



BRAINWARE UNIVERSITY

Term End Examination 2021 - 22

Programme – Bachelor of Pharmacy

Course Name – Biopharmaceutics and Pharmacokinetics

Course Code - BP604T

(Semester VI)

Time allotted : 1 Hrs.30 Min.

Full Marks : 75

[The figure in the margin indicates full marks.]

Group-A

(Multiple Choice Type Question)

1 x 75=75

Choose the correct alternative from the following :

- (1) What are the characteristics of continuous release systems?

a) Release the drug along the entire length of GIT	b) Prolonged their residence in the GIT and release
c) Release only at a specific drug	d) Release as soon as comes in contact to the saliva
- (2) What is the characteristic of dissolution for controlled release systems?

a) Release the drug along the entire length of GIT	b) Prolonged their residence in the GIT and release
c) Release only at a specific drug	d) Very slow dissolution rate
- (3) What is the characteristic of encapsulation or coating dissolution-controlled release systems?

a) Microencapsulation using slowly dissolving materials	b) Prolonged their residence in the GIT and release
c) Release only at a specific drug	d) Employ waxes to control the rate of dissolution
- (4) What are the characteristics of diffusion-controlled release systems?

a) Release the drug along the entire length of GIT	b) Diffusion of the dissolved drug
c) Release only at a specific drug	d) Employ waxes to control the rate of dissolution
- (5) What are the characteristics of Matrix diffusion-controlled release systems?

a) Release the drug along the entire length of GIT	b) Drug disperse in an insoluble matrix of rigid hydrophobic materials
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- c) Release only at a specific drug
- d) Employ waxes to control the rate of dissolution
- (6) What are the characteristics of reservoir devices-controlled release systems?
- a) Release the drug along the entire length of GIT
- b) Drug disperse in the insoluble matrix of rigid hydrophobic materials
- c) Hollow systems containing drug surrounded by a polymer membrane
- d) Employ waxes to control the rate of dissolution
- (7) What is the characteristic of pH-independent formulations?
- a) Buffering agents that adjust pH to the desired value
- b) Drug disperse in the insoluble matrix of rigid hydrophobic materials
- c) Hollow systems containing drug surrounded by a polymer membrane
- d) Formation of complexes between the drug and anion/cation exchange resins
- (8) What is the driving force for Passive Diffusion?
- a) Concentration gradient only
- b) Electrochemical gradient only
- c) Charge equilibration and concentration gradient
- d) Concentration and Electrochemical gradient
- (9) What is the driving force of pore transport?
- a) Hydrostatic pressure
- b) Concentration gradient
- c) Electrochemical gradient
- d) Charge equilibration
- (10) What will be the best definition for “carrier”?
- a) Nonpolar drugs can be transported through carrier-mediated transport
- b) Carrier binds reversibly and not covalently with the molecules
- c) It discharges the molecules and gets destroyed itself
- d) The carrier is a protein
- (11) What influences the permeation of drugs in an Ionic or Electrochemical diffusion?
- a) Charge on the membrane
- b) Charge on the particle
- c) Concentration gradient
- d) Equilibration of charge
- (12) What is the major difference between facilitated diffusion and passive diffusion?
- a) Carrier-mediated transport
- b) Downhill transport
- c) Energy is used
- d) Inhibition by metabolic poisons
- (13) Which drugs are absorbed through pore transport?
- a) High lipophilicity
- b) Water-soluble drugs of molecular weight less than 100 Dalton
- c) Oily droplets
- d) Affinity for carriers
- (14) Which of these absorption methods involves engulfing of the extracellular drug?
- a) Endocytosis
- b) Passive diffusion
- c) Facilitated diffusion
- d) Ion-pair transport
- (15) What is the other name of “cell eating”?
- a) Transcytosis
- b) Phagocytosis
- c) Pinocytosis
- d) Endocytosis
- (16) Proteins interact with which part of the cell membrane?
- a) Hydrophobic tail
- b) Polar head
- c) Non polar head
- d) Hydrophilic tail
- (17) What helps in the passing of inorganic ions?

- a) Ion channels
c) Aqueous filled pores
- b) Voltage gated channels
d) Diffusion
- (18) The cell membrane is _____ in nature.
- a) Impermeable
c) Permeable
- b) Semipermeable
d) Permeable to only gases
- (19) What is the most important characteristic of a drug to be absorbed after oral administration?
- a) Dissolved in HCL
c) Can pass through the cell membrane
- b) Dissolved in alkaline solution
d) Form aggregate and settle down
- (20) Gastrointestinal route is an example of which of the major drug delivery routes?
- a) The enteral route
c) The topical route
- b) The parenteral route
d) The intravenous route
- (21) The onset of drug action depends on the rate of:
- a) Drug absorption
c) pH
- b) Drug dissociation
d) GI motility
- (22) Movement of ions through the pores in cell membrane can be controlled by-
- a) Counter ion transport
c) Both a & b
- b) Expenditure of intracellular energy
d) None of these
- (23) What happens when an obese person is given with a lipophilic drug?
- a) Drug aggregation will begin
c) High adipose tissue take up most of the lipophilic drug
- b) He cannot absorb lipophilic drugs
d) A large amount of drug is needed as the person's weight is more
- (24) Who has poorly developed BBB?-
- a) Infants
c) Aged
- b) Adults Of age more than 20
d) Children at puberty
- (25) What should be the molecular weight of the drug molecules so that they can easily pass through the membrane?
- a) 600-800 Dalton
c) 300-500 Dalton
- b) 500-600 Dalton
d) 200-400 Dalton
- (26) Which type of drug cannot enter the cell membrane in the below picture?
- a) Ionized drug
c) Hydrolyzed drug
- b) Unionized drug
d) Unhydrated drug
- (27) Which drugs cannot pass the capillary endothelial barrier?
- a) Molecular size less than 600 Dalton
c) Drugs bound to a chemical moiety
- b) Drugs bound to blood components
d) All drugs can pass
- (28) Which of the following drug cannot pass through the plasma membrane barrier?
- a) Drug size less than 50 Dalton
c) Polar or ionized drugs of size greater than 50 Dalton
- b) Lipophilic drugs 50-600 Dalton
d) Drug size more than 600 Dalton
- (29) What is the name of the specialized cells that support the blood-brain barrier tissue?
- a) Astrocytes
c) Fat cells
- b) Dendrites
d) Endothelial cells
- (30) Why dopamine cannot be administered for the disease parkinsonism?

- a) Don't have a medicine
b) It is not the medicine
c) Cannot cross the blood-brain barrier
d) Forms aggregate and thus cannot cross the BBB
- (31) In equation, $X=Vd \cdot C$, what does Vd denotes?
a) Density
b) Volume of blood
c) Volume of body
d) Volume of distribution
- (32) How can you determine the extracellular fluid volume?
a) Evans blue
b) Na^+
c) D20
d) Tritiated water
- (33) Which one of the below does not belong to the 4 classes of lipoprotein?
a) Chylomicrons
b) Very low-density lipoproteins
c) High-density lipoprotein
d) Fatty acids
- (34) Which drugs bind to RBC membrane?
a) Pentobarbital
b) Acetazolamide
c) Imipramine
d) Phenytoin
- (35) Which one of the following is the principal organ for drug excretion?
a) Lungs
b) Liver
c) Kidneys
d) Sweat glands
- (36) Which compounds are excreted through the lungs?
a) Lipophilic
b) Gaseous
c) Liquid and hydrophilic
d) Solid less than 100 Dalton
- (37) Which of the following compounds are used as agents to determine glomerular filtration rate?
a) Calcium ion
b) Albumin
c) Creatinine
d) Calcium carbonate
- (38) What is the equation for clearance?
a) Elimination rate / plasma drug concentration
b) Plasma drug concentration/elimination rate
c) $1 / \text{Plasma drug concentration}$
d) $1 / \text{Elimination rate}$
- (39) What will be the renal clearance ratio of a drug whose renal clearance is 40 ml/min and the clearance of creatinine is 95 ml/min?
a) 0.421
b) 2.38
c) 0.010
d) 0.025
- (40) Which drugs cannot be filtered through glomerulus?
a) Drugs bound to plasma proteins
b) Unbound
c) Free drug
d) Below molecular weight of 300 Dalton
- (41) What is the equation of bioavailable fraction?
a) $1/\text{Bioavailable dose}$
b) $1/\text{Administered dose}$
c) $\text{Bioavailable dose} / \text{Administered dose}$
d) $\text{Administered dose} / \text{Bioavailable dose}$
- (42) Which of the following is not an important parameter of plasma level time studies?
a) C_{max}
b) T_{max}
c) The area under the plasma level-time curve
d) Steady state level

- a) Physiologic model
c) Noncompartment model
- b) Compartment mode
d) Mammillary model
- (55) In pharmacokinetics, the term 'rate' refers to a change in which of the following measurements over time.
- a) Drug dose
c) Concentration of drug in plasma
- b) Drug elimination
d) Drug metabolism
- (56) Instantaneous distribution to most body tissues and fluids is assumed in which of the following models?
- a) One-compartment model
c) Multicompartment model
- b) Two-model
d) Non-compartmental model
- (57) The amount of drug per unit of volume is defined as the:
- a) Volume of distribution
c) Rate
- b) Concentration
d) Absorption
- (58) Which data is needed to decide on that the drug is suitable to prepare retard preparation?
- a) Clearance
c) Biological half life
- b) Area under the curve
d) Absorption rate constant
- (59) Which method is not suitable to calculate area under the curve?
- a) Least square method
c) Trapezoidal rule
- b) Weighing
d) Integration of curve
- (60) Which factors has no effect on bioavailability?
- a) Maximum plasma level
c) Tmax
- b) Therapeutic range
d) Quantity of food
- (61) Which marker is used to estimate volume of plasma?
- a) Evans blue
c) HTO
- b) Cr-51
d) Antipyrine
- (62) Unit for rate of infusion
- a) Mg/L
c) mg/h
- b) mg
d) mg.L/h
- (63) Clearance is determined as the ratio of
- a) Rate of Absorption to Plasma drug concentration
c) Rate of Elimination to Plasma drug concentration
- b) Rate of Elimination to Volume of distribution
d) Rate of Elimination to Plasma drug concentration
- (64) The loading dose of a drug is usually based on
- a) Total clearance of the drug
c) Fraction of drug excreted unchanged in urine
- b) Plasma protein binding percentage
d) Apparent volume of distribution and desired steady state drug concentration in plasma
- (65) Which is not a factor influencing the plasma elimination half life of a drug?
- a) Apparent volume of distribution
c) Protein binding
- b) Clearance
d) Route of administration
- (66) The objective of pharmacokinetic model is to quantify the drug content in-
- a) Distribution
c) Disintegration
- b) Dissolution
d) Diffusion

- (67) A system showing dose dependent pharmacokinetics, will follow-
- a) Linear pharmacokinetics
 - b) Non-linear pharmacokinetics
 - c) Zero order
 - d) Pseudo first order
- (68) Which of the following statement is correct with respect to non-linear pharmacokinetics?
- a) First order
 - b) First order followed by zero order
 - c) Pseudo first order
 - d) Zero order
- (69) For determining in vivo Michaelis- Menten constant, two doses of following are used-
- a) Inhalation
 - b) Infusion
 - c) I.V. bolus
 - d) Oral
- (70) The disadvantages of in vivo method of determining K_m and V_m are
- a) Clearance changes
 - b) Compartment model changes
 - c) Drug is eliminated by more than one capacity limited elimination
 - d) Unpredicted dose level
- (71) Double reciprocal plot of Michaelis- Menten equation is also called as-
- a) Hanes- Woolf plot
 - b) Lineweaver- Burke plot
 - c) Scatchard plot
 - d) Metabolism
- (72) Which of the following is not involved in non-linear pharmacokinetics?
- a) Binding to proteins and tissue
 - b) Release and dissolution
 - c) Enzymes or carrier systems
 - d) Diffusion and permeability
- (73) According to chrono-pharmacokinetics, which of the following factors is responsible for variation in drug distribution?
- a) Protein binding
 - b) Extracellular fluid
 - c) Red blood cells
 - d) Tissue binding
- (74) Which of the following is not a cause of non-linear pharmacokinetics?
- a) Saturation of plasma protein binding
 - b) Saturation of carrier molecules
 - c) Enzyme inhibition
 - d) Enzyme induction
- (75) Chrono-pharmacokinetics involves the study of ADME with reference to -
- a) Dosing interval
 - b) Time of administration
 - c) Time of the day
 - d) Sample collection time