Seat No.: Enrol	ment No.
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## GUJARAT TECHNOLOGICAL UNIVERSITY

## M. Pharm. - SEMESTER- III • EXAMINATION – SUMMER-2016

Subject Code:1931601 Date: 27/05/2016 Subject Name: Regulatory Affairs-II Time: 2:30 PM to 5:30 PM Total Marks: 80

**Instructions:** 

1.	Attempt	anv	five	questions.

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

(a)	Discuss the regulatory aspects that affect drug product design, manufacture and distribution in Brazil			
<b>(b)</b>	What are Bolar exemptions? Give a brief account on recent developments	05		
(c)	Write a note on: Hatch Waxman Act.	05		
(a)	Discuss in brief the regulatory requirements for pharmaceutical and bulk drug manufacture.	06		
<b>(b)</b>	Explain the procedures for certification and licensing of a pharmaceutical product.	05		
<b>(c)</b>	Give a brief account on importance of regulatory drug analysis.	05		
(a)	Explain herbal drug regulations in India. State the differences of the same from the regulatory aspects of other countries.	06		
<b>(b)</b>	Which are the most recent regulations laid for the contract manufacturing in India?	05		
(c)	Discuss in brief the most recent amendments to Drugs and Cosmetics Act, 1940.	05		
(a)	Describe in brief the disrtibution records, batch release documents and complaints & recalls documents.	06		
(b) (c)	What are the quality and safety aspects for the cosmetic products? Write a note on: BOP Analysis.	05 05		
(a) (b)	Explain PCT. Discuss the advantages and procedure of PCT. What is the importance of investigator's brochure? Give a brief outline of	06 05		
(c)	clinical research protocols.  Discuss the salient features of FDA guidelines for clinical trials in India.	05		
(a)	Write a note on specific requirements, content and format of NDA.	06		
(b) (c)	Discuss the new trends in patenting biotechnology based products.  Give a brief outline of the procedure for importing and exporting the pharmaceutical goods.	05 05		
(a)	Discuss the different factors affecting the international business environment.	06		
(b) (c)	Explain in brief exporting to US and prelitigation considerations.  Write a note on: Effective Pharma Patent Drafting.	05 05		
	(b) (c) (a) (b) (c)	distribution in Brazil.  (b) What are Bolar exemptions? Give a brief account on recent developments relating to the bolar exemptions.  (c) Write a note on: Hatch Waxman Act.  (a) Discuss in brief the regulatory requirements for pharmaceutical and bulk drug manufacture.  (b) Explain the procedures for certification and licensing of a pharmaceutical product.  (c) Give a brief account on importance of regulatory drug analysis.  (a) Explain herbal drug regulations in India. State the differences of the same from the regulatory aspects of other countries.  (b) Which are the most recent regulations laid for the contract manufacturing in India?  (c) Discuss in brief the most recent amendments to Drugs and Cosmetics Act, 1940.  (a) Describe in brief the disrtibution records, batch release documents and complaints & recalls documents.  (b) What are the quality and safety aspects for the cosmetic products?  (c) Write a note on: BOP Analysis.  (a) Explain PCT. Discuss the advantages and procedure of PCT.  (b) What is the importance of investigator's brochure? Give a brief outline of clinical research protocols.  (c) Discuss the salient features of FDA guidelines for clinical trials in India.  (a) Write a note on specific requirements, content and format of NDA.  (b) Discuss the new trends in patenting biotechnology based products.  (c) Give a brief outline of the procedure for importing and exporting the pharmaceutical goods.  (a) Discuss the different factors affecting the international business environment.  Explain in brief exporting to US and prelitigation considerations.		

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