Seat No.:	Enrolment No.

GUJARAT TECHNOLOGICAL UNIVERSITY

B.Pharm – SEMESTER - VII • EXAMINATION – SUMMER-2017

Subj	ect N	Code: 2270015 Name: Quality by Design (QbD) and Process Analytical Technology (PAT) 230 PM to 05:30 PM Total Marks: 80	
1. 2. 3.	Atte Mak	mpt any five questions. The suitable assumptions wherever necessary. The suitable assumptions wherever necessary. The suitable assumptions wherever necessary.	
Q.1	(a) (b) (c)	Define: Quality by Design. Explain the need of QbD for Pharma sector. Explain the QTPP (quality target product profile) parameter with example. Draw a flow chart of quality risk management process.	06 05 05
Q.2	(a) (b) (c)	Explain the following terminology: a) CQA (Critical Quality Attributes) b) CPP (Critical Process Parameter) Discuss advantage and limitation of QbD. Discuss Scope and principles of PAT.	06 05 05
Q.3	(a) (b) (c)	Classify the Optimization techniques. Compare the Traditional and QbD approach for Pharmaceutical product supply. Explain the Clinical Study Reports of CTD.	06 05 05
Q.4	(a) (b) (c)	Explain the elements of QbD. Explain the need of optimization for Pharmaceutical development. Discuss challenges for implementation of QbD.	06 05 05
Q.5	(a) (b) (c)	Define: a) Design Space b) Level c) Factor Explain scope and principle of Quality Risk Management. Explain: Failure Mode Effects Analysis (FMEA).	06 05 05
Q. 6	(a) (b) (c)	Explain Yate's method for optimization with example. Explain process control tool for PAT. Enlist the different parts of CTD.	06 05 05
Q.7	(a)	Discuss Quality target product profile with respect to Modified release dosage form.	06

Discuss management responsibility for Pharmaceutical Quality Management.

05

05

Discuss Scope and principles of PAT.

(b)

(c)