LJ UNIVERSITY

LJ INSTITUTE OF PHARMACY

SEMESTER: VII

Subject Name: Industrial Pharmacy-II

Subject Code: BP702TT

Scope: This course is designed to impart fundamental knowledge on pharmaceutical product development and translation from laboratory to market.

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

- 1. Know the process of pilot plant and scale up of pharmaceutical dosage forms
- 2. Understand the process of technology transfer from lab scale to commercial batch
- 3. Know different Laws and Acts that regulate pharmaceutical industry
- 4. Understand the approval process and regulatory requirements for drug products.

Teaching scheme and examination scheme:

Teaching Scheme				Evaluation Scheme			
Theory	Tutorial	Practical	Total	Theory		Practical	
				External	Internal	External	Internal
3	1	0	4	75	25	0	0

	Course Contents	Hours				
P	Pilot plant scale up techniques: General considerations – including	1 0				
	significance of personnel requirements, space requirements, raw materials, Pilot plant scale up					
	considerations for solids, liquid orals, semi solids and relevant documentation, SUPAC					
$\overline{}$	guidelines, Introduction to platform technology					
	Technology development and transfer: WHO guidelines for Technology	1				
	Fransfer(TT): Terminology, Technology transfer protocol, Quality risk management,	0				
	Fransfer from R & D to production (Process, packaging and					
	cleaning), Granularity of TT Process (API, excipients, finished products,					
	backaging materials) Documentation, Premises and equipments, qualification and validation,					
	quality control, analytical method transfer, Approved regulatory bodies and agencies,					
	Commercialization - practical aspects and					
•	problems (case studies), TT agencies in India - APCTD, NRDC, TIFAC, BCIL, TBSE/SIDBI;					
	TT related documentation - confidentiality agreement, licensing,					
_	MoUs, legal issues	1				
	Regulatory affairs: Introduction, Historical overview of Regulatory Affairs,	0				
	Regulatory authorities, Role of Regulatory affairs department, Responsibility of	U				
	Regulatory Affairs Professionals Regulatory requirements for drug approval: Drug Development Teams, Non-Clinical					
	Drug Development, Pharmacology, Drug Metabolism and Toxicology, General considerations					
	of Investigational New Drug (IND) Application, Investigator's Brochure (IB) and New Drug					
	Application (NDA), Clinical research / BE studies, Clinical Research Protocols, Biostatistics					
	n Pharmaceutical Product Development, Data Presentation for FDA Submissions,					
	Management of Clinical Studies.					
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	Quality management systems: Quality management & Certifications: Concept	8			
4.	of Quality, Total Quality Management, Quality by Design (QbD), Six Sigma concept, Out				
	of Specifications (OOS), Change control, Introduction to ISO 9000				
	series of quality systems standards, ISO 14000, NABL, GLP				
5.	Indian Regulatory Requirements: Central Drug Standard Control	7			
	Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities,				
	Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval				
	procedures for New Drugs.				
Total hours					

Recommended Books (Latest Editions)

- 1. Regulatory Affairs from Wikipedia, the free encyclopedia modified on 7th April available at http,//en.wikipedia.org/wiki/Regulatory_ Affairs.
- 2. International Regulatory Affairs Updates, 2005. available at http://www.iraup.com/about.php
- 3. Douglas J Pisano and David S. Mantus. Text book of FDA Regulatory Affairs A Guide for Prescription Drugs, Medical Devices, and Biologics' Second Edition.
- 4. Regulatory Affairs brought by learning plus, inc. available at http://www.cgmp.com/ra.htm.