

Total No. of Questions :6]

[Total No. of Pages :2

P1831

[3957] - 101

M.Pharmacy (Sem. - I)

**ADVANCED ANALYTICAL TECHNIQUES
(2008 Pattern)**

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question 1 and 4 are compulsory.*
- 2) *Attempt any one question from the remaining in section I and any one question from the remaining questions of section II.*
- 3) *Answers to the two sections should be written in separate books.*
- 4) *Draw diagrams wherever necessary.*
- 5) *Figures to the right indicate full marks.*

SECTION - I

- Q1) a)** Suggest suitable chemical structural formula for following spectroscopic data. **[8]**
MF $C_7H_{12}O_4$
IR : 1745 cm^{-1}
Proton NMR :
3.4 singlet 2H
1.3 triplet 6H
4.2 quartate 4H
- b) Explain with example how inductive effect, mesomeric effect, resonance and ring strain affect absorption of IR radiation. **[8]**
- c) What is Bragg's law? Write its significance. **[4]**
- Q2) a)** Discuss principle, instrumentation and applications of thermogravimetric analysis. **[8]**
- b) Write about fundamental law of absorption and its applications in quantitative analysis. **[6]**
- c) Write principle and applications of ESR spectroscopy **[6]**
- Q3) a)** Discuss modes of vibrations in linear and non linear molecule. **[6]**
- b) Comment on hyphenated techniques **[8]**
- c) Explain effect of polarity of solvent on absorption maxima in UV spectroscopy. **[6]**

P.T.O.

SECTION - II

- Q4)** a) Explain in detail the working of various pumps used in HPLC. [10]
b) Give different modes of fragmentation for amino acids and alcohols in EIMS. [10]

Q5) Following equation is a fundamental equation in HPLC technique. Explain clearly and in detail the meaning of various terms involved in the equation and how resolution [Rs] can be controlled through modification of these terms given in the equation. [20]

$$R_s = \frac{1}{4} (\alpha - 1) \sqrt{n} (K' / 1 + K')$$

- Q6)** a) How will you differentiate / identify between the following pairs using mass spectrometry [MS] and / or Proton NMR spectrometry. [10]
i) para-n-Propyltoluene and 1'-Methylpropylbenzene.
ii) para-i-Propylethylbenzene and 1-methyl-3, 5-diethylbenzene.
b) Explain why and how derivatization is carried out in HPLC and GC. [10]



Total No. of Questions :8]

[Total No. of Pages :2

P1839

[3957] - 108

M.Pharmacy (Sem. - I & II)

**QUALITY CONTROL & ASSURANCE OF PHARMACEUTICALS
(2008 Pattern)**

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory.*
- 2) *Solve any two from the remaining questions for each section.*
- 3) *Answers to the two sections should be written in separate books.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

Q1) Discuss management of rejected & recovered materials in pharmaceutical processing. **[10]**

Q2) a) Explain the concept of quality culture & importance of staff training in maintaining it. **[8]**

b) Discuss hazards of mix-ups and cross contaminations. **[7]**

Q3) a) Define key personnel and explain the responsibilities of key personnel in pharmaceutical industry. **[8]**

b) Quality control of packaging materials. **[7]**

Q4) Write short note on: **[15]**

- a) Good manufacturing practises.
- b) Sanitation of manufacturing premises.
- c) SOP on Personnel hygiene.

P.T.O.

SECTION - II

- Q5)** What is the significance of pharmaceutical manufacturing documentation? Explain in detail batch production & control record. **[10]**
- Q6)** a) Enlist component of HVAC system & detail the construction of HEPA filter unit. **[8]**
b) Explain the significance & procedure of cleaning validation. **[7]**
- Q7)** a) Explain the significance & procedure for pharmaceutical plant audit. **[8]**
b) Explain quality control of biological products. **[7]**
- Q8)** Write short note on: **[15]**
- a) Validation master plan.
b) Drug master file.
c) Sanitation in aseptic area.



Total No. of Questions :8]

[Total No. of Pages :2

P1840

[3957] - 112

M.Pharmacy (Sem. - I & II)

CHEMISTRY OF MEDICINAL NATURAL PRODUCTS

(2008 Pattern)

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Attempt any two questions from remaining for section I and section II each.*
- 2) *Figures to the right indicates full marks.*
- 3) *Answers to the two sections should be written in separate answer books.*

SECTION - I

Q1) Describe the chemistry and structural elucidation of Caffeine. [10]

Q2) a) Discuss the methods of isolation of essential oil and separation of terpenoids from essential oil. [8]

b) Explain the biosynthesis of fatty acids. [7]

Q3) a) Explain the biosynthetic pathway of isoprenoid compounds. [8]

b) Describe the structural elucidation of morphine. [7]

Q4) Write note on following (any two): [15]

- a) Methods of extraction of alkaloids.
- b) Isolation and purification of glycosides.
- c) Shikimic acid Pathway.

P.T.O.

SECTION II

- Q5)** Describe the chemistry and structural elucidation of solasodine. [10]
- Q6)** a) Discuss in detail the chemistry of plant steroids. [8]
b) Classify flavonoids. Discuss the properties of flavonoids. [7]
- Q7)** a) Explain in detail the chemistry of disaccharides. [8]
b) Describe the structural elucidation of ephedrine. [7]
- Q8)** Write note on following (any two): [15]
a) General reactions of monosaccharides.
b) Chemistry of diosgenin.
c) Chemistry of carotenoids.



Total No. of Questions :8]

[Total No. of Pages :2

P1841

[3957] - 117

M.Pharmacy (Sem. - I & II)

NATURAL PRODUCTS MANAGEMENT

(2008 Pattern)

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Out of the remaining solve any two questions from section I and any two questions from section II.*
- 2) *Answers to the two sections should be written in separate answer books.*

SECTION - I

Q1) Write about the appraisal of farm resources, capital resources, management factors, land resources and enterpreural aspects of farm analysis & farm planning. **[10]**

- Q2)** a) Discuss mechanization/modernization of Natural Products Market. **[8]**
b) Comment on - processing of an agricultural marketing. **[7]**

- Q3)** Write about
a) Farm planning & budgeting. **[8]**
b) Application of research in farm management. **[7]**

Q4) Write on - 'Co-operative processing / efforts among collectors & growers to store, transport & market the natural products'. **[15]**

P.T.O.

SECTION - II

Q5) Write about ex-situ and in-situ cultivation & conservation of medicinal plants. **[10]**

Q6) a) Write about cultivation economics and project proposals for few prioritized medicinal plants of India. **[8]**

b) Give an account on the legal requirements & processing techniques for marketing of raw materials and value added products in relation to herbal cosmetics. **[7]**

Q7) Discuss the general requirements to establish extraction unit based on herbs/herbal products. **[15]**

Q8) Write an essay on 'IPR in relation to medicinal herbs and herbal products'. **[15]**



Total No. of Questions :8]

[Total No. of Pages :2

P1842

[3957] - 118

M.Pharmacy (Sem. - I & II)

MEDICINAL PLANT BIOTECHNOLOGY

(2008 Pattern)

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Out of remaining attempt any two questions from section I and section II.*
- 2) *Figures to the right indicate full marks.*

SECTION - I

Q1) Give detail account of pharmaceutical applications of enzyme immobilization. **[10]**

Q2) a) Write about purification of enzymes. **[8]**

b) Describe methods of protoplast fusion. **[7]**

Q3) a) Give an account of hairy root culture and multiple shoot culture. **[8]**

b) Enlist different physical methods of DNA mediated gene transfer. **[7]**

Q4) Write notes on : **[15]**

a) Somaclonal variation and synthetic seeds.

b) Ti plasmid.

c) Micropropagation.

P.T.O.

SECTION - II

Q5) Give an account of enzyme reactor. **[10]**

Q6) a) Describe method of gene transfer using vectors of *Agrobacterium*. **[8]**

b) Write about RAPD markers for genetic mapping. **[7]**

Q7) a) Discuss RFLP genetic maps in plants. **[8]**

b) Give an account of physical maps using In situ hybridization. **[7]**

Q8) Write notes on: **[15]**

a) Mutation.

b) Hybridization.

c) Applications of PCR.



Total No. of Questions :8]

[Total No. of Pages :2

P1843

[3957] - 207

M.Pharmacy (Sem. - II)

(Spl. Pharmacology)

MOLECULAR PHARMACOLOGY

(2008 Pattern)

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory.*
- 2) *Solve any two questions from the remaining in section I and section II.*
- 3) *Figures to the right indicate full marks.*
- 4) *Write answers for section I and II in separate answer sheets.*

SECTION - I

Q1) Enlist various endogenous bioactive molecules & discuss role of COX-2 regulators in inflammation. **[10]**

Q2) Classify Dopaminergic receptors. Add a note on various drugs acting on them. **[15]**

Q3) Define apoptosis. Describe it's pharmacological and clinical implications. **[15]**

Q4) Write a note on (any three): **[15]**

- a) Sodium channel modulators.
- b) Cellular signaling mechanism of drug action.
- c) Neuropeptides.
- d) Angiotensin receptors.

P.T.O.

SECTION - II

- Q5)** What is chronopharmacology? Discuss implications of chronopharmacology to drug therapy. **[10]**
- Q6)** Enlist various laboratory animals. Write a note on application of transgenic mouse in experimental pharmacology. **[15]**
- Q7)** Define Immunopharmacology with respect to cellular cytotoxicity. **[15]**
- Q8)** Describe potential of human genome mapping in drug research. **[15]**

Total No. of Questions :8]

[Total No. of Pages :2

P1844

[3957] - 208

M.Pharmacy (Sem. - II)

(Spl. Pharmacognosy)

PHYTOCHEMISTRY & PHYTOPHARMACEUTICALS

(2008 Pattern)

Time : 3 Hours]

[Max. Marks :80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory.*
- 2) *Out of the remaining attempt any two questions from section I and any two questions from the section II.*
- 3) *Answers to the two sections should be written in separate books.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

Q1) Describe in detail methods of extraction, isolation, characterization and structure elucidation of Caffeine. **[10]**

Q2) Give the Instrumental identification of following phytoconstituents. **[15]**

- a) Gingerol
- b) Curcumin
- c) Vasicine

Q3) Explain in detail the chemistry of Saponins and give the Pharmaceutical profile of Glycyrrhizinic acid. **[15]**

Q4) Write short notes (any two): **[15]**

- a) IR spectral analysis of Rutin and Atropine.
- b) Extraction and isolation of Sennosides.
- c) Pharmaceutical significance of Taxol.

P.T.O.

SECTION - II

Q5) Explain the principle, procedure, and importance of following parameters in evaluation of Natural products as per WHO guidelines. **[10]**

- a) Determination of Arsenic and Heavy metals.
- b) Pesticide residue.

Q6) Describe in detail various pharmacological screening methods for evaluation of **[15]**

- a) Hepatoprotective activity.
- b) Anti epileptic activity.

Q7) Explain in detail the various methods and related equipment for extraction of herbal drugs. **[15]**

Q8) Write short notes (any two): **[15]**

- a) Pharmacological screening of Anti oxidants.
- b) Evaluation of Herbal extracts.
- c) Physical evaluation of crude drugs.



Total No. of Questions : 6]

[Total No. of Pages : 1

P1832

[3957]-102

M.Pharmacy

RESEARCH METHODOLOGY

(2008 Pattern) (Sem. - I)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Attempt any two questions from Section - I and any two questions from Section - II.*
- 2) *Answers to the two sections should be written in separate answer books.*
- 3) *All questions carry equal marks.*

SECTION - I

Q1) What is the purpose of research? Enlist the different types or research. Give the elaborated account of historical, descriptive and patent oriented research. **[20]**

- Q2)** a) Give the elaborated account on questionnaire contents and working. **[10]**
- b) Enlist the different tools used in research for data collection. Add a note on Interview method along with merits and demerits. **[10]**

- Q3)** Write notes on any two of the following: **[20]**
- a) Use of computer packages in documentation.
 - b) Student 't' test.
 - c) Importance of methodology and results and discussion in thesis writing.

SECTION - II

Q4) What is the meaning of hypothesis. Describe the various sources of hypothesis. Add a note on role of hypothesis in research. **[20]**

Q5) Give the salient features of techniques involved in oral presentation of research outcome. **[20]**

- Q6)** Write notes on any two of the following: **[20]**
- a) Use of bibliography in research.
 - b) Correlation data.
 - c) Use of visual aids in oral presentation.



Total No. of Questions : 6]

[Total No. of Pages : 1

P1834

[3957]-103

M.Pharmacy

(Spl. Pharmaceutics)

ADVANCED PHARMACEUTICS - I

(2008 Pattern) (Sem. - I)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Answer two questions from Section - I and two questions from Section - II.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

Q1) Elaborate the preformulation studies of semisolids. **[20]**

Q2) Describe the biodegradable polymers and their significance. **[20]**

Q3) Write short notes (Any Two): **[20]**

- a) Superdisintegrants.
- b) Liquid crystals.
- c) Overages and ICH guidelines.

SECTION - II

Q4) Explain the drug release modeling through polymer matrix and laminates. **[20]**

Q5) Discuss the significance of following optimization techniques: Simplex method, EVOP, Grid search method. **[20]**

Q6) Write short notes (Any Two): **[20]**

- a) Validation of pharmaceutical processes.
- b) Dissolution models.
- c) Methods of microencapsulation.



Total No. of Questions : 8]

[Total No. of Pages : 2

P1835

[3957]-104

M.Pharmacy

(Spl. Pharmaceutical Chemistry)

ADVANCED PHARMACEUTICAL CHEMISTRY

(2008 Pattern) (Sem. - I)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question No. 1 and 5 is compulsory. Out of the remaining attempt any two questions from each Section - I and Section - II.*
- 2) *Write answer to Section - I and Section - II in separate answer book.*

SECTION - I

Q1) What are conformational isomers? Explain with examples how the pharmacological properties of drugs changes with conformational Isomerism. **[10]**

Q2) Discuss the mechanism, stereochemistry and applications of Grignard Reaction taking example of medicinal agent. **[15]**

Q3) Write notes on any two: **[15]**

- a) Pinacol rearrangement.
- b) Diazomethane and its synthetic applications.
- c) Free radical reaction.

Q4) Explain synthone approach of designing drug synthesis. Develop the synthetic route for Terfenadine or Rosiglitazone using synthone approach. **[15]**

SECTION - II

Q5) What are chiral drugs? Explain how the chirality of medicinal agents affects the pharmacodynamic and pharmacokinetic properties. **[10]**

P.T.O.

Q6) Discuss the mechanism, stereochemistry and applications of Wittig Reaction taking example of medicinal agent. **[15]**

Q7) What are reduction reactions? Explain Birch reduction. **[15]**

Q8) Write note on any Two: **[15]**

- a) Ionic liquids and supercritical liquids.
- b) Solvent free reactions by microwave and ultrasound energy.
- c) Oppenauer oxidation.



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Total No. of Questions : 8]

[Total No. of Pages : 2

P1836

[3957]-105

M.Pharmacy

(Spl. Pharmacology)

ADVANCED PHARMACOLOGY

(Pre-clinical Evaluation of Drugs)

(2008 Pattern) (Sem. - I)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Answer any two questions from the remaining.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*

SECTION - I

Q1) Explain in detail 3 Rs in Animal Experimentation. Discuss how these 3 Rs can be achieved while preparation of research protocol as per form - B. [10]

Q2) Discuss in detail organization of preclinical screening programme. Explain in detail all the safety assessment tests. [15]

Q3) Explain the latest structure and function of Institutional Animal Ethical Committee as per CPCSEA. Write in detail structure, layout of an animal house. Add a note on handling and breeding techniques of laboratory animals. [15]

Q4) Write a Note on Any Two: [15]

- a) Patch Clamp Technique.
- b) Limitations of *In vitro* testing of drugs.
- c) Alternatives to Animal Studies.

P.T.O.

SECTION - II

- Q5)** Discuss the design and procedures for screening of diuretic agents. [10]
- Q6)** Discuss the animal models for evaluation of antiparkinsonian agents. [15]
- Q7)** Write in detail principle of design and animal models for screening of drugs used in treatment of cardiac arrhythmia. [15]
- Q8)** Write a Note on Any Two: [15]
- a) Screening of Histamin antagonists.
 - b) Evaluation of androgens.
 - c) Methods to test drugs acting as laxatives.



Total No. of Questions : 8]

[Total No. of Pages : 2

P1837

[3957]-106

M.Pharmacy

(Spl. Pharmacognosy)

ADVANCED PHARMACOGNOSY

(2008 Pattern) (Sem. - I)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Out of the remaining attempt 2 questions from Section - I and 2 questions from Section - II.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

Q1) Describe biotic and abiotic elicitors-induced production of secondary metabolites using plant cell culture. **[10]**

Q2) a) What is chemotaxonomy? What are its advantages & limitations over other methods of classifications? Explain the term divergence & convergence. **[7]**

b) Describe the terpenes as chemotaxonomic marker with suitable examples. **[8]**

Q3) Enlist various strategies used to enhance secondary metabolite production through tissue culture techniques. Describe genetic manipulation using plant cell culture. **[15]**

Q4) Write a Note on Any Two: **[15]**

- a) Coloring pigments derived from plants.
- b) Photosensitizing agent.
- c) Applications of biopolymers as pharmaceutical excipients.
- d) Biofuel.

P.T.O.

SECTION - II

- Q5)** Write various in vitro & in vivo models used in the evaluation of anticancer activity with suitable examples. **[10]**
- Q6)** Enlist techniques used in the study of plant biosynthesis. Describe sequential analysis technique along with various methods used for detection and measurement of radio labeled precursors. **[15]**
- Q7)** a) Explain the antidiabetic role of flavonoids. **[7]**
b) Review the plants having hepatoprotective activity. **[8]**
- Q8)** Write a Note on Any Three: **[15]**
- a) Flavonoids as anti-inflammatory agents.
 - b) Role of high throughput screening (HTS) in drug discovery.
 - c) Bioreactor for the production of secondary metabolites.
 - d) Camptothecin.



Total No. of Questions : 6]

[Total No. of Pages : 1

P1838

[3957]-107

M.Pharmacy

(Spl. Quality Assurance Techniques)

ADVANCED QUALITY ASSURANCE TECHNIQUES

(2008 Pattern) (Sem. - I) (Theory)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Solve any two questions from Section - I and two questions from Section - II.*
- 2) *All questions carry equal marks.*

SECTION - I

Q1) What is materials management in pharmaceutical industry? Write in details about every aspect of materials management. **[20]**

Q2) Explain how validation is important in pharmaceutical manufacturing and discuss various steps involved in validation of equipments with reference to tablet dosage form. **[20]**

Q3) a) Quality control of sterile product.
b) Steps in Environmental Protection. **[20]**

SECTION - II

Q4) Explain different components of quality assurance and discuss their importance in pharmaceutical manufacturing. **[20]**

Q5) Enlist various important facilities and discuss their relevance in building construction to provide suitable atmosphere for Pharmaceutical Industry. **[20]**

Q6) a) IPQC tests for Tablets and Capsules.
b) Plant level document. **[20]**



Total No. of Questions : 8]

[Total No. of Pages :2

P1850

[3957]-110

M.Pharmacy

BIOPHARMACEUTICS AND PHARMACOKINETICS

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 and 5 are compulsory. Out of the remaining attempt 2 questions from section I and 2 questions from section II.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*
- 4) *Figures to the right indicate full marks.*
- 5) *Use of logarithmic tables slide rule, Mollier charts, electronic pocket calculator and steam tables is allowed.*

SECTION - I

- Q1)** Describe Wagner Nelson method for determination of absorption rate constant. What is the limitation of this method? **[10]**
- Q2)** What is the significance of P gP(permeability glycoprotein) as an efflux transporter system in Development of targeted drug delivery system for brain? Explain with suitable example the drug design accordingly. **[15]**
- Q3)** Explain in detail the method of determination of hybrid first order constants for any drug that follows two compartment model and administered as i. v. bolus dose. **[15]**
- Q4)** Write short notes on any three. **[15]**
- a) Design and evaluation of bioequivalence studies.
 - b) In vivo/biological models for permeability studies.
 - c) Blood placental barrier.
 - d) Model independent approach in bioequivalence studies.

SECTION - II

- Q5)** How non linear kinetics of a drug is detected? Explain the causes of non linearity and significance. Give 'New Drug Application' requirements for a drug that follows Non linearity. **[10]**
- Q6)** What are drug displacement interactions due to protein binding? Why all displacement interactions are not clinically significant? Explain with examples. **[15]**

P.T.O.

Q7) The elimination half life of the Tobramycin was reported to be 2.15 hours and the volume of distribution was reported to be 33.5% of body weight. What is the dose for an 80 Kg individual if a steady state level of 2.5 $\mu\text{g/mL}$ is desired?

Assume that the drug is given by iv bolus injection every 8 hours. **[15]**

Q8) Write short notes on any three. **[15]**

- a) Determination and significance of 'Area under the curve'.
- b) Kinetics of protein binding.
- c) Significance of V_{max} and K_m .
- d) Apparent volume of distribution.



Total No. of Questions : 8]

[Total No. of Pages :2

P1851

[3957]-111

M.Pharmacy

STERILE PRODUCTS FORMULATION & TECHNOLOGY

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Out of the remaining attempt 2 questions from section I and 2 questions from section II.*
- 2) *Answers to the two sections should be written in separate answer books.*
- 3) *Neat diagrams must be drawn wherever necessary.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

- Q1)** Discuss physicochemical properties of drugs affecting design of parenteral dosage form. **[10]**
- Q2)** Explain the physiological considerations of LVP's. Discuss formulation and manufacturing process of LVP's. **[15]**
- Q3)** a) Discuss physiological factors affecting formulation of ophthalmic products. **[8]**
b) Explain Nanoparticles in parenteral drug delivery. **[7]**
- Q4)** Write a short notes on (any two): **[15]**
a) Liposomes in parenteral drug delivery.
b) Glass as a Parenteral packaging material.
c) Pyrogen Testing of sterile dosage forms.

SECTION - II

- Q5)** Describe construction, working and validation of HEPA filter. **[10]**
- Q6)** Describe in detail process selection and process specification in sterilization of parenterals. **[15]**
- Q7)** Explain overview of GMP and regulatory guidelines in parenteral manufacturing. **[15]**

P.T.O.

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Total No. of Questions : 8]

[Total No. of Pages :1

P1852

[3957]-116

M.Pharmacy

TRADITIONAL SYSTEMS OF MEDICINE AND AYURVEDIC FORMULATIONS

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 & 5 are compulsory. Answer any two questions from the remaining.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Figures to the right indicate full marks.*

SECTION - I

- Q1)** Write note on Ayurvedic cosmetic formulations. **[10]**
- Q2)** What is homeopathic system of medicine? Write about history, principles, homeopathic dilutions and herbs used in homeopathy. **[15]**
- Q3)** What is 'Guggulu'? Explain the process of sodhana. What are the characteristic of 'Sodhita guggulu' and how it is preserved? **[15]**
- Q4)** What is ethnopharmacognosy? How knowledge is affected by habitat change, species loss and the cultivation and hybridization of the plant? **[15]**

SECTION - II

- Q5)** What is Unani system of medicine? Write about its history and Unani medicines in Asia. **[10]**
- Q6)** Write brief note on standardization of Ayurvedic dosage forms using physical & chemical methods. **[15]**
- Q7)** Write note on acupuncture and moxibustion as Chinese system of medicine and its safety. **[15]**
- Q8)** Write down the differences between Ayurvedic medicine and homeopathic medicine with respect to history, philosophy and preparation of medicine. **[15]**



Total No. of Questions : 8]

[Total No. of Pages :1

P1853

[3957]-201

M.Pharmacy

DRUG REGULATORY AFFAIRS

(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 & 5 are compulsory. Out of the remaining attempt 2 questions from section I and 2 questions from section II.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

- Q1)** Explain in detail organisation structure and working of WHO and US - FDA. **[10]**
- Q2)** a) Explain the functions of Drug Inspector. **[8]**
b) Comment on salient features of pollution control Act. **[7]**
- Q3)** Comment on the provisions related to cosmetics under Drug & Cosmetics Act. **[15]**
- Q4)** Write in detail on Narcotic & Psychotropic Substances Act 1985. **[15]**

SECTION - II

- Q5)** Give the salient features of NDA. **[10]**
- Q6)** Explain the concept of 'Novelty' as applicable to patents. Give suitable examples. **[15]**
- Q7)** Compare and contrast the GMP of U.S.FDA and Indian regulatory body. **[15]**
- Q8)** Write short notes on (any three) **[3 × 5 =15]**
- a) DMF
 - b) Copyright
 - c) Industrial safety
 - d) Latest edition of I.P.



Total No. of Questions : 8]

[Total No. of Pages :2

P1854

[3957]-202

M.Pharmacy

(Spl. Pharmaceutics)

FORMULATIONS AND DEVELOPMENT

(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 & 5 are compulsory.*
- 2) *Solve any two questions from the remaining in Section - I and Section - II.*
- 3) *Figures to the right indicate full marks.*
- 4) *Answers to the two sections should be written in separate answer books.*

SECTION - I

- Q1)** How do multiple emulsions differ from micro emulsions? Explain with the help of ternary phase diagrams the selection of various phases in the formulation of microemulsions and SMEDDS. **[10]**
- Q2)** Discuss in detail various types of gastroretentive drug delivery systems. **[15]**
- Q3)** How can pulsatile delivery system be formulated? **[15]**
- Q4)** Write short notes on any three (5 marks each): **[15]**
- a) Emulgels based on liposomes.
 - b) Mouth dissolving tablets.
 - c) Sublingual formulations.
 - d) Penetration enhancers in semisolid preparations.

SECTION - II

- Q5)** What are the different types of containers used for aerosol Preparations? Discuss in detail the two phase and three phase system mode of operation of aerosols. **[10]**
- Q6)** Explain in detail the packaging material development for regulated markets for conventional and novel drug delivery systems. **[15]**
- Q7)** Discuss in detail veterinary specialized dose dispensers. Add a note on the need and problems of designing veterinary dosage forms. **[15]**

P.T.O.

Total No. of Questions : 8]

[Total No. of Pages :1

P1855

[3957]-203

M.Pharmacy

(Spl. Pharmaceutics)

NOVEL DRUG DELIVERY SYSTEMS

(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 & 5 are compulsory.*
- 2) *Solve any two questions from the remaining in Section - I and Section - II.*
- 3) *Figures to the right indicate full marks.*
- 4) *Answers to the two sections should be written in separate answer books.*

SECTION - I

- Q1)** Explain mechanisms of floating drug delivery? Describe its evaluation. [10]
- Q2)** What are the basic components of transdermal drug delivery system? Describe development of TDDS based on adhesive - dispersion type system. [15]
- Q3)** What is chronotherapeutics? Describe formulation and evaluation of pulsatile delivery system. [15]
- Q4)** Write notes (any two) [15]
- a) Biodegradable microspheres.
 - b) Long acting contraceptive formulations.
 - c) Mechanism of transmucosal transport of drugs.

SECTION - II

- Q5)** Describe evaluation procedures for colon targeted drug delivery. [10]
- Q6)** Describe different approaches to targeting drug delivery to brain. [15]
- Q7)** Describe the protein and peptide drug delivery. Give its limitations. [15]
- Q8)** Write notes (any three) [15]
- a) Drug targeting using monoclonal antibodies.
 - b) Stabilization of protein and peptide drugs.
 - c) Microbial approach for colon specific drug delivery.
 - d) Lacriserts.



Total No. of Questions : 6]

[Total No. of Pages : 2

P1856

[3957]-204

M.Pharmacy

ADVANCED MEDICINAL CHEMISTRY

(Spl. Pharmaceutical Chemistry)

(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) Write any two questions from Section - I and two questions from Section - II.
- 2) Write answer to Section - I and Section - II in separate answer book.

SECTION - I

Q1) a) Write the microbial conversions of prostaglandins giving suitable examples. **[15]**

b) Write a note on CADD. **[5]**

Q2) a) Discuss in detail different types of receptors. Highlight the features of models of cholinergic receptors. **[15]**

b) Explain the enzyme immobilization techniques. **[5]**

Q3) Write the synthetic steps with reaction conditions and mechanism involved in following synthesis (Any Two): **[20]**

a) Dapsone.

b) Ziprasidone.

c) Cetrizine.

SECTION - II

Q4) a) Discuss the various theories proposed for drug-receptor interactions. **[15]**

b) Give brief note on Gene Therapy. **[5]**

P.T.O.

- Q5)** a) Explain the various aspects of combinatorial chemistry. [10]
b) Explain the role of QSAR in drug design. [10]

- Q6)** Write notes on any two: [20]
a) GABA receptors.
b) Enzyme inhibition.
c) HTS.



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Total No. of Questions : 6]

[Total No. of Pages : 2

P1857

[3957]-205

M.Pharmacy
(Spl. Pharmaceutical Chemistry)
DRUG DESIGN
(2008 Pattern) (Sem. - II) (Theory)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 and 4 are compulsory.*
- 2) *Answer any one question from Section - I and any one question from Section - II from the remaining.*
- 3) *Answers to the two sections should be written on separate books.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

Q1) Explain in detail QSAR with its advantage and its application. Discuss Hansch's LFER model in detail. **[20]**

- Q2)** a) Explain the significance of ADME in drug design. **[10]**
b) Explain in detail analog approach for drug design with suitable example. **[10]**

Q3) Explain various approaches of drug design in detail with suitable examples. **[20]**

SECTION - II

Q4) The concept of antagonism and enzyme inhibition were proved to be excellent tools in the process of drug design - Explain, with suitable examples. **[20]**

- Q5)** a) Explain the conformational search techniques in CADD. **[10]**
b) Explain the concept of prodrugs in drug design. **[10]**

P.T.O.

Q6) Write short notes on *any four*:

[20]

- a) Rigid docking.
- b) Excluded volume & shape analysis.
- c) Artificial neural network in drug design.
- d) Quantum mechanics.
- e) Force fields.



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Total No. of Questions : 8]

[Total No. of Pages : 2

P1858

[3957]-209

M.Pharmacy
(Spl. Pharmacognosy)
INDUSTRIAL PHARMACOGNOSY
(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 and 5 are compulsory. Out of the remaining attempt 2 questions from Section - I and 2 questions from Section - II.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*

SECTION - I

Q1) Describe international/world wide trade of medicinal plants & derived products. **[10]**

Q2) Elaborate production and utilization of medicinal plants in India, with suitable examples. **[15]**

Q3) Explain role of medicinal and aromatic plants in future economic growth & development of herbal medicine industry. **[15]**

Q4) Write short note on: **[15]**

- a) Herbal Drug Regulation.
- b) Indian spices & their export potential.
- c) Phytopharmaceutical production in India.

P.T.O.

SECTION - II

Q5) Classify plant based industry. Elaborate scope of herbal drugs referring to various plant based industries. **[10]**

Q6) Elaborate the requirements of an ideal herbal extraction unit. Comment with reference to infrastructure & staff requirements. **[15]**

Q7) Elaborate utilization of Aromatic plant & derived products with reference to Indian trade. **[15]**

Q8) Write short note on: **[15]**

- a) Global regulatory status of herbal medicine.
- b) Production technique of Cinchona alkaloid.
- c) IPR & Herbal patents.



Total No. of Questions : 6]

[Total No. of Pages : 2

P1859

[3957]-210

M.Pharmacy

(Spl. Quality Assurance Tech.)

PHARMACEUTICAL VALIDATION

(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question Nos. 1 and 4 are compulsory. Out of the remaining attempt one question from Section - I and one question from Section - II.*
- 2) *Answers to the two sections should be written in separate books.*
- 3) *Neat diagrams must be drawn wherever necessary.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

Q1) a) Define validation; elaborate its benefits, Types and component. **[10]**

b) Explain in detail Validation Master Plan. **[10]**

Q2) What is importance of Equipment validation? Elaborate URS, DQ, IQ OQ and PQ of Fluid bed dryer and Autoclave. **[20]**

Q3) a) Define analytical method validation. Discuss validation parameters with respect to HPLC method. **[10]**

b) Give qualification of UV/Visible spectrophotometer. **[10]**

SECTION - II

Q4) a) Write importance of process validation. Elaborate validation of ampoules and vials. **[10]**

b) Write short note on vendor certification. **[10]**

P.T.O.

Q5) a) What is significance of cleaning validation? Discuss cleaning validation of Double cone mixer. **[10]**

b) Explain Validation of HAVAC system. **[10]**

Q6) Write short note: **[20]**

a) Computer System Validation.

b) Validation of integrated line by media fill test.



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Total No. of Questions : 8]

[Total No. of Pages : 2

P1860

[3957]-211

M.Pharmacy

(Spl. Quality Assurance Tech.)

QUALITY PLANNING AND ANALYSIS

(Theory) (2008 Pattern) (Sem. - II) (Revised Course)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory.*
- 2) *Answer any two questions from Section - I and any two questions from Section - II from the remaining.*
- 3) *Answers to the two sections should be written on separate books.*
- 4) *Figures to the right indicate full marks.*

SECTION - I

- Q1)** What is Sampling? Enumerate different sampling plans. Justify situation and conditional criteria. Where it is applied? Discuss the characteristics of good sampling plan. **[12]**
- Q2)** 'Poor Quality and High Cost' Vs 'Quality improvement and cost reduction' if studied meticulously can help in industrial progress. Comment. **[14]**
- Q3)** Describe and justify the relevant importance of planning to maintain & achieve quality in manufacturing operations. **[14]**
- Q4)** Write short notes on (Any two): **[14]**
- a) Contribution of Prof. Juran, Prof. Deming and Prof. Crosby in Quality field.
 - b) Motivation.
 - c) Quality Surveys.

SECTION - II

- Q5)** Define Audit and explain its scope in improving the quality of different systems and operations. What are the characteristics of a good auditor? Where surprise audit is carried out? **[12]**

P.T.O.

Q6) Explain the term: Inspection, testing & measurement. How will you decide to what extent inspection is necessary and its accuracy? **[14]**

Q7) Define Statistics. Discuss the advantages of Statistics on Process Control. Explain different statistical control charts used in the industry. **[14]**

Q8) Write short notes on (Any two): **[14]**

- a) Sporadic & Chronic Quality problems.
- b) SKIP - LOT Sampling Plan & its Utility.
- c) Quality Improvement Programme (QIP).



Total No. of Questions : 8]

[Total No. of Pages : 2

P1861

[3957]-114

M.Pharmacy

CLINICAL TRIALS

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Question No. 1 and 5 are compulsory. Solve any two questions from the remaining in Section - I and Section - II.*
- 2) *Figures to the right indicate full marks.*
- 3) *Write answers for Section - I and Section - II in separate answer sheets.*

SECTION - I

- Q1)** Discuss collection, monitoring and interpretation of data in clinical trials. **[10]**
- Q2)** Describe ethical issues in clinical trials with special mention of Helsinki declaration. **[15]**
- Q3)** Explain importance of ICH-GCP guidelines in clinical trial. **[15]**
- Q4)** Write a note on (Any Two): **[15]**
- a) Case report forms.
 - b) Role of CRO in clinical trials.
 - c) Computer application in data analysis.

SECTION - II

- Q5)** Enlist various parts of clinical trial design. Add a note on risk and benefit calculations. **[10]**
- Q6)** Name various stakeholders of clinical trials. Discuss responsibilities of physicians. **[15]**

P.T.O.

Q7) Justify role of therapeutic drug monitoring towards quality control in clinical trials. **[15]**

Q8) Write a note on (Any Two): **[15]**

- a) Hypothesis in clinical trial design.
- b) Institutional review board.
- c) NDA.



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Total No. of Questions : 6]

[Total No. of Pages : 2

P1862

[3957]-113

M.Pharmacy

ACTIVE PHARMACEUTICAL INGREDIENTS

MANUFACTURING TECHNOLOGY

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Answer any two questions from Section - I and any two questions from Section - II.*
- 2) *All questions carry equal marks.*
- 3) *Neat diagrams must be drawn wherever is necessary.*

SECTION - I

Q1) Describe the technology of animation by reduction using iron and acid for the manufacturing of aromatic amines, illustrate with examples. **[20]**

Q2) Describe in detail manufacture of following drugs with process and instrumentation diagram (any two): **[20]**

- a) Rifampicin.
- b) Adrenaline.
- c) Benzocaine.

Q3) Write short notes on any two: **[20]**

- a) Fluidized bed dryers.
- b) Industrial centrifuges.
- c) Counter current extractions.

SECTION - II

Q4) Discuss in detail the acylation and esterification process in the manufacture of pharmaceuticals. **[20]**

P.T.O.

- Q5)** a) Give detailed account of Health hazard in manufacturing facility with respect to Bioethics and Bio-safety. [10]
b) Write notes on any two: [10]
i) Atmospheric contaminants.
ii) Detection and sampling.
iii) Environment protection laws related to Pharma Industry.

- Q6)** Write notes on any two: [20]
a) Oxidation in drug synthesis.
b) Crystallizers used in pharmaceuticals.
c) Stoichiometry in drug synthesis.



Total No. of Questions : 6]

[Total No. of Pages : 1

P1863

[3957]-109

M.Pharmacy

PHARMACEUTICAL PLANT DESIGN AND OPERATIONS

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) Answer 2 questions from Section - I and 2 questions from Section - II.*
- 2) Answers to the two sections should be written in separate books.*
- 3) Neat diagrams must be drawn wherever necessary.*

SECTION - I

Q1) Discuss the design, layout and operational facilities for tablets. **[20]**

Q2) Explain in detail regulatory requirements of pharma facilities with reference to revised schedule M & factory act. **[20]**

Q3) Explain in detail design, layout and operational facilities with services and utilities for sterile products powders ready for reconstitution. **[20]**

SECTION - II

Q4) Discuss in detail designing of plant support services. **[20]**

Q5) Discuss in detail design of effluent treatment plant. **[20]**

Q6) Explain the design of utility services as water stream compressed air & other gases. **[20]**



Total No. of Questions : 8]

[Total No. of Pages :1

P1864

[3957]-115

M.Pharmacy

SAFETY PHARMACOLOGY

(2008 Pattern) (Sem. - I & II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) *Q 1 & Q 5 are compulsory.*
- 2) *Attempt any 2 questions from the remaining in each section.*
- 3) *Figures to the right indicate full marks.*

SECTION - I

- Q1)** Define safety Pharmacology. What is its scope? [10]
- Q2)** Enlist the regulatory requirements for the new drug safety assessment as per ICH, OECD and USFDA guidelines. [15]
- Q3)** Write notes on: [15]
- a) Analysis of safety pharmacological data.
 - b) EMEA guidelines on safety assessment of new drug.
- Q4)** Give the principles and study design of acute, sub - acute and chronic toxicities in pre - clinical studies. [15]

SECTION - II

- Q5)** Define Pharmacovigilance. What are its objectives and functions? [10]
- Q6)** Write notes on: [15]
- a) Safety testing for dermatological products.
 - b) Risk - benefit assessment.
- Q7)** Describe the pre - clinical safety studies on genotoxicity, reproductive and ocular toxicity. [15]
- Q8)** What is adverse event monitoring during clinical trials? Discuss the data collection, reporting methods, assessment and analysis of the same. [15]



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Total No. of Questions : 8]

[Total No. of Pages : 1

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[3957]-206

M.Pharmacy.

(Spl. Pharmacology)

CLINICAL PHARMACOLOGY

(2008 Pattern) (Sem. - II)

Time : 3 Hours]

[Max. Marks : 80

Instructions to the candidates:

- 1) Question number 1 and 5 are compulsory. Out of the remaining attempt any two questions from Section-I and two questions from Section-II.*
- 2) Answers to the two Sections should be written in separate book.*
- 3) Figures to the right indicate full marks.*

SECTION - I

- Q1)** Discuss the management of peptic ulcers. **[10]**
- Q2)** Explain the mechanism of resistance to antibiotics. Add a note on measures to minimize the antibiotic resistance. **[15]**
- Q3)** Explain the pharmacotherapy of congestive heart failure. **[15]**
- Q4)** Write notes on : **[15]**
- a) Responsibilities of Investigator.
 - b) Management of coagulation disorder.

SECTION - II

- Q5)** Discuss the process of new drug development process. Add a note on ethics in clinical trial. **[10]**
- Q6)** Describe the management of asthma. **[15]**
- Q7)** Discuss the current concepts in the management of cancer. **[15]**
- Q8)** Write a note on : **[15]**
- a) Renal dialysis.
 - b) Immunosuppressants.

