BHARATI VIDYAPEETH (DEEMED TO BE UNIVERSITY), PUNE, INDIA PhD Entrance Test – 2022

SECTION-II: Pharmaceutics - 50 Marks

Section II	
1 Physical Pharmaceutics covering the following aspects:	
	Solids: Handling of solids, pharmaceutical granulation, compression and
	compaction properties of binary mixtures, lubricant sensitivity,
	characterization of granules and compacts.
	Dissolution: Theory of dissolution, concept of drug release. Dissolution test
	apparatus: different designs, factors affecting dissolution rate. Dissolution of
	different dosage forms: solids, suspensions, topicals, suppositories and
	controlled release systems.
	Pharmaceutical aspects of solubilization: Solubilization of drugs by
	following approaches: use of surfactants for solubilization; solid
	dispersions, cyclodextrin inclusion complexes, cosolvency etc.
2	Pharmaceutical preformulations:
	Drug Excipient interactions: different methods
	Drug Stabilityand Kinetics:General considerations & concepts, half-life
	determination, Influence of temperature, light, solvent, catalytic species and
	other factors, Accelerated stability study, expiration dating.
3	Novel Drug Delivery Systems covering the following aspects:
	Design, development, manufacture and evaluation of the following:
	Oral Drug Delivery Systems: Osmotic DDS, Ion exchange controlled
	DDS, Hydrodynamically balanced DDS Myeogol DDS: Physical spiced basis of myeogol delivery with reference to
	Mucosal DDS: Physiological basis of mucosal delivery with reference to oral mucosal, nasal, vaginal and rectal routes. Bioadhesion and bioadhesive
	polymers, DDS for mucosal administration. In-vitro, ex-vivo and in-vivo
	evaluationtechniques
	Transdermal DDS: Percutaneous absorption and penetration enhancers,
	development of transdermal gels, patches with reference to components and
	evaluation. Iontophoretic and Sonophoretic DDS.
	Microspheres: Methods to obtain microcapsules/ microspheres, their
	evaluation and applications
	Nanoparticulate and Colloidal systems: Polymeric and lipid
	nanoparticles, liposomes, niosomes, and polymeric micelles.
4	Biopharmaceutics and Pharmacokinetics covering the following
	aspects:
	Absorption, Distribution, Metabolism, Excretion
	Pharmacokinetics : Pharmacokinetics models, Laplace transformations and
	concept of compartment modeling. One compartment model: intravenous
	injection, intravenous infusion, extravascular route
	Bio-availability and Bioequivalence: Study design, protocols and
	regulatoryrequirements and statistical consideration in data analysis.

References:

- 1. A.Martin, P.Bustamante and A.H.Chun; Physical Pharmacy; Waverly.
- 2. N.G.Stanley-Wood, Enlargement and compaction of particle solids; Butterworths.
- 3. D.M.Parikh, Handbook of Pharmaceutical Granulation Technology; Marcel Dekker.
- 4. H.G.Brittain; Physical Characterization of Pharmaceutical solids; Marcel Dekker.
- 5. Lieberman, Rieser and Banker; Pharmaceutical dosage forms; Disperse system; Marcel Dekker.
- 6. M.N. Rubinstein, Pharmaceutical Technology, Drug Stability, John Wiley and Sons.
- 7. Martin R hodes, Principles of Powder Technology, John Wiley and Sons.
- 8. James J. Wells, Pharmaceutical Preformulation, Ellis Horwood Ltd.
- 9.P. H. List and P. C. Schmidt; Pharmaceutical technology, C R Spress.
- 10. Robinson, Novel Drug Delivery System, Marcel Dekker.
