NUTRACEUTICALS IN INDIA AND INTELLECTUAL PROPERTY RIGHTS

(MRA 104T)

Scope

This course is designed to impart fundamental knowledge on regulations and legislation in India w.r.t. Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals. It prepares the students for basic regulatory requirements in India of Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals. for manufacture, import & registration, export, sale, marketing authorization, clinical trials and intellectual property rights.

Objectives

Upon the completion of the course the student shall be able to:

- Know different Acts and guidelines that regulate Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals industry in India.
- Understand the approval process and regulatory requirements for Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals

THEORY 60 Hrs

- Biologicals & Herbals, and Food & Nutraceuticals
 Acts and Rules (with latest amendments):
 Hrs
 - Drugs and Cosmetics Act 1940 and Rules 1945: DPCO and NRFA
 - 2. Other relevant provisions (rules schedules and guidelines for approval of Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals in India

Other relevant Acts: Narcotics Drugs and Psychotropic Substances Act; Medicinal and Toilet Preparations (Excise Duties) Act, 1955; Pharmacy Act, 1948; Drugs and Magic Remedies (Objectionable Advertisements) Act, 1955; Prevention of Cruelty to Animals Act.

Regulatory requirements and approval procedures for Drugs & Cosmetics Medical Devices, Biologicals & Herbals, and Hrs Food & Nutraceuticals

CDSCO (Central Drug Standard Control Organization) and State Licensing Authority: Organization, Responsibilities

- Rules, regulations, guidelines and standards for regulatory filing of Drugs & Cosmetics, Medical Devices, Biologicals & Herbals, and Food & Nutraceuticals
- Format and contents of Regulatory dossier filing Clinical trial/investigations
- 3 Indian Pharmacopoeial Standards, BIS standards and ISO and 12 other relevant standards

 Hrs
- 4 Bioavailability and Bioequivalence data (BA &bF), BCS 12 Classification of Drugs, Regulatory Requirements for Hrs Bioequivalence study Stability requirements: ICH and WHO

Guidelines for Drug testing in animals/Freclinical Studies

Animal testing: Rationale for conducting studies, CPCSEA Guidelines
Ethical guidelines for human participants
ICMR-DBT Guidelines for Stem Cell Research

5 Intellectual Property Rights: Patent, Trademark, Copyright, 12 Industrial Designs and Geographical Indications, Indian Patent Hrs Scenario. 128 vs Regulatory Affairs

REFERENCES

- Manual of Patent Practice & Procedure, 3rd Edition, by The Patent Office of India
- 2. Patent Failure How Judges, Bureaucrats, and Lawyers put innovators at risk by James Bessen and Michael J. Meurer
 - Principles and Practice of Clinical Trial Medicine by Richard Chin and Bruce Y. Lee
 - 4. Ethical Guidelines for Biomedical Research on Human Participants by Indian Council of Medical Research New delhi 2006.
 - 5. CPCSEA Guidelines for Laboratory Animal Facility by Committee for the purpose of control and supervision on experiments on animals (CPCSEA)

- 6. ICH E6 Guideline Good Clinical Practice|| by ICH Harmonised Tripartite
- 7. Guidance for Industry on Submission of Clinical Trial Application for Evaluating Safety and Efficacy by CDSCO (Central Drug Standard Control Organisation)
- al .als | JSCO GERMANTIACS, SCO GERMANTI 8. Guidance for Industry on Requirement of Chemical & Pharmaceutical Information including Stability Study Data before approval of clinical trials /

REGULATORY AFFAIRS PRACTICAL - I

(MRA 105P)

- 1. Case studies (4 Nos.) of each of Good Pharmaceutical Practices.
- 2. Documentation for in process and finished products Quality control tests for Solid, liquid, Semisolid and Sterile preparations.
- 3. Preparation of SOPs, Analytical reports (Stability and validation)
- Protocol preparation for documentation of various types of records (BMR, MFR, DR)
- 5. Labeling comparison between brand & generics.
- 6. Preparation of clinical trial protocol for registering trial in India
- 7. Registration for conducting BA/BE studies in India
- 8. Import of drugs for research and developmental activities
- 9. Preparation of regulatory dossier as per Indian CTD forma* a 10 submission in SUGAM
- 10. Registering for different Intellectual Property Rights in India
- 11. GMP Audit Requirements as per CDSCO
- 12. Preparation and documentation for Indian Patent application.
- 13. Preparation of checklist for registration of IND as per ICH CTD format.
- 14. Preparation of checklist for registration of NDA as per ICH CTD format.
- 15. Preparation of checklist for registration of Al/DA as per ICH CTD format.
- 16. Case studies on response with scientific rationale to USFDA Warning Letter
- 17. Preparation of submission checklist of IMPD for EU submission.
- 18. Comparison study of marketing cuthorization procedures in EU.
- 19. Comparative study of DMF system in US, EU and Japan
- 20. Preparation of regulatory submission using eCTD software
- 21. Preparation of Clinical Trial Application (CTA) for US submission
- 22. Preparation of Clinical Trial Application (CTA) for EU submission
- 23. Comparison of Cinical Trial Application requirements of US, EU and Japan of a dosage form
- 24. Regulatory requirements checklist for conducting clinical trials in India.
- 25. Regulatory requirements checklist for conducting clinical trials in Europe.
- 26. Regulatory requirements checklist for conducting clinical trials in USA