Seat No.:

Enrolment No.\_\_\_\_\_

Date: 09-01-2015

## **GUJARAT TECHNOLOGICAL UNIVERSITY**

## M. Pharm. – SEMESTER – I • EXAMINATION – WINTER • 2014

Subject Code: 910204

## Subject Name: Good Manufacturing and Good Laboratory Practice **Total Marks: 80**

Time: 10.30 am - 01.30 pm

**Instructions:** 

- 1. Attempt any five questions.
- 2. Make suitable assumptions wherever necessary.
- 3. Figures to the right indicate full marks.

Q.1	(a)	Define the terms GMP, Quality control & Quality assurance and explain their interrelationship.	06
	<b>(b</b> )	Enumerate GLP guidelines practiced in India and briefly explain scope of GLP activities.	05
	(c)	Enumerate different GMP guidelines for Drug Substance and Drug product.	05
Q.2	(a)	Define the followings in relation to pharmaceutical manufacturing facility: Plant Layout, Environment Control, Clean Area, Sanitation, Cross Contamination, Utilities.	06
	(b) (c)	Explain the terms Clean-in-place (CIP) and Sterilize-in-place (SIP). Enumerate different documents related to production of drugs and enlist in- process tests for tablet dosage form.	05 05
Q.3	(a)	What is meant by SOP? Write names of five SOPs used in production department for different unit operations and describe any one of them.	06
	(b)	Explain Line clearance and Reconciliation of printed packaging materials in relation to Packaging and labeling operations.	05
	(c)	Explain role of production, QC and QA in batch release procedure.	05
Q.4	(a)	Define the followings with respect to QC operations : Sampling, Standard Test Procedure, Retention Samples.	06
	(b) (c)	Enlist various sections of warehouse and Good warehouse practices. Write a note on Vendor approval.	05 05
Q.5	(a)	Describe the objective, scope and procedure for WHO certification scheme for pharmaceutical products.	06
	(b) (c)	Write a note on self-inspection and quality audit. Give specifications for any one raw material and work-in-process material.	05 05
Q. 6	(a)	Define the following terms in relation to GLP: Sponsor, Quality Assurance Unit, Animal care, Protocol for conduct of study.	06
	(b) (c)	How will you determine the specifications of Non-pharmaceutical product? Write a note on complaint and recall.	05 05
Q.7	(a) (b) (c)	Write different tests and methods for testing of primary packing materials. Enlist different documents related to QC laboratory testing. Explain importance and different types of Training imparted to an employee in pharmaceutical plant.	06 05 05

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