Seat No.: Enrolment No GUJARAT TECHNOLOGICAL UNIVERSITY	
Subject Code: 930104 Date: 11-12-	
J J	2013
Subject Name: Validation & Product Development	0.0
Time: 10.30 am - 01.30 pm Total Mark	s: 80
Instructions:	
1. Attempt any five questions.	
2. Make suitable assumptions wherever necessary.	
3. Figures to the right indicate full marks.	
Q-1 (a) Define Process Validation, discuss its options & explain its importance in	
Pharmaceutical industry.	(6)
(b) Explain the significance of instrument validation & describe the procedure	
For validation of UV-Visible spectrophotometer.	(5)
(c) Discuss the Regulatory Perspectives of analytical method validation and	
Discuss the type of analytical procedures to be validated.	(5)
Q-2 (a) Explain in detail, the Cause & Effect diagram for process of coated tablet	
Manufacturing.	(6)
(b) What is SOP? What is its importance? Describe the SOP for Dissolution Te	• •
Apparatus.	(5)
(c) Explain the importance & advantages of Vendor Certification.	(5)
(c) Explain the importance & advantages of vendor certification.	(\mathbf{J})
Q-3 (a) What is Cleaning Validation? Explain in detail the general procedure for	
Cleaning validation.	(6)
(b) Discuss in detail, the methods to determine Accuracy & Precision of any	(0)
New Instrumental analytical method.	(5)
(c) What is Protocol? Give the general protocol format for Prospective Process	(5)
Validation.	(5)
	(0)
Q-4 (a) Mention the types of water used in Pharmaceutical Industry & Explain the	
Validation of Water for Injection plant.	(6)
(b) Give the flow - chart for Retrospective Process Validation.	(5)
(c) Give a brief note on precautions to be taken while performing Scale-up	
Operation.	(5)
Q-5 (a) What is Calibration? What is its importance? Give the Master Plan for	
Calibration of HPLC system.	(6)
(b) Give a brief note on Software System Validation.	(5)
(c) Enlist the differences between Validation Protocol & Validation Master	
Plan.	(5)
Q-6 (a) Explain in detail IQ,OQ and PQ for Hot Air Oven.	(6)
(b) Describe the Validation Master Plan for Injectable Preparations.	(5)
(c) Enlist the differences between Validated & Non-validated manufacturing	(\mathbf{J})
Process.	(5)
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Q-7 (a) Give detailed explanation about Product Development steps for Sustained	
Release tablet formulation.	(6)
(b) Draw the detailed Cause & Effect diagram for Ointment manufacturing.	(5)
(c) Write a short-note on Aseptic area validation.	(5)