

GUJARAT TECHNOLOGICAL UNIVERSITY**B. PHARM. - SEMESTER – VIII EXAMINATION – OCTOBER 2012****Subject code: 280001****Date: 25/10/2012****Subject Name: Dosage Form Design-II****Time: 02.30pm - 05.30pm****Total Marks: 80****Instructions:**

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

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| Q.1 | (a) | What criteria are necessary for selection of a drug as candidate for formulations of controlled release form? Explain giving optimum ranges for the Bio-pharmaceutics and pharmacokinetics parameters of the drug. | 06 |
| | (b) | What is the major objective of controlled drug delivery system? Give advantages and disadvantages of such a system. | 05 |
| | (c) | Explain loading dose and maintenance dose used in controlled release formulation. | 05 |
| Q.2 | (a) | Discuss the formulation of floating drug delivery systems | 06 |
| | (b) | Explain formulation of different types of trans dermal drug delivery system. | 05 |
| | (c) | Write a note on colon targeted drug delivery system. | 05 |
| Q.3 | (a) | Explain the method of residuals for the calculation of absorption rate constant from oral data. | 06 |
| | (b) | What are pharmacokinetic models? Explain in detail compartment models. | 05 |
| | (c) | Define and explain extraction ratio and discuss hepatic clearance in detail. | 05 |
| Q.4 | (a) | Enlist different methods for determination of pharmacokinetics parameters from urinary excretion data and explain any one method in detail. | 06 |
| | (b) | Define clinical pharmacokinetics and explain dosage adjustment in patients with renal failure. | 05 |
| | (c) | Explain pharmacokinetic drug interactions giving suitable examples. | 05 |
| Q.5 | (a) | Discuss the formulation of parenteral emulsions and suspensions | 06 |
| | (b) | Write a note on implantable osmotic drug delivery system | 05 |
| | (c) | How are liposomes classified? Why are they considered versatile carriers for parenteral drug delivery. | 05 |
| Q. 6 | (a) | Explain lag time, burst effect and reservoir systems with respect to controlled release formulations | 06 |
| | (b) | Explain Hixson and Crowell's cube root law of dissolution. | 05 |
| | (c) | Write a note on dissolution and diffusion controlled release system | 05 |
| Q. 7 | (a) | Explain how one can detect nonlinear pharmacokinetics? Explain Michaelis-Menten equation for capacity limited process. | 06 |
| | (b) | Explain Wagner-Nelson method in detail. | 05 |
| | (c) | Write a note on formulation of trans mucosal drug delivery system | 05 |