

29- The biogeneric drug is handled equally like small molecule APIs in the regulatory authorities ( T / E )

According to that, FOPs is

- a) biosimilar    b) biogeneric    c) interchangeable    d) small molecule

FOPs stand for follow on proteins

30 – The 1<sup>st</sup> level of the ATC code is designed by digits ( T / E )

This is because digits are in levels 2 and 5

31- Priority review is granted by the RA (regulatory authority) when sponser submit ANDA for a generic drug (T / E)

This is because Priority review is for breakthrough drug (submit NDA) not generic drugs

32- Daclizumab is of ..... origin

- a) humanized    b) rat    c) human    d) murine

✓ 33- Biotechnology can produce small molecule APIs (T / F ). An example is:

- a) 11- $\beta$ -hydroxylation of steroids    b) leeches    c) r-GH    d) Hirudine

34- Deconvolution algorism is used by:

- a) solid state CC    b) CC synthesis in solution    c) Dynamic CC  
d) target directed fragment assembly    e) none of the above

This is because it can be used with all of the above

35- Fragment based SAR is one of the analogue approach in the DD (T / E).

This is because it is of the one type of the QSAR studies

36- Deconvolution method is followed during:

- a) CC synthesis    b) HTS    c) A followed by B    d) none

This is because it's used to identify the most active compound of the library formed in the CC using HTS tests.

# Drug Design Mcq With Answers

**RJ Alexander**



## **Drug Design Mcq With Answers:**

**TEXT BOOK OF MEDICINAL CHEMISTRY-I** Dr. Sohini Dixit, Dr. Mukesh Bansal, Krutika Warthi, Santhosh Kumar Banoth, Ajeet Singh, 2025-12-10 The Textbook of Medicinal Chemistry I is a comprehensive guide designed for pharmacy and medicinal chemistry students. It introduces learners to the evolution of medicinal chemistry, highlighting the history, growth, and significance of the discipline in modern drug discovery. The book begins with a thorough explanation of physicochemical properties like ionization, solubility, partition coefficient, hydrogen bonding, protein binding, chelation, bioisosterism, and stereochemistry, all of which dictate the biological action of drugs. It then progresses to drug metabolism, covering the principles of Phase I and Phase II reactions, as well as the factors including stereochemical aspects that influence metabolic pathways. A major portion of the text is dedicated to drugs acting on the autonomic nervous system. Students are introduced to adrenergic neurotransmitters with detailed discussions on the biosynthesis and catabolism of catecholamines, alongside receptor classification and distribution. Sympathomimetic agents are presented systematically, including their classification, mechanisms of action, uses, and structure-activity relationships (SAR). Both direct and indirect acting sympathomimetic agents are highlighted, along with those having mixed mechanisms. Adrenergic antagonists are given equal emphasis, covering both alpha and beta blockers with their SAR and clinical significance. The cholinergic system is another core area, with explanations of acetylcholine biosynthesis, catabolism, and receptor subtypes. Parasympathomimetic agents are classified and elaborated upon, including direct acting agents such as carbachol and pilocarpine, and indirect acting cholinesterase inhibitors, both reversible and irreversible. The text also details cholinesterase reactivators like pralidoxime. Cholinergic blocking agents are discussed in depth, ranging from natural solanaceous alkaloids such as atropine and scopolamine to synthetic derivatives like dicyclomine and ipratropium, with their SAR, mechanisms, and therapeutic uses clearly explained. Moving into central nervous system drugs, the book explores sedatives and hypnotics, particularly benzodiazepines and barbiturates. Their SAR, mechanisms, and clinical applications are thoroughly described, along with miscellaneous sedative agents such as meprobamate and paraldehyde. Antipsychotics are then presented, classified into phenothiazines, ring analogues, butyrophenones, and other groups, with special focus on their SAR and therapeutic uses. Anticonvulsants receive dedicated coverage, as well, including barbiturates, hydantoins, succinimides, benzodiazepines, and newer agents like carbamazepine, valproic acid, and gabapentin, alongside their mechanisms of action. The section on general anesthetics explains classification, mechanisms, and clinical applications, discussing inhalation anesthetics such as halothane and sevoflurane, ultra short acting barbiturates like methohexital, and dissociative agents like ketamine. Narcotic and non-narcotic analgesics form another crucial segment, where the SAR of morphine analogues is highlighted. Students gain insights into opioids, opioid antagonists, and non-opioid analgesics, including anti-inflammatory agents like aspirin, ibuprofen, mefenamic acid, and diclofenac.

**TEXT BOOK OF INDUSTRIAL PHARMACY-II** Dr. Ritu Gulia, Ms Poonam Sandhu, Ms. Tilottama

Bhattacharya, Dr. Viram Parmar, Ms. Aditi Sharma, 2025-05-20 The Textbook of Industrial Pharmacy II is a comprehensive guide tailored for students researchers and professionals engaged in the pharmaceutical industry focusing on critical areas of drug manufacturing and regulation It delves into pilot plant scale up techniques highlighting key factors such as personnel and space requirements raw materials and process adaptation from laboratory to industrial scale for solids liquids and semi solids The book emphasizes the importance of proper documentation and introduces SUPAC guidelines and platform technologies which are essential for ensuring consistent quality and compliance It also offers an in depth discussion on technology development and transfer TT referencing WHO guidelines and addressing granular processes for APIs excipients packaging materials and finished products The documentation equipment qualification validation and regulatory agency roles are thoroughly covered including insight into Indian TT bodies like APCTD and NRDC A dedicated section on regulatory affairs explores their evolution functions and the responsibilities of professionals in the field It examines the steps involved in drug approval starting from preclinical development through IND and NDA submissions and stresses the significance of clinical protocols biostatistics and data presentation in gaining FDA approval Furthermore the book discusses quality management systems detailing modern quality tools like TQM QbD Six Sigma and standard systems such as ISO 9000 ISO 14000 NABL and GLP essential for ensuring regulatory compliance and product excellence Lastly it elaborates on Indian regulatory requirements shedding light on the organizational structure and role of CDSCO and State Licensing Authorities with a focus on obtaining the Certificate of Pharmaceutical Product COPP and navigating the approval procedures for new drugs This book is a valuable academic and practical resource for understanding the multidisciplinary scope of industrial pharmacy and its regulatory landscape TEXT BOOK OF MOLECULAR PHARMACEUTICS (NANO TECHNOLOGY & TARGETED DDS)

Dr Ravi Kumar Kota, Mr K. Sampath Kumar, Mrs. Nallagandla Rajitha, Dr L. Jyothi Rani, Mrs M. Anusha, 2025-12-18 Text Book of Molecular Pharmaceutics Nanotechnology Targeted DDS introduces students to advanced concepts of site specific drug delivery It explains why targeted drug delivery systems are needed to improve therapeutic efficacy and reduce side effects The book describes the basic concepts and key events involved when a targeted carrier travels from the administration site to the target tissue Biological processes that govern drug targeting such as recognition circulation extravasation and cellular uptake are clearly highlighted Special emphasis is given to tumor targeting strategies including the principles of selective accumulation in cancer tissues Brain specific drug delivery is discussed with focus on overcoming the blood brain barrier and improving CNS drug transport A dedicated unit on targeting methods presents their introduction basic design principles and classification It explains different approaches for the preparation of targeting systems and the critical parameters that influence their performance Types of nanoparticles used in drug delivery are described along with their advantages over conventional dosage forms The text outlines common techniques for nanoparticle preparation and methods used to characterize them Different types of liposomes are presented as versatile carriers for drugs

proteins and genes Students are guided through the preparation of liposomes and the evaluation tests required to ensure their quality The book defines microcapsules and microspheres and explains their structural and functional features It describes the types preparation techniques and evaluation parameters of microcapsules and microspheres in detail An introduction to monoclonal antibodies is provided covering their production and role in targeted therapy Novel carrier systems such as niosomes aquasomes phytosomes and electrosomes are explained with their methods of preparation The pharmaceutical and clinical applications of these novel carriers are highlighted to show their practical importance Pulmonary drug delivery systems are discussed with special reference to aerosols as efficient lung targeted dosage forms The role of propellants containers and different types of aerosols in pulmonary delivery is clearly explained Formulation and evaluation of aerosol preparations are described to help students understand product development aspects The book also introduces intranasal route delivery systems as a simple and noninvasive approach for local and systemic therapy Types preparation methods and evaluation of intranasal formulations are systematically discussed A final unit on nucleic acid based therapeutic delivery systems provides a foundation in gene therapy It covers ex vivo and in vivo gene therapy viral and nonviral gene transfer systems and liposomal gene delivery The text further discusses biodistribution and pharmacokinetics of these advanced systems It concludes with therapeutic antisense molecules and aptamers presenting them as promising drugs of the future

**All India Ayush Post Graduate Entrance Test [AIAPGET] Ayurveda Question Bank [MCQ] Book 1500+ Questions With Detail Explanations** DIWAKAR EDUCATION HUBY,2024-09-09 All India Ayush Post Graduate Entrance Test AIAPGET Ayurveda Question Bank MCQ Book 1500 Questions With Detail Explanations Highlight of MCQ Book Cover All Chapters of Ayurveda As Per Syllabus Given Detail Explanations of Each MCQ Design by Expert Faculties As Per Updated Syllabus [CUET PG Life Science Book 2026 Solved Previous Year Paper 2022 to 2025 and Chapter Wise 1500 MCQ As Per Updated Syllabus](#) ,2026-01-07 **Generative AI for Business Leaders (One-Sitting Guide)**

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**Biomedical Informatics** Andreas Holzinger, 2014-05-06 This book provides a broad overview of the topic Bioinformatics with focus on data information and knowledge From data acquisition and storage to visualization ranging through privacy regulatory and other practical and theoretical topics the author touches several fundamental aspects of the innovative interface between Medical and Technology domains that is Biomedical Informatics Each chapter starts by providing a useful inventory of definitions and commonly used acronyms for each topic and throughout the text the reader finds several real world examples methodologies and ideas that complement the technical and theoretical background This new edition includes new sections at the end of each chapter called future outlook and research avenues providing pointers to future challenges At the beginning of each chapter a new section called key problems has been added where the author discusses possible traps and unsolvable or major problems *NEET PG - Community Medicine (Preventive and Social Medicine) (Question Bank 3300 MCQs Chapterwise)* Mocktime Publication, NEET PG Community Medicine Preventive and Social

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