Seat No.:	Enrolment No
GUJARAT TECHNOLOGICAL UNIVERSITY M. Pharm - SEMESTER-III • EXAMINATION – SUMMER-2016	

Date: 27/05/2016

Subject Name: Drug regulation and Regulatory Authority Time: 2:30 PM to 5:30 PM **Total Marks: 80** 

**Instructions:** 

Subject Code: 1931501

1. Attempt any five questions.

- 2. Make suitable assumptions wherever necessary.
- 3. Figures to the right indicate full marks.

Q.1	(a) (b) (c)	Explain in detail Drug Master File. Write a note on medical device registration as per 510k. Explain the format and contents of NDA.	06 05 05
Q.2	(a) (b) (c)	Write a note on WHO certification scheme on Quality of Pharmaceutical products.  Explain the Procedure for price fixing of Drug substance and Drug Product.  Write a note on National Formulary of India.	06 05 05
Q.3	(a) (b) (c)	Explain the Standard format of Monograph as per IP. Give Emphasis on IP Review Process. Write a note on Site Master File.	06 05 05
Q.4	(a) (b) (c)	Write a note on GCP as per US.  Define Pharmacovigilance. Explain ADR reporting system in India and US.  Write a note on Toxicological studies as per ICH guidelines.	06 05 05
Q.5	(a) (b) (c)	Classify medical device. Explain in detail PMA process in detail.  Define Haemovigilance. Explain Adverse event reporting system as per India.  Explain labeling requirements for drugs as per D &C act.	06 05 05
Q. 6	(a) (b) (c)	Explain Objectives and functions of CDL and DTAB. Write a note on NDA as per EUDRA guidelines. Explain the process involved in Harmonization of Pharmacopieal standards.	06 05 05
Q.7	(a) (b) (c)	Explain the clinical evaluation process as per US and Europe. Explain WHO guidelines with emphasis on International registration. Write a note on electronic CTD.	06 05 05

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