

**B. PHARM.
SIXTH SEMESTER
BIOPHARMACEUTICS & PHARMACOKINETICS
BP604T [REPEAT]**

**SET
A**

Duration : 3 hrs.

Full Marks : 75

(PART-A: Objective)

Time : 30 min.

Marks : 20

Choose the correct answer from the following:

1×20=20

- Uphill transport is commonly known as:
 - Active transport
 - Passive transport
 - Pore transport
 - Ion-pair transport
- Duration of washout period for crossover design is:
 - 4 weeks
 - 1 month
 - 2 months
 - 1 week
- _____ is the organ that mainly comprises Peripheral compartment in Two Compartment model:
 - Kidney
 - Muscles
 - Liver
 - Lungs
- An example of Permeation enhancers used in Blood-Brain Barrier is"
 - Mannitol
 - Dihydropyridine
 - DMSO
 - Immunoglobulins
- The most frequently used Compartment model is:
 - Physiological model
 - Mammillary model
 - Catenary model
 - Distribution Parameter model
- Pharmacokinetic methods of Bioavailability measurement involves which studies:
 - Plasma level-time studies
 - Urinary excretion studies
 - Both (a) & (b)
 - Therapeutic studies
- Line-Weaver-Burke Plot is also known as:
 - Scatchard Plot
 - Klotz Plot
 - Hitchcock Plot
 - Direct Plot
- Elimination Half life is also known as:
 - Renal clearance
 - Rate constant
 - Plasma clearance
 - Biological half life
- The unit of C_{max} is expressed in:
 - mcg/ml
 - mg
 - mg/min
 - µg
- Surface Renewal Theory is also known as:
 - Film Theory
 - Interfacial Barrier model
 - Limited Solvation Theory
 - Danckwert's Model

(PART-B :Descriptive)

Time : 2 hrs. 30 min.

Marks : 35

[Answer any seven (7) questions]

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|---|-------------|
| 1. Explain about Kinetics of Protein Binding with proper graphs. | 5 |
| 2. Explain about Two Compartment IV Infusion Open Model | 5 |
| 3. What is Bioavailability? What are the Pharmacodynamic methods of Bioavailability measurement? | 5 |
| 4. What are the causes of Non linearity in Drug Absorption? | 5 |
| 5. What is Compartment analysis? Discuss about 5 advantages of Compartment modeling. | 1+4=5 |
| 6. Discuss One Compartment Open Model IV Bolus for estimation of Pharmacokinetic parameters. | 5 |
| 7. Discuss about any 5 Patient related factors influencing Drug Absorption. | 5 |
| 8. What is IVIVC? What are the levels in IVIVC? | 2+3=5 |
| 9. What is Pharmacokinetics? Discuss about the Pharmacokinetic Parameters with proper explanation of Plasma Drug Concentration Time Graph | 1+3+1
=5 |

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PART-C: Long type questions

[Answer any two (2) questions]

1. Discuss about Michaelis-Menten equation. Give a detailed explanation about the different methods of estimation of K_m and V_{max} . 10
2. Discuss in details about 10 methods to enhance Bioavailability. 10
3. What is Drug Absorption? Describe in details about the mechanisms of Drug Absorption with proper diagram. 1+9=10

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