## GUJARAT TECHNOLOGICAL UNIVERSITY M.Pharm PHARMACEUTICAL REGULATORY AFFAIRS SEMESTER: I

## Subject Name: GOOD REGULATORY PRACTICES Subject Code: MRA101T

**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**: Upon completion of this course the student should be able to

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

Sr	Course Contents	Total Hrs
No		
1	Current Good Manufacturing Practices: Introduction, US Cgmp Part 210 and	12
	Part 211.EC Principles of GMP (Directive 91/356/EEC) Article 6 to Article	
	14 and WHO cGMP guidelines GAMP-5; Medical device and IVDs Global	
	Harmonization Task Force(GHTF) Guidance docs	
2	Good Laboratory Practices: Introduction, USFDA GLP Regulations (Subpart	12
	A to Subpart K), Controlling the GLP inspection process, Documentation,	
	Audit, goals of Laboratory Quality Audit, Audit tools, Future of GLP	
	regulations, relevant ISO and Quality Council of India(QCI) Standards	
3	Good Automated Laboratory Practices: Introduction to GALP, Principles of	12
	GALP, GALP Requirements, SOPs of GALP, Training Documentation,21	
	CFR Part 11, General check list of 21CFR Part 11, Software Evaluation	
	checklist, relevant ISO and QCI Standards.	
4	Good Distribution Practices: Introduction to GDP, Legal GDP requirements	12
	put worldwide, Principles, Personnel, 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	
5	Quality management systems: Concept of Quality, Total Quality	12
	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