### Pharmacy Council of India New Delhi

Rules & Syllabus for the Bachelor of Pharmacy (B. Pharm) Course

[Framed under Regulation 6, 7 & 8 of the Bachelor of Pharmacy (B. Pharm) course regulations 2014]



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#### **CHAPTER-I: REGULATIONS**

#### 1. Short Title and Commencement

These regulations shall be called as "The Revised Regulations for the B. Pharm. Degree Program (CBCS)of the Pharmacy Council of India, New Delhi". They shall come into effect from the Academic Year 2016-17. The regulations framed are subject to modifications from time to time by Pharmacy Council of India.

#### 2. Minimum qualification for admission

#### First year B. Pharm:

Candidate shall have passed 10+2 examination conducted by the respective state/central government authorities recognized as equivalent to 10+2 examination by the Association of Indian Universities (AIU) with English as one of the subjects and Physics, Chemistry, Mathematics (P.C.M) and or Biology (P.C.B / P.C.M.B.) as optional subjects individually. Any other qualification approved by the Pharmacy Council of India as equivalent to any of the above examinations.

#### 2.2. B. Pharm lateral entry (to third semester):

A pass in D. Pharm. course from an institution approved by the Pharmacy Council of India under section 12 of the Pharmacy Act.

#### 3. Duration of the program

The course of study for B.Pharm shall extend over a period of eight semesters (four academic years) and six semesters (three academic years) for lateral entry students. The curricula and syllabi for the program shall be prescribed from time to time by Pharmacy Council of India, New Delhi.

#### 4. Medium of instruction and examinations

Medium of instruction and examination shall be in English.

#### 5. Working days in each semester

Each semestershall consist of not less than 100 working days. The odd semesters shall be conducted from the month of June/July to November/December and the even semesters shall be conducted from December/January to May/June in every calendar year.

#### 6. Attendance and progress

A candidate is required to put in at least 80% attendance in individual courses considering theory and practical separately. The candidate shall complete the prescribed course satisfactorily to be eligible to appear for the respective examinations.

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#### 7. Program/Course credit structure

As per the philosophy of Credit Based Semester System, certain quantum of academic work viz. theory classes, tutorial hours, practical classes, etc. are measured in terms of credits. On satisfactory completion of the courses, a candidate earns credits. The amount of credit associated with a course is dependent upon the number of hours of instruction per week in that course. Similarly, the credit associated with any of the other academic, co/extra-curricular activities is dependent upon the quantum of work expected to be put in for each of these activities per week.

#### Credit assignment

#### Theory and Laboratory courses

Courses are broadly classified as Theory and Practical. Theory courses consist of lecture (L) and /or tutorial (T) hours, and Practical (P) courses consist of hours spent in the laboratory. Credits (C) for a course is dependent on the number of hours of instruction per week in that course, and is obtained by using a multiplier of one (1) for lecture and tutorial hours, and a multiplier of half (1/2) for practical (laboratory) hours. Thus, for example, a theory course having three lectures and one tutorial per week throughout the semester carries a credit of 4. Similarly, a practical having four laboratory hours per week throughout semester carries a credit of 2.

#### Minimum credit requirements

The minimum credit points required for award of a B. Pharm. degree is 208. These credits are divided into Theory courses, Tutorials, Practical, Practice School and Projectover the duration of eight semesters. The credits are distributed semester-wise as shown in Table IX. Courses generally progress in sequences, building competencies and their positioning indicates certain academic maturity on the part of the learners. Learners are expected to follow the semester-wise schedule of courses given in the syllabus.

The lateral entry students shall get 52 credit points transferred from their D. Pharm program. Such students shall take up additional remedial courses of 'Communication Skills' (Theory and Practical) and 'Computer Applications in Pharmacy' (Theory and Practical) equivalent to 3 and 4 credit points respectively, a total of 7 credit points to attain 59 credit points, the maximum of I and II semesters.

#### 8. Academic work

A regular record of attendance both in Theory and Practical shall be maintained by the teaching staff of respective courses.

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#### 9. Course of study

The course of study for B. Pharm shall include Semester Wise Theory & Practical as given in Table - I to VIII. The number of hours to be devoted to each theory, tutorial and practical course in any semester shall not be less than that shown in Table - I to VIII.

Table-I: Course of study for semester I

	1 able-1: Course of study for s	emester 1		
Course code	Name of the course	No. of hours	Tuto rial	Credit points
BP101T	Human Anatomy and Physiology I— Theory	3	1	4
BP102T	Pharmaceutical Analysis I – Theory	3	1	4
BP103T	Pharmaceutics I – Theory	3	1	4
BP104T	Pharmaceutical Inorganic Chemistry – Theory	3	1	4
BP105T	Communication skills – Theory *	2	-	2
BP106RBT BP106RMT	Remedial Biology/ Remedial Mathematics – Theory*	2 .	-	2
BP107P	Human Anatomy and Physiology – Practical	4	-	2
BP108P	Pharmaceutical Analysis I – Practical	4	-	2
BP109P	Pharmaceutics I – Practical	4	-	2
BP110P	Pharmaceutical Inorganic Chemistry – Practical	4	-	2
BP111P	Communication skills – Practical*	2	-	1
BP112RBP	Remedial Biology – Practical*	2	-	1
	Total	32/34 <sup>\$</sup> /36 <sup>#</sup>	4	27/29 <sup>\$</sup> /30 <sup>#</sup>

<sup>\*</sup>Applicable ONLY for the students who have studied Mathematics / Physics / Chemistry at HSC and appearing for Remedial Biology (RB)course.

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<sup>&</sup>lt;sup>\$</sup>Applicable ONLY for the students who have studied Physics / Chemistry / Botany / Zoology at HSC and appearing for Remedial Mathematics (RM)course.

<sup>\*</sup> Non University Examination (NUE)

Table-VIII: Course of study for semester VIII

Course code	Name of the course	No. of hours	Tutorial	Credit points
BP801T	Biostatistics and Research Methodology	3	1	4
BP802T	Social and Preventive Pharmacy	3	1	4
BP803ET	Pharma Marketing Management			
BP804ET	Pharmaceutical Regulatory Science			
BP805ET	Pharmacovigilance			
BP806ET	Quality Control and Standardization of Herbals	3 + 3 =	1 + 1 = 2	4 + 4 =
BP807ET	Computer Aided Drug Design	6		8
BP808ET	Cell and Molecular Biology			
BP809ET	Cosmetic Science			
BP810ET	Experimental Pharmacology			
BP811ET	Advanced Instrumentation Techniques			
BP812ET	Dietary Supplements and Nutraceuticals			
BP813PW	Project Work	12	-	6
	Total	24	4	22

Table-IX: Semester wise credits distribution

Semester	Credit Points
I	27/29 <sup>\$</sup> /30 <sup>#</sup>
II	29
III	26
IV	28
V	26
VI	26
VII	24
VIII	22
Extracurricular/ Co curricular activities	01*
Total credit points for the program	209/211 <sup>\$</sup> /212 <sup>#</sup>

<sup>\*</sup> The credit points assigned for extracurricular and or co-curricular activities shall be given by the Principals of the colleges and the same shall be submitted to the University. The criteria to acquire this credit point shall be defined by the colleges from time to time.

\*Applicable ONLY for the students studied Mathematics / Physics / Chemistry at HSC and appearing for Remedial Biology course.

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<sup>\$</sup>Applicable ONLY for the students studied Physics / Chemistry / Botany / Zoology at HSC and appearing for Remedial Mathematics course.

Tables-X: Schemes for internal assessments and end semester examinations semester wise

## Semester I

<sup>\*</sup>Applicable ONLY for the students studied Mathematics / Physics / Chemistry at HSC and appearing for Remedial Biology (RB)course.



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<sup>&</sup>lt;sup>\$</sup>Applicable ONLY for the students studied Physics / Chemistry / Botany / Zoology at HSC and appearing for Remedial Mathematics (RM)course.

<sup>\*</sup> Non University Examination (NUE)

## Semester VIII

BP812PW   Proj	BP811ET Adv	BP810ET Experii	BP809ET Cos	BP808ET Cell an Theory	BP807ET Compu	Theory	BP806ET Star	Qua	BP805ET Phai	- 4	RD80AFT Phai	Theory	RP803FT Phai	BP802T Soci	BP801T Bios	0000	code	Course
Project Work	Advanced Instrumentation Techniques – Theory	Experimental Pharmacology – Theory	Cosmetic Science – Theory	Cell and Molecular Biology – Theory	Computer Aided Drug Design – Theory	ory	Standardization of Herbals –	Quality Control and	Pharmacovigilance - Theory	Science - Theory	Pharmaceutical Regulatory	ory	Pharmaceutical Marketing -	Social and Preventive Pharmacy  - Theory	Biostatistics and Research Methodology – Theory		Name of the course	
1					= 20	10 + 10								10	10	Mode	Continuous	
1					30	15 + 15 =								15	15	Marks	Sessional Exams	Internal Assessment
•					2 Hrs	1+1=								1 Hr	1 Hr	Duration	l Exams	sessment
1					30	25 + 25 =								25	25	I OLAI	Total	
150					001 =	75 + 75								75	75	INTALL	Mark	<b>End Seme</b>
4 Hrs					1113	3+3=0	) - )							3 Hrs	3 Hrs	Duranon	Duration	End Semester Exams
150					200	100+								100	100	LYLGE INS	Marks	Total

No.		トイイン	くつグ	
	\			

Total

40

60

4 Hrs

100

450

16 Hrs

550

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#### BP 106RBT.REMEDIAL BIOLOGY (Theory)

30 Hours

**Scope:** To learn and understand the components of living world, structure and functional system of plant and animal kingdom.

Objectives: Upon completion of the course, the student shall be able to

- know the classification and salient features of five kingdoms of life
- understand the basic components of anatomy & physiology of plant
- know understand the basic components of anatomy & physiology animal with special reference to human

UNIT I 07 Hours

#### Living world:

- Definition and characters of living organisms
- Diversity in the living world
- Binomial nomenclature
- Five kingdoms of life and basis of classification. Salient features of Monera, Potista, Fungi, Animalia and Plantae, Virus,

#### Morphology of Flowering plants

- Morphology of different parts of flowering plants Root, stem, inflorescence, flower, leaf, fruit, seed.
- General Anatomy of Root, stem, leaf of monocotyledons & Dicotylidones.

UNIT II 07 Hours

#### Body fluids and circulation

- Composition of blood, blood groups, coagulation of blood
- Composition and functions of lymph
- Human circulatory system
- Structure of human heart and blood vessels
- Cardiac cycle, cardiac output and ECG

#### **Digestion and Absorption**

- Human alimentary canal and digestive glands
- Role of digestive enzymes
- Digestion, absorption and assimilation of digested food

#### Breathing and respiration

- Human respiratory system
- Mechanism of breathing and its regulation
- Exchange of gases, transport of gases and regulation of respiration
- Respiratory volumes

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UNIT III 07 Hours

#### Excretory products and their elimination

- Modes of excretion
- Human excretory system- structure and function
- Urine formation
- Rennin angiotensin system

#### Neural control and coordination

- Definition and classification of nervous system
- Structure of a neuron
- Generation and conduction of nerve impulse
- Structure of brain and spinal cord
- Functions of cerebrum, cerebellum, hypothalamus and medulla oblongata

#### Chemical coordination and regulation

- Endocrine glands and their secretions
- Functions of hormones secreted by endocrine glands

#### **Human reproduction**

- Parts of female reproductive system
- Parts of male reproductive system
- Spermatogenesis and Oogenesis
- Menstrual cycle

UNIT IV 05 Hours

#### Plants and mineral nutrition:

- Essential mineral, macro and micronutrients
- Nitrogen metabolism, Nitrogen cycle, biological nitrogen fixation

#### **Photosynthesis**

• Autotrophic nutrition, photosynthesis, Photosynthetic pigments, Factors affecting photosynthesis.

UNIT V 04 Hours

Plant respiration: Respiration, glycolysis, fermentation (anaerobic).

#### Plant growth and development

• Phases and rate of plant growth, Condition of growth, Introduction to plant growth regulators

#### Cell - The unit of life

• Structure and functions of cell and cell organelles. Cell division

#### **Tissues**

• Definition, types of tissues, location and functions.

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#### **Text Books**

- a. Text book of Biology by S. B. Gokhale
- b. A Text book of Biology by Dr. Thulajappa and Dr. Seetaram.

#### **Reference Books**

- a. A Text book of Biology by B.V. Sreenivasa Naidu
- b. A Text book of Biology by Naidu and Murthy
- c. Botany for Degree students By A.C.Dutta.
- d.Outlines of Zoology by M. Ekambaranatha ayyer and T. N. Ananthakrishnan.
- e. A manual for pharmaceutical biology practical by S.B. Gokhale and C. K. Kokate

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#### BP112RBP.REMEDIAL BIOLOGY (Practical)

30 Hours

- 1. Introduction to experiments in biology
  - a) Study of Microscope
  - b) Section cutting techniques
  - c) Mounting and staining
  - d) Permanent slide preparation
- 2. Study of cell and its inclusions
- 3. Study of Stem, Root, Leaf, seed, fruit, flower and their modifications
- 4. Detailed study of frog by using computer models
- 5. Microscopic study and identification of tissues pertinent to Stem, Root Leaf, seed, fruit and flower
- 6. Identification of bones
- 7. Determination of blood group
- 8. Determination of blood pressure
- 9. Determination of tidal volume

#### **Reference Books**

- 1. Practical human anatomy and physiology. by S.R.Kale and R.R.Kale.
- 2. A Manual of pharmaceutical biology practical by S.B.Gokhale, C.K.Kokate and S.P.Shriwastava.
- 3. Biology practical manual according to National core curriculum .Biology forum of Karnataka. Prof .M.J.H.Shafi

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#### BP 106RMT.REMEDIAL MATHEMATICS (Theory)

30 Hours

**Scope:** This is an introductory course in mathematics. This subject deals with the introduction to Partial fraction, Logarithm, matrices and Determinant, Analytical geometry, Calculus, differential equation and Laplace transform.

Objectives: Upon completion of the course the student shall be able to:-

- 1. Know the theory and their application in Pharmacy
- 2. Solve the different types of problems by applying theory
- 3. Appreciate the important application of mathematics in Pharmacy

#### **Course Content:**

UNIT – I 06 Hours

#### • Partial fraction

Introduction, Polynomial, Rational fractions, Proper and Improper fractions, Partial fraction, Resolving into Partial fraction, Application of Partial Fraction in Chemical Kinetics and Pharmacokinetics

#### Logarithms

Introduction, Definition, Theorems/Properties of logarithms, Common logarithms, Characteristic and Mantissa, worked examples, application of logarithm to solve pharmaceutical problems.

#### • Function:

Real Valued function, Classification of real valued functions,

#### • Limits and continuity:

Introduction , Limit of a function, Definition of limit of a function ( $\in$  -  $\delta$ 

definition), 
$$\lim_{x\to a} \frac{x^n - a^n}{x - a} = na^{n-1}$$
,  $\lim_{\theta \to 0} \frac{\sin \theta}{\theta} = 1$ ,

UNIT –II 06 Hours

#### • Matrices and Determinant:

Introduction matrices, Types of matrices, Operation on matrices, Transpose of a matrix, Matrix Multiplication, Determinants, Properties of determinants, Product of determinants, Minors and co-Factors, Adjoint or adjugate of a square matrix, Singular and non-singular matrices, Inverse of a matrix, Solution of system of linear of equations using matrix method, Cramer's rule, Characteristic equation and roots of a square matrix, Cayley–Hamilton theorem, Application of Matrices in solving Pharmacokinetic equations

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UNIT – III 06 Hours

Calculus

**Differentiation**: Introductions, Derivative of a function, Derivative of a constant, Derivative of a product of a constant and a function, Derivative of the sum or difference of two functions, Derivative of the product of two functions (product formula), Derivative of the quotient of two functions (Quotient formula) – **Without Proof**, Derivative of  $x^n$  w.r.tx, where n is any rational number, Derivative of  $e^x$ , Derivative of  $\log_e x$ , Derivative of  $e^x$ , Derivative of trigonometric functions from first principles (without **Proof**), Successive Differentiation, Conditions for a function to be a maximum or a minimum at a point. Application

UNIT – IV

06 Hours

Analytical Geometry

Introduction: Signs of the Coordinates, Distance formula,

**Straight Line**: Slope or gradient of a straight line, Conditions for parallelism and perpendicularity of two lines, Slope of a line joining two points, Slope – intercept form of a straight line

Integration:

Introduction, Definition, Standard formulae, Rules of integration, Method of substitution, Method of Partial fractions, Integration by parts, definite integrals, application

UNIT-V 06 Hours

- **Differential Equations**: Some basic definitions, Order and degree, Equations in separable form, Homogeneous equations, Linear Differential equations, Exact equations, **Application in solving Pharmacokinetic equations**
- Laplace Transform: Introduction, Definition, Properties of Laplace transform, Laplace Transforms of elementary functions, Inverse Laplace transforms, Laplace transform of derivatives, Application to solve Linear differential equations, Application in solving Chemical kinetics and Pharmacokinetics equations

#### **Recommended Books (Latest Edition)**

- 1. Differential Calculus by Shanthinarayan
- 2. Pharmaceutical Mathematics with application to Pharmacy by Panchaksharappa Gowda D.H.
- 3. Integral Calculus by Shanthinarayan
- 4. Higher Engineering Mathematics by Dr.B.S.Grewal

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#### BP803ET. PHARMA MARKETING MANAGEMENT (Theory)

45 Hours

#### Scope:

The pharmaceutical industry not only needs highly qualified researchers, chemists and, technical people, but also requires skilled managers who can take the industry forward by managing and taking the complex decisions which are imperative for the growth of the industry. The Knowledge and Know-how of marketing management groom the people for taking a challenging role in Sales and Product management.

Course Objective: The course aims to provide an understanding of marketing concepts and techniques and their applications in the pharmaceutical industry.

Unit I 10 Hours

#### Marketing:

Definition, general concepts and scope of marketing; Distinction between marketing & selling; Marketing environment; Industry and competitive analysis; Analyzing consumer buying behavior; industrial buying behavior.

#### Pharmaceutical market:

Quantitative and qualitative aspects; size and composition of the market; demographic descriptions and socio-psychological characteristics of the consumer; market segmentation& targeting.Consumer profile; Motivation and prescribing habits of the physician; patients' choice of physician and retail pharmacist.Analyzing the Market;Role of market research.

Unit II 10 Hours

#### **Product decision:**

Classification, product line and product mix decisions, product life cycle,product portfolio analysis; product positioning; New product decisions; Product branding, packaging and labeling decisions, Product management in pharmaceutical industry.

Unit III 10 Hours

#### **Promotion:**

Methods, determinants of promotional mix, promotional budget; An overview of personal selling, advertising, direct mail, journals, sampling, retailing, medical exhibition, public relations, online promotional techniques for OTC Products.

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Unit IV 10 Hours

#### Pharmaceutical marketing channels:

Designing channel, channel members, selecting the appropriate channel, conflict in channels, physical distribution management: Strategic importance, tasks in physical distribution management.

#### Professional sales representative (PSR):

Duties of PSR, purpose of detailing, selection and training, supervising, norms for customer calls, motivating, evaluating, compensation and future prospects of the PSR.

Unit V 10 Hours

#### **Pricing:**

Meaning, importance, objectives, determinants of price; pricing methods and strategies, issues in price management in pharmaceutical industry. An overview of DPCO (Drug Price Control Order) and NPPA (National Pharmaceutical Pricing Authority).

#### **Emerging concepts in marketing:**

Vertical & Horizontal Marketing; RuralMarketing; Consumerism; Industrial Marketing; Global Marketing.

#### **Recommended Books: (Latest Editions)**

- 1. Philip Kotler and Kevin Lane Keller: Marketing Management, Prentice Hall of India, New Delhi
- 2. Walker, Boyd and Larreche: Marketing Strategy- Planning and Implementation, Tata MC GrawHill, New Delhi.
- 3. Dhruv Grewal and Michael Levy: Marketing, Tata MC Graw Hill
- 4. Arun Kumar and N Menakshi: Marketing Management, Vikas Publishing, India
- 5. Rajan Saxena: Marketing Management; Tata MC Graw-Hill (India Edition)
- 6. Ramaswamy, U.S & Nanakamari, S: Marketing Managemnt:Global Perspective, IndianContext,Macmilan India, New Delhi.
- 7. Shanker, Ravi: Service Marketing, Excell Books, New Delhi
- 8. Subba Rao Changanti, Pharmaceutical Marketing in India (GIFT Excel series) Excel Publications.

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#### BP804 ET: PHARMACEUTICAL REGULATORY SCIENCE (Theory)

45Hours

**Scope:** This course is designed to impart the fundamental knowledge on the regulatory requirements for approval of new drugs, and drug products in regulated markets of India & other countries like US, EU, Japan, Australia, UK etc. It prepares the students to learn in detail on the regulatory requirements, documentation requirements, and registration procedures for marketing the drug products.

Objectives: Upon completion of the subject student shall be able to;

- 1. Know about the process of drug discovery and development
- 2. Know the regulatory authorities and agencies governing the manufacture and sale of pharmaceuticals
- 3. Know the regulatory approval process and their registration in Indian and international markets

#### **Course content:**

Unit I 10Hours

#### New Drug Discovery and development

Stages of drug discovery, Drug development process, pre-clinical studies, non-clinical activities, clinical studies, Innovator and generics, Concept of generics, Generic drug product development.

Unit II 10Hours

#### **Regulatory Approval Process**

Approval processes and timelines involved in Investigational New Drug (IND), New Drug Application (NDA), Abbreviated New Drug Application (ANDA). Changes to an approved NDA / ANDA.

#### Regulatory authorities and agencies

Overview of regulatory authorities of India, United States, European Union, Australia, Japan, Canada (Organization structure and types of applications)

Unit III 10Hours

#### Registration of Indian drug product in overseas market

Procedure for export of pharmaceutical products, Technical documentation, Drug Master Files (DMF), Common Technical Document (CTD), electronic Common Technical



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Document (eCTD), ASEAN Common Technical Document (ACTD)research.

Unit IV 08Hours

#### Clinical trials

Developing clinical trial protocols, Institutional Review Board / Independent Ethics committee - formation and working procedures, Informed consent process and procedures, GCP obligations of Investigators, sponsors & Monitors, Managing and Monitoring clinical trials, Pharmacovigilance - safety monitoring in clinical trials

Unit V 07Hours

#### **Regulatory Concepts**

Basic terminology, guidance, guidelines, regulations, Laws and Acts, Orange book, Federal Register, Code of Federal Regulatory, Purple book

#### Recommended books (Latest edition):

1. Drug Regulatory Affairs by Sachin Itkar, Dr. N.S. Vyawahare, Nirali Prakashan.

- 2. The Pharmaceutical Regulatory Process, Second Edition Edited by Ira R. Berry and Robert P. Martin, Drugs and the Pharmaceutical Sciences, Vol. 185. Informa Health care Publishers.
- 3. New Drug Approval Process: Accelerating Global Registrations By Richard A Guarino, MD, 5<sup>th</sup> edition, Drugs and the Pharmaceutical Sciences, Vol. 190.
- 4. Guidebook for drug regulatory submissions / Sandy Weinberg. By John Wiley & Sons. Inc.
- 5. FDA Regulatory Affairs: a guide for prescription drugs, medical devices, and biologics /edited by Douglas J. Pisano, David Mantus.
- 6. Generic Drug Product Development, Solid Oral Dosage forms, Leon Shargel and Isader Kaufer, Marcel Dekker series, Vol.143
- 7. Clinical Trials and Human Research: A Practical Guide to Regulatory Compliance By Fay A. Rozovsky and Rodney K. Adams
- 8. Principles and Practices of Clinical Research, Second Edition Edited by John I. Gallin and Frederick P. Ognibene
- 9. Drugs: From Discovery to Approval, Second Edition By Rick Ng

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#### BP 805T: PHARMACOVIGILANCE (Theory)

45 hours

**Scope:** This paper will provide an opportunity for the student to learn about development of pharmacovigilance as a science, basic terminologies used in pharmacovigilance, global scenario of Pharmacovigilance, train students on establishing pharmacovigilance programme in an organization, various methods that can be used to generate safety data and signal detection. This paper also develops the skills of classifying drugs, diseases and adverse drug reactions.

#### **Objectives:**

At completion of this paper it is expected that students will be able to (know, do, and appreciate):

- 1. Why drug safety monitoring is important?
- 2. History and development of pharmacovigilance
- 3. National and international scenario of pharmacovigilance
- 4. Dictionaries, coding and terminologies used in pharmacovigilance
- 5. Detection of new adverse drug reactions and their assessment
- 6. International standards for classification of diseases and drugs
- 7. Adverse drug reaction reporting systems and communication in pharmacovigilance
- 8. Methods to generate safety data during pre clinical, clinical and post approval phases of drugs' life cycle
- 9. Drug safety evaluation in paediatrics, geriatrics, pregnancy and lactation
- 10. Pharmacovigilance Program of India (PvPI) requirement for ADR reporting in India
- 11. ICH guidelines for ICSR, PSUR, expedited reporting, pharmacovigilance planning
- 12. CIOMS requirements for ADR reporting
- 13. Writing case narratives of adverse events and their quality.

#### **Course Content**

Unit I 10 Hours

#### Introduction to Pharmacovigilance

- History and development of Pharmacovigilance
- Importance of safety monitoring of Medicine
- WHO international drug monitoring programme
- Pharmacovigilance Program of India(PvPI)

#### Introduction to adverse drug reactions

- Definitions and classification of ADRs
- Detection and reporting
- Methods in Causality assessment
- Severity and seriousness assessment
- Predictability and preventability assessment
- Management of adverse drug reactions

Basic terminologies used in pharmacovigilance

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- Terminologies of adverse medication related events
- Regulatory terminologies

#### **Unit II**

10 hours

#### Drug and disease classification

- Anatomical, therapeutic and chemical classification of drugs
- International classification of diseases
- Daily defined doses
- International Non proprietary Names for drugs

#### Drug dictionaries and coding in pharmacovigilance

- WHO adverse reaction terminologies
- MedDRA and Standardised MedDRA queries
- WHO drug dictionary
- Eudravigilance medicinal product dictionary

#### Information resources in pharmacovigilance

- Basic drug information resources
- Specialised resources for ADRs

#### Establishing pharmacovigilance programme

- Establishing in a hospital
- Establishment & operation of drug safety department in industry
- Contract Research Organisations (CROs)
- Establishing a national programme

#### **Unit III**

10 Hours

#### Vaccine safety surveillance

- Vaccine Pharmacovigilance
- Vaccination failure
- Adverse events following immunization

#### Pharmacovigilance methods

- Passive surveillance Spontaneous reports and case series
- Stimulated reporting
- Active surveillance Sentinel sites, drug event monitoring and registries
- Comparative observational studies Cross sectional study, case control study and cohort study
- Targeted clinical investigations

#### Communication in pharmacovigilance

- Effective communication in Pharmacovigilance
- Communication in Drug Safety Crisis management
- Communicating with Regulatory Agencies, Business Partners, Healthcare facilities & Media

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#### Safety data generation

- Pre clinical phase
- Clinical phase
- Post approval phase (PMS)

#### ICH Guidelines for Pharmacovigilance

- Organization and objectives of ICH
- Expedited reporting
- Individual case safety reports
- Periodic safety update reports
- Post approval expedited reporting
- Pharmacovigilance planning
- Good clinical practice in pharmacovigilance studies

#### Unit V

7 hours

#### Pharmacogenomics of adverse drug reactions

• Genetics related ADR with example focusing PK parameters.

#### Drug safety evaluation in special population

- Paediatrics
- Pregnancy and lactation
- Geriatrics

#### **CIOMS**

- CIOMS Working Groups
- CIOMS Form

#### CDSCO (India) and Pharmacovigilance

- D&C Act and Schedule Y
- Differences in Indian and global pharmacovigilance requirements

#### Recommended Books (Latest edition):

- 1. Textbook of Pharmacovigilance: S K Gupta, Jaypee Brothers, Medical Publishers.
- 2. Practical Drug Safety from A to Z By Barton Cobert, Pierre Biron, Jones and Bartlett Publishers.
- 3. Mann's Pharmacovigilance: Elizabeth B. Andrews, Nicholas, Wiley Publishers.
- 4. Stephens' Detection of New Adverse Drug Reactions: John Talbot, Patrick Walle, Wiley Publishers.
- 5. An Introduction to Pharmacovigilance: Patrick Waller, Wiley Publishers.
- 6. Cobert's Manual of Drug Safety and Pharmacovigilance: Barton Cobert, Jones & Bartlett Publishers.
- 7. Textbook of Pharmacoepidemiolog edited by Brian L. Strom, Stephen E Kimmel, Sean Hennessy, Wiley Publishers.
- 8. A Textbook of Chical Pharmacy Practice -Essential Concepts and Skills:G. Parthasarathi, Karin NyfortHansen, Milap C. Nahata
- 9. National Formulary of India
- 10. Text Book of Medicine by Yashpal Munjal

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#### BP 806 ET. QUALITY CONTROL AND STANDARDIZATION OF HERBALS (Theory)

**Scope:** In this subject the student learns about the various methods and guidelines for evaluation and standardization of herbs and herbal drugs. The subject also provides an opportunity for the student to learn cGMP, GAP and GLP in traditional system of medicines.

Objectives: Upon completion of the subject student shall be able to;

- 1. know WHO guidelines for quality control of herbal drugs
- 2. know Quality assurance in herbal drug industry
- 3. know the regulatory approval process and their registration in Indian and international markets
- 4. appreciate EU and ICH guidelines for quality control of herbal drugs

Unit I 10 hours

Basic tests for drugs – Pharmaceutical substances, Medicinal plants materials and dosage forms

WHO guidelines for quality control of herbal drugs.

Evaluation of commercial crude drugs intended for use

Unit II 10 hours

Quality assurance in herbal drug industry of cGMP, GAP, GMP and GLP in traditional system of medicine.

WHO Guidelines on current good manufacturing Practices (cGMP) for Herbal Medicines WHO Guidelines on GACP for Medicinal Plants.

Unit III 10 hours

EU and ICH guidelines for quality control of herbal drugs.

Research Guidelines for Evaluating the Safety and Efficacy of Herbal Medicines

Unit IV 08 hours

Stability testing of herbal medicines. Application of various chromatographic techniques in standardization of herbal products.

Preparation of documents for new drug application and export registration GMP requirements and Drugs & Cosmetics Act provisions.

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Unit V 07 hours

Regulatory requirements for herbal medicines.

WHO guidelines on safety monitoring of herbal medicines in pharmacovigilance systems Comparison of various Herbal Pharmacopoeias.

Role of chemical and biological markers in standardization of herbal products

#### Recommended Books: (Latest Editions

- 1. Pharmacognosy by Trease and Evans
- 2. Pharmacognosy by Kokate, Purohit and Gokhale
- 3. Rangari, V.D., Text book of Pharmacognosy and Phytochemistry Vol. I, Carrier Pub., 2006.
- 4. Aggrawal, S.S., Herbal Drug Technology. Universities Press, 2002.
- 5. EMEA. Guidelines on Quality of Herbal Medicinal Products/Traditional Medicinal Products,
- 6. Mukherjee, P.W. Quality Control of Herbal Drugs: An Approach to Evaluation of Botanicals. Business Horizons Publishers, New Delhi, India, 2002.
- 7. Shinde M.V., Dhalwal K., Potdar K., Mahadik K. Application of quality control principles to herbal drugs. International Journal of Phytomedicine 1(2009); p. 4-8.
- 8. WHO. Quality Control Methods for Medicinal Plant Materials, World Health Organization, Geneva, 1998. WHO. Guidelines for the Appropriate Use of Herbal Medicines. WHO Regional Publications, Western Pacific Series No 3, WHO Regional office for the Western Pacific, Manila, 1998.
- 9. WHO. The International Pharmacopeia, Vol. 2: Quality Specifications, 3rd edn. World Health Organization, Geneva, 1981.
- 10. WHO. Quality Control Methods for Medicinal Plant Materials. World Health Organization, Geneva, 1999.
- 11. WHO. WHO Global Atlas of Traditional, Complementary and Alternative Medicine. 2 vol. set. Vol. 1 contains text and Vol. 2, maps. World Health Organization, Geneva, 2005.
- 12. WHO. Guidelines on Good Agricultural and Collection Practices (GACP) for Medicinal Plants. World Health Organization, Geneva, 2004.

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#### BP 807 ET. COMPUTER AIDED DRUG DESIGN (Theory)

45 Hours

Scope: This subject is designed to provide detailed knowledge of rational drug design process and various techniques used in rational drug design process.

Objectives: Upon completion of the course, the student shall be able to understand

- Design and discovery of lead molecules
- The role of drug design in drug discovery process
- The concept of QSAR and docking
- Various strategies to develop new drug like molecules.
- The design of new drug molecules using molecular modeling software

#### **Course Content:**

**UNIT-I** 

10 Hours

#### Introduction to Drug Discovery and Development

Stages of drug discovery and development

#### Lead discovery and Analog Based Drug Design

Rational approaches to lead discovery based on traditional medicine, Random screening, Non-random screening, serendipitous drug discovery, lead discovery based on drug metabolism, lead discovery based on clinical observation.

Analog Based Drug Design: Bioisosterism, Classification, Bioisosteric replacement. Any three case studies

**UNIT-II** 

10 Hours

#### Quantitative Structure Activity Relationship (OSAR)

SAR versus QSAR, History and development of QSAR, Types of physicochemical parameters, experimental and theoretical approaches for the determination of physicochemical parameters such as Partition coefficient, Hammet's substituent constant and Tafts steric constant. Hansch analysis, Free Wilson analysis, 3D-QSAR approaches like COMFA and COMSIA.

**UNIT-III** 

10 Hours

#### Molecular Modeling and virtual screening techniques

Virtual Screening techniques: Drug likeness screening, Concept of pharmacophore mapping and pharmacophore based Screening,

Molecular docking: Rigid docking, flexible docking, manual docking, Docking based screening. De novo drug design.

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UNIT-IV 08 Hours

#### Informatics & Methods in drug design

Introduction to Bioinformatics, chemoinformatics. ADME databases, chemical, biochemical and pharmaceutical databases.

UNIT-V 07 Hours

**Molecular Modeling:** Introduction to molecular mechanics and quantum mechanics. Energy Minimization methods and Conformational Analysis, global conformational minima determination.

#### **Recommended Books (Latest Editions)**

- 1. Robert GCK, ed., "Drug Action at the Molecular Level" University Prak Press Baltimore.
- 2. Martin YC. "Quantitative Drug Design" Dekker, New York.
- 3. Delgado JN, Remers WA eds "Wilson & Gisvolds's Text Book of Organic Medicinal & Pharmaceutical Chemistry" Lippincott, New York.
- 4. Foye WO "Principles of Medicinal chemistry 'Lea & Febiger.
- 5. Koro lkovas A, Burckhalter JH. "Essentials of Medicinal Chemistry" Wiley Interscience.
- 6. Wolf ME, ed "The Basis of Medicinal Chemistry, Burger's Medicinal Chemistry" John Wiley & Sons, New York.
- 7. Patrick Graham, L., An Introduction to Medicinal Chemistry, Oxford University Press.
- 8. Smith HJ, Williams H, eds, "Introduction to the principles of Drug Design" Wright Boston.
- 9. Silverman R.B. "The organic Chemistry of Drug Design and Drug Action" Academic Press New York.

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#### BP808ET: CELL AND MOLECULAR BIOLOGY (Elective subject)

45 Hours

#### Scope:

- Cell biology is a branch of biology that studies cells their physiological properties, their structure, the organelles they contain, interactions with their environment, their life cycle, division, death and cell function.
- This is done both on a microscopic and molecular level.
- Cell biology research encompasses both the great diversity of single-celled organisms like bacteria and protozoa, as well as the many specialized cells in multi-cellular organisms such as humans, plants, and sponges.

**Objectives:** Upon completion of the subject student shall be able to;

- Summarize cell and molecular biology history.
- Summarize cellular functioning and composition.
- Describe the chemical foundations of cell biology.
- Summarize the DNA properties of cell biology.
- Describe protein structure and function.
- Describe cellular membrane structure and function.
- Describe basic molecular genetic mechanisms.
- Summarize the Cell Cycle

#### **Course content:**

Unit I 10Hours

- a) Cell and Molecular Biology: Definitions theory and basics and Applications.
- b) Cell and Molecular Biology: History and Summation.
- c) Properties of cells and cell membrane.
- d) Prokaryotic versus Eukaryotic
- e) Cellular Reproduction
- f) Chemical Foundations an Introduction and Reactions (Types)

Unit II 10 Hours

- a) DNA and the Flow of Molecular Information
- b) DNA Functioning
- c) DNA and RNA
- d) Types of RNA
- e) Transcription and Translation

Unit III 10 Hours

- a) Proteins: Defined and Amino Acids
- b) Protein Structure

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- c) Regularities in Protein Pathways
- d) Cellular Processes
- e) Positive Control and significance of Protein Synthesis

**Unit IV** 

08 Hours

- a) Science of Genetics
- b) Transgenics and Genomic Analysis
- c) Cell Cycle analysis
- d) Mitosis and Meiosis
- e) Cellular Activities and Checkpoints

Unit V

07 Hours

- a) Cell Signals: Introduction
- b) Receptors for Cell Signals
- c) Signaling Pathways: Overview
- d) Misregulation of Signaling Pathways
- e) Protein-Kinases: Functioning

#### Recommended Books (latest edition):

- 1. W.B. Hugo and A.D. Russel: Pharmaceutical Microbiology, Blackwell Scientific publications, Oxford London.
- 2. Prescott and Dunn., Industrial Microbiology, 4<sup>th</sup> edition, CBS Publishers & Distributors, Delhi.
- 3. Pelczar, Chan Kreig, Microbiology, Tata McGraw Hill edn.
- 4. Malcolm Harris, Balliere Tindall and Cox: Pharmaceutical Microbiology.
- 5. Rose: Industrial Microbiology.
- 6. Probisher, Hinsdill et al: Fundamentals of Microbiology, 9th ed. Japan
- 7. Cooper and Gunn's: Tutorial Pharmacy, CBS Publisher and Distribution.
- 8. Peppler: Microbial Technology.
- 9. Edward: Fundamentals of Microbiology.
- 10. N.K.Jain: Pharmaceutical Microbiology, Vallabh Prakashan, Delhi
- 11. Bergeys manual of systematic bacteriology, Williams and Wilkins- A Waverly company
- 12. B.R. Glick and J.J. Pasternak: Molecular Biotechnology: Principles and Applications of RecombinantDNA: ASM Press Washington D.C.
- 13. RA Goldshy et. al., : Kuby Immunology.

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#### BP809ET. COSMETIC SCIENCE(Theory)

45Hours

UNITI 10Hours

Classification of cosmetic and cosmeceutical products

Definition of cosmetics as per Indian and EU regulations, Evolution of cosmeceuticals from cosmetics, cosmetics as quasi and OTC drugs

Cosmetic excipients: Surfactants, rheology modifiers, humectants, emollients,

preservatives. Classification and application

Skin: Basic structure and function of skin.

Hair: Basic structure of hair. Hair growth cycle.

Oral Cavity: Common problem associated with teeth and gums.

**UNIT II** 10 Hours

Principles of formulation and building blocks of skin care products:

Face wash,

Moisturizing cream, Cold Cream, Vanishing cream and their advantages and disadvantages. Application of these products in formulation of cosmecuticals.

Antiperspants & deodorants- Actives & mechanism of action.

Principles of formulation and building blocks of Hair care products:

Conditioning shampoo, Hair conditioner, anti-dandruff shampoo.

Hair oils.

Chemistry and formulation of Para-phylene diamine based hair dye.

Principles of formulation and building blocks of oral care products:

Toothpaste for bleeding gums, sensitive teeth. Teeth whitening, Mouthwash.

**UNIT III** 10 Hours

Sun protection, Classification of Sunscreens and SPF.

Role of herbs in cosmetics:

Skin Care: Aloe and turmeric Hair care: Henna and amla. Oral care: Neem and clove

Analytical cosmetics: BIS specification and analytical methods for shampoo, skin-

cream and toothpaste.

**UNIT IV** 08 Hours.

Principles of Cosmetic Evaluation: Principles of sebumeter, corneometer. Measurement of TEWL, Skin Color, Hair tensile strength, Hair combing properties Soaps, and syndet bars. Evolution and skin benfits.

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**UNIT V** 07 Hours

Oily and dry skin, causes leading to dry skin, skin moisturisation. Basic understanding of the terms Comedogenic, dermatitis.

Cosmetic problems associated with Hair and scalp: Dandruff, Hair fall causes Cosmetic problems associated with skin: blemishes, wrinkles, acne, prickly heat and body odor.

Antiperspirants and Deodorants- Actives and mechanism of action

#### References

- 1) Harry's Cosmeticology, Wilkinson, Moore, Seventh Edition, George Godwin.
- 2) Cosmetics Formulations, Manufacturing and Quality Control, P.P. Sharma, 4<sup>th</sup> Edition, Vandana Publications Pvt. Ltd., Delhi.
- 3) Text book of cosmelicology by Sanju Nanda & Roop K. Khar, Tata Publishers.

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#### BP810 ET. PHARMACOLOGICAL SCREENING METHODS

45 Hours

**Scope:** This subject is designed to impart the basic knowledge of preclinical studies in experimental animals including design, conduct and interpretations of results.

#### **Objectives**

Upon completion of the course the student shall be able to,

- Appreciate the applications of various commonly used laboratory animals.
- Appreciate and demonstrate the various screening methods used in preclinical research
- Appreciate and demonstrate the importance of biostatistics and researchmethodology
- Design and execute a research hypothesis independently

Unit –I	08 Hours
Laboratory Animals:	
Study of CPCSEA and OECD guidelines for maintenance, breeding	
and conduct of experiments on laboratory animals, Common lab	
animals: Description and applications of different species and strains	
of animals. Popular transgenic and mutant animals.	
Techniques for collection of blood and common routes of drug	•
administration in laboratory animals, Techniques of blood collection	
and euthanasia.	
Unit –II	10 Hours
Preclinical screening models	
a. Introduction: Dose selection, calculation and conversions,	
preparation of drug solution/suspensions, grouping of animals and	
importance of sham negative and positive control groups.	
Rationale for selection of animal species and sex for the study.	
b. Study of screening animal models for	
Diuretics, nootropics, anti-Parkinson's, antiasthmatics,	
Preclinical screening models: for CNS activity- analgesic,	
antipyretic, anti-inflammatory, general anaesthetics, sedative and	
hypnotics, antipsychotic, antidepressant, antiepileptic,	
antiparkinsonism, alzheimer's disease	



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#### [PUBLISHED IN THE GAZETTE OF INDIA, No.19, PART III, SECTION 4]

Ministry of Health and Family Welfare (Pharmacy Council of India)

New Delhi, 10<sup>th</sup> May, 2008.

#### Pharm.D. Regulations 2008

Regulations framed under section 10 of the Pharmacy Act, 1948 (8 of 1948).

(As approved by the Government of India, Ministry of Health vide, letter No.V.13013/1/2007-PMS, dated the 13<sup>th</sup> March, 2008 and notified by the Pharmacy Council of India).

No.14-126/2007-PCI.— In exercise of the powers conferred by section 10 of the Pharmacy Act, 1948 (8 of 1948), the Pharmacy Council of India, with the approval of the Central Government, hereby makes the following regulations, namely:-

#### **CHAPTER-I**

- 1. Short title and commencement. -(1) These regulations may be called the Pharm.D. Regulations 2008.
  - (2) They shall come into force from the date of their publication in the official Gazette.
- 2. Pharm.D. shall consist of a certificate, having passed the course of study and examination as prescribed in these regulations, for the purpose of registration as a pharmacist to practice the profession under the Pharmacy Act, 1948.

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#### **CHAPTER-II**

- 3. Duration of the course.
  - a) Pharm.D: The duration of the course shall be six academic years (five years of study and one year of internship or residency) full time with each academic year spread over a period of not less than two hundred working days. The period of six years duration is divided into two phases
    - Phase I consisting of First, Second, Third, Fourth and Fifth academic year.
    - Phase II consisting of internship or residency training during sixth year involving posting in speciality units. It is a phase of training wherein a student is exposed to actual pharmacy practice or clinical pharmacy services and acquires skill under supervision so that he or she may become capable of functioning independently.
  - b) Pharm.D. (Post Baccalaureate): The duration of the course shall be for three academic years (two years of study and one year internship or residency) full time with each academic year spread over a period of not less than two hundred working days. The period of three years duration is divided into two phases
    - Phase I consisting of First and Second academic year.
    - Phase II consisting of Internship or residency training during third year involving posting in speciality units. It is a phase of training wherein a student is exposed to actual pharmacy practice or clinical pharmacy services, and acquires skill under supervision so that he or she may become capable of functioning independently.
- 4. Minimum qualification for admission to. –
- a) Pharm.D. Part-I Course A pass in any of the following examinations -
- (1) 10+2 examination with Physics and Chemistry as compulsory subjects along with one of the following subjects:

Mathematics or Biology.

- (2) A pass in D.Pharm course from an institution approved by the Pharmacy Council of India under section 12 of the Pharmacy Act.
- (3) Any other qualification approved by the Pharmacy Council of India as equivalent to any of the above examinations.

Provided that a student should complete the age of 17 years on or before 31<sup>st</sup> December of the year of admission to the course.

Provided that there shall be reservation of seats for the students belonging to the Scheduled Castes, Scheduled Tribes and other Backward Classes in accordance with the instructions issued by the Central Government/State Government/Union Territory Administration as the case may be from time to time.

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Principal. M, Pharm Ph. A GIET SCHOOL OF PHARMACY, NH. 16, Chaitanya Knowledge City PAJAHMUNDRY-533 296: (AP b) Pharm.D. (Post Baccalaureate) Course -

A pass in B.Pharm from an institution approved by the Pharmacy Council of India under section 12 of the Pharmacy Act:

Provided that there shall be reservation of seats for the students belonging to the Scheduled Castes, Scheduled Tribes and other Backward Classes in accordance with the instructions issued by the Central Government/State Government/Union Territory Administration as the case may be from time to time.

- 5. Number of admissions in the above said programmes shall be as prescribed by the Pharmacy Council of India from time to time and presently be restricted as below
  - i) Pharm.D. Programme -30 students.
  - ii) Pharm.D. (Post Baccalaureate) Programme 10 students.
- 6. Institutions running B.Pharm programme approved under section 12 of the Pharmacy Act, will only be permitted to run Pharm.D. programme. Pharm.D. (Post Baccalaureate) programme will be permitted only in those institutions which are permitted to run Pharm.D. programme.
- 7. Course of study. The course of study for Pharm.D. shall include the subjects as given in the Tables below. The number of hours in a week, devoted to each subject for its teaching in theory, practical and tutorial shall not be less than that noted against it in columns (3), (4) and (5) below.

#### TABLES

#### First Year:

S.No.	Name of Subject	No. of hours of Theory	No. of hours of Practical	No. of hours of Tutorial
(1)	(2)	(3)	(4)	(5)
1.1	Human Anatomy and Physiology	3	3	1
1.2	Pharmaceutics	2	3	1
1.3	Medicinal Biochemistry	3	3	1
1.4	Pharmaceutical Organic Chemistry	3	3	1
1.5	Pharmaceutical Inorganic Chemistry	2	3.	1
1.6	Remedial Mathematics/ Biology	3	3*	1
	Total hours	16	18	6 = (40)

\* For Biology



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#### Sixth Year:

Internship or residency training including postings in speciality units. Student should independently provide the clinical pharmacy services to the allotted wards.

- (i) Six months in General Medicine department, and
- (ii) Two months each in three other speciality departments
- 8. Syllabus. The syllabus for each subject of study in the said Tables shall be as specified in Appendix -A to these regulations.
- 9. Approval of the authority conducting the course of study. (1) No person, institution, society or university shall start and conduct Pharm.D or Pharm.D. (Post Baccalaureate) programme without the prior approval of the Pharmacy Council of India.
  - (2) Any person or pharmacy college for the purpose of obtaining p ermission under sub-section (1) of section 12 of the Pharmacy Act, shall submit a scheme as prescribed by the Pharmacy Council of India.
  - (3) The scheme referred to in sub-regulation (2) above, shall be in such form and contain such particulars and be preferred in such manner and be accompanied withsuch fee as may be prescribed:

Provided that the Pharmacy Council of India shall not approve any institution under these regulations unless it provides adequate arrangements for teaching in regard to building, accommodation, labs., equipments, teaching staff, non-teaching staff, etc., as specified in Appendix-B to these regulations.

- 10. Examination. -(1) Every year there shall be an examination to examine the students.
  - (2) Each examination may be held twice every year. The first examination in a year shall be the annual examination and the second examination shall be supplementary examination.
  - (3) The examinations shall be of written and practical (including oral nature) carrying maximum marks for each part of a subject as indicated in Tables below:

#### TABLES

#### First Year examination:

S.No.	Name of Subject	Maximu	ım marks for	Theory	Maximun	n marks for P	racticals
		Examination	Sessional	Total	Examination	Sessional	Total
1.1	Human Anatomy and Physiology	70	30	100	70	30	100
1.2	Pharmaceutics	70	30	100	70	30	100
1.3	Medicinal Biochemistry	70	30	100	70	30	100
1.4	Pharmaceutical Organic Chemistry	70	30	100	70	30	100
1.5	Pharmaceutical Inorganic Chemistry	70	30	100	70	30	100
1.6	Remedial Mathematics/ Biology	70	30	100	70*	30*	100*
				600		2/	600 = 1200

\* for Biology.

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#### 1.6 REMEDIAL MATHEMATICS/BIOLOGY (THEORY)

Theory: 3 Hrs./Week

#### **REMEDIAL MATHEMATICS:**

1. Scope and objectives: This is an introductory course in mathematics. This subjects deals with the introduction to matrices, determinants, trigonometry, analytical geometry, differential calculus, integral calculus, differential equations, laplace transform.

#### 2. Upon completion of the course the student shall be able to: -

- a. Know Trignometry, Analytical geometry, Matrices, Determinant, Integration, Differential equation, Laplace transform and their applications;
- b. solve the problems of different types by applying theory; and
- c. appreciate the important applications of mathematics in pharmacy.

#### 3. Course materials:

#### **Text books**

- a. Differential calculus By Shantinarayan
- b. Text book of Mathematics for second year pre-university by Prof.B.M.Sreenivas

#### Reference books

- a. Integral calculus By Shanthinarayan
- b. Engineering mathematics By B.S.Grewal
- c. Trigonometry Part-I By S.L.Loney

#### 4. Lecture wise programme :

#### **Topics**

- 1 Algebra: Determinants, Matrices
- 2 Trigonometry: Sides and angles of a triangle, solution of triangles
- 3 Analytical Geometry: Points, Straight line, circle, parabola
- 4 **Differential calculus:** Limit of a function, Differential calculus, Differentiation of a sum, Product, Quotient Composite, Parametric, exponential, trigonometric and Logarithmic function. Successive differentiation, Leibnitz's theorem, Partial differentiation, Euler's theorem on homogeneous functions of two variables
- 5 **Integral Calculus:** Definite integrals, integration by substitution and by parts, Properties of definite integrals.
- 6 **Differential equations:** Definition, order, degree, variable separable, homogeneous, Linear, heterogeneous, linear, differential equation with constant coefficient, simultaneous linear equation of second order.
- 7 Laplace transform: Definition, Laplace transform of elementary functions, Properties of linearity and shifting.

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#### **BIOLOGY:**

1. Scope and objectives: This is an introductory course in Biology, which gives detailed study of natural sources such as plant and animal origin. This subject has been introduces to the pharmacy course in order to make the student aware of various naturally occurring drugs and its history, sources, classification, distribution and the characters of the plants and animals. This subject gives basic foundation to Pharmacognosy.

#### 2. Course materials:

#### Text books

- a. Text book of Biology by S.B. Gokhale
- b. A Text book of Biology by Dr. Thulajappa and Dr. Seetaram.

#### Reference books

- a. A Text book of Biology by B.V.Sreenivasa Naidu
- b. A Text book of Biology by Naidu and Murthy
- c. Botany for Degree students By A.C.Dutta.
- d. Outlines of Zoology by M. Ekambaranatha ayyer and T.N. Ananthakrishnan.
- e. A manual for pharmaceutical biology practical by S.B.Gokhale and C.K.Kokate.

#### 3. Lecture wise programme:

#### Topic

#### PART-A

- 01 Introduction
- 02 General organization of plants and its inclusions
- 03 Plant tissues
- 04 Plant kingdom and its classification
- 05 Morphology of plants
- 06 Root, Stem, Leaf and Its modifications
- 07 Inflorescence and Pollination of flowers
- 08 Morphology of fruits and seeds
- 09 Plant physiology
- 10 Taxonomy of Leguminosae, umbelliferae, Solanaceae, Lilliaceae, Zinziberaceae, Rubiaceae
- 11 Study of Fungi, Yeast, Penicillin and Bacteria

#### **PART-B**

- 01 Study of Animal cell
- 02 Study animal tissues
- 03 Detailed study of frog
- 04 Study of Pisces, Raptiles, Aves
- 05 Genearal organization of mammals
- 06 Study of poisonous animals

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#### 1.6 BIOLOGY (PRACTICAL)

#### Practical: 3 Hrs./Week

#### Title:

- 1. Introduction of biology experiments
- 2. Study of cell wall constituents and cell inclusions
- 3. Study of Stem modifications
- 4. Study of Root modifications
- 5. Study of Leaf modifications
- 6. Identification of Fruits and seeds
- 7. Preparation of Permanent slides
- 8. T.S. of Senna, Cassia, Ephedra, Podophyllum.
- 9. Simple plant physiological experiments
- 10. Identification of animals
- 11. Detailed study of Frog
- 12. Computer based tutorials

#### **Scheme of Practical Examination:**

	Sessionals	Annual
Identification	04	10
Synopsis	04	10
Major Experiment	07	20
Minor Experiment	03	15
Viva	02	15
Max Marks	20	70
Duration	03hrs	04hrs

Note: Total sessional marks is 30 (20 for practical sessional plus 10 marks for regularity, promptness, viva-voce and record maintenance.

Or. M.D. DHANA RAJU,
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#### ANDHRA UNIVERSITY



#### **MASTER OF PHARMACY**

(2020)

Regulations and Syllabus
Four semester pattern
With effect from 2020-21



Principal. M.Pharm. Ph. GIET SCHOOL OF PHARMACY, NH-16, Chaitanya Knowledge City PAJAHMUNDRY-533 296: 11

#### M.PHARM (2020) REGULATIONS AND SYLLABUS

#### **INDEX:**

- 1. Admission, instruction and attendance
- 2. Examinations Sessional and Semester end
- 3. Eligibility criteria for appointment as examiner for M.Pharm examination
- 4. Regulations for pursuing M.Pharm III and IV Semester project
- 5. Declaration of results and classification:
- 6. Grading system:
- 7. Guidelines for paper setting and model papers.

#### 1. Admission, instruction and attendance

The degree of Master of Pharmacy of the Andhra University will be conferred on a candidate who has satisfied the following conditions:

- 1.1. The candidate must have passed the B.Pharm. Degree examination of this University or B.Pharm. Degree examinations of any other University recognized by the Academic Council as equivalent thereto in First or Second class; and must have qualified in any entrance examination, if prescribed.
- 1.2. Every student, selected for admission to PG Pharmacy program in any PCI approved institution should have obtained registration with the State Pharmacy Council or should obtain the same within one month from the date of his/her admission, failing which the admission of the candidate shall be cancelled.
- 1.3. The candidate should have undergone a regular course of study as prescribed hereunder extending over a period of four semesters, ordinarily consecutive, and satisfied the academic requirements as prescribed hereinafter. The course of instruction and periods of study shall be as given in the scheme of instruction and in the syllabus.
- 1.4. The subjects of specializations for Master of Pharmacy Course shall be as follows:
  - 1. Pharmaceutical Analysis
  - 2. Pharmaceutical Chemistry
  - 3. Pharmaceutics
  - 4. Pharmaceutical Biotechnology
  - 5. Pharmacology
  - 6. Pharmacognosy
  - 7. Pharmaceutical Regulatory Affairs
  - 8. Pharmaceutical Quality Assurance
  - 9. Industrial Pharmacy
  - 10. Pharmacy Practice
- 1.5. Instruction and examination in each academic year is spread over two semesters with a minimum of 96 working days in each semester (192 in any given academic year). The odd semesters shall be conducted from the month of July to November and the even semesters shall be conducted from the month of December to April.
- 1.6. Each period of instruction is of 45 minutes duration. Eight periods of instruction are provided on each day and there are six working days in a week (Monday to Saturday).
- 1.7. Attendance Requirements: A regular course of study during an academic semester means a minimum of average attendance of 80% of all the courses of the semester computed by totaling the number of periods of lectures and practicals, as the case may be held in every course. In special cases where sufficient causes were shown, the Vice-

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Table 1: Pharmaceutical Analysis (MPA)

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			110	Interna	Internal Assessment	ıt	S of the second	
Code	Course	Credits	week	Continuous mode	Sessional Exam	Total	End Exam	Total
I Semester								
MPA 101T	Modern Pharmaceutical Analytical Techniques	4	4	10	20	30	70	100
MPA 102T	Advanced Pharmaceutical Analysis	4	4	10	20	30	70	100
MPA 103T	Pharmaceutical Validation (Common paper for MPA and MQA)	4	4	10	20	30	70	100
MPA 104T	Food Analysis	4	4	10	20	30	70	100
MPA 105P	Pharmaceutical Analysis Practical - I	2	9	15	15	30	70	100
MPA 106P	Pharmaceutical Analysis Practical - II	2	9	15	15	30	70	100
MPA 107	Seminar*	2	4	50	1	1		20
	Total	22	32	-	-			029
II Semester								
MPA 201T	Advanced Instrumental Analysis	4	4	10	20	30	70	100
MPA 202T	Modern Bio-Analytical Techniques	4	4	10	20	30	70	100
MPA 203T	Quality Control and Quality Assurance (Common paper for MPA and MQA)	4	4	10	20	30	70	100
MPA 204T	Herbal and Cosmetic Analysis	4	4	10	20	30	70	100
MPA 205P	Pharmaceutical Analysis Practical - III	2	9	15	15	30	70	100
MPA 206	Comprehensive Viva	2				-	-	50
MPA 207	Seminar*	2	2	50	1	1	1	50
	Total	22	265/	840		1	1	009
	1		13	H				

Or. M.D. DHANA KAJU,
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Table 3: Pharmaceutics (MPH)

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			/ Vonesa / II	Interna	Internal Assessment	ıt	Commonton	
Code	Course	Credits	week	Continuous mode	Sessional Exam	Total	Semester End Exam	Total
I Semester							*	
MPH 101T	Modern Pharmaceutical Analytical Techniques	4	4	10	20	30	70	100
MPH 102T	Advanced Biopharmaceutics & Pharmacokinetics (Common paper for MPH and MIP)	4	4	10	20	30	70	100
MPH 103T	Modern Pharmaceutics	4	4	10	20	30	70	100
MPH 104T	Regulatory Affairs	4	4	10	20	30	70	100
MPH 105P	Pharmaceutics Practical – I	2	9	15	15	30	70 7	100
MPH 106P	Pharmaceutics Practical – II	2	9	15	. 15	30	70	100
MPH 107	Seminar*	2	4	50	-	ł	1	50
	Total	22	32	1	1	1	1	029
II Semester								
MPH 201T	Molecular Pharmaceutics (Nano Technology and Targeted DDS)	4	4	10	20	30	70	100
MPH 202T	Drug Delivery Systems (DDS)	4	4	10	20	30	70	100
MPH 203T	Computer Aided Drug Development (CADD)	4	4	10	20	30	70	100
MPH 204T	Pharmaceutical and Cosmetic Product Development	4	4	10	20	30	70	100
MPH 205P	Pharmaceutics Practical - III	2	9	15	15	30	70	100
MPH 206	Comprehensive Viva	2	-	-	1	1	1	50
MPH 207	Seminar*	2	052	50				50
	Total	155	26.5	1		1		009
	or. M.D. DHANA RAJI	H-WE	AND HAUNDRY BY					

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Table 5: Pharmacology (MPL)

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,			/SamoH	Interna	Internal Assessment	nt	o d	
Code	Course	Credits	week	Continuous mode	Sessional Exam	Total	Semester End Exam	Total
I Semester								
MPL 101T	Modern Pharmaceutical Analytical Techniques	4	4	10	20	30	70	100
MPL 102T	Advanced Pharmacology – I	4	4	10	20	30	70	100
MPL 103T	Pharmacokinetics and Drug Metabolism	4	4	10	20	30	70	100
MPL 104T	Cellular and Molecular Pharmacology	4	4	10	20	30	70	100
MPL 105P	Pharmacology Practical - I	2	9	15	15	30	70	100
MPL 106P	Pharmacology Practical – II	2	9	15	15	30	70	100
MPL 107	Seminar*	2	4	50	1	1	1	50
	Total	22	32	I	ŀ	1	1	029
II Semester								
MPL 201T	Advanced Pharmacology - II	4	4	10	20	30	70	100
MPL 202T	Pharmacological and Toxicological Screening Methods	4	4	10	20	30	70	100
MPL 203T	Principles of Drug Discovery	4	4	10	20	30	70	1000
MPL 204T	Clinical Research and Pharmacovigilance	4	4	10	20	30	70	100
MPL 205P	Pharmacology Practical – III	2	9	15	15	30	70	100
MPL 206	Comprehensive Viva	2	1	1	1	1	1	50
MPL 207	Seminar*	2	2	50	1	1	1	50
	Total	22	26	-	1	1	1	009

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Table 8: Pharmaceutical Quality Assurance (MQA)

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			/ 500 800 1	Interna	Internal Assessment	ıt	C comp of the comp	
Code	Course	Credits	mours/ week	Continuous	Sessional Exam	Total	End Exam	Total
I Semester								
MQA 101T	Modern Pharmaceutical Analytical Techniques	4	4	10	20	30	70	100
MQA 102T	Quality Management System	4	4	10	20	30	70	100
MQA 103T	Pharmaceutical Validation (Common paper for MPA and MQA)	4	4	10	20	30	70	100
MQA 104T	Product Development and Technology Transfer	4	4	10	20	30	70	100
MQA 105P	Pharmaceutical Quality Assurance Practical - I	2	9	15	15	30	70	100
MQA 106P	Pharmaceutical Quality Assurance Practical - II	2	9	15	15	30	70	100
MQA 107	Seminar*	2	4	50	1	1	1	50
	Total	22	32	1	1	ŀ	l	029
II Semester								
MQA 201T	Hazards and Safety Management	4	4	10	20	30	70	100
MQA 202T	Audits and Regulatory Compliance	4	4	10	20	30	70	100
MQA 203T	Quality Control and Quality Assurance (Common paper for MPA and MQA)	4	4	10	20	30	70	100
MQA 204T	Pharmaceutical Manufacturing Technology	4	4	10	20	30	70	100
MQA 205P	Pharmaceutical Quality Assurance Practical - III	2	9	15	15	30	70	100
MQA 206	Comprehensive Viva	2				-	-	50
MQA 207	Seminar*	2	2	50				50
	Total	22	26	!	-	1	-	009
	132 10							

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or. M.D. DHANA RAJU,

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# II/II III Semester

3312	3312 Seminar on the objectives and work plan of the proposed project to be	
	completed within one month from the commencement of the project	(w)
3313	3313 Mid-term project review at the end of third semester 50	
3314	Seminar on Selected research topic 100	
	Total 200	

# II/II IV Semester

3415	3415 Thesis Evaluation	100
3416	Thesis Viva-voce	100
	Total	200
	Grand total	1500



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# I. PHARMACEUTICAL TECHNOLOGY

## I/II I Semester

Course		No. of hours/week	s/week	Max. Marks	Questions to be answered
No.		Theory	Practical		in the semester end examination
3101	Biostatistics Theory (Common for all	5		100	5 out of 7
	specializations)				
3102	Biopharmaceutics and Pharmacokinetics	2	1	100	5 out of 7
	Theory				
3103	Biopharmaceutics and Pharmacokinetics	1	10	100	
	Practical				
3104	Advanced Physical Pharmaceutics Theory	2	1	100	5 out of 7
3105	Advanced Physical Pharmaceutics Practical	1	10	100	
3106	Comprehensive viva			50	
	Total			550	

## I/II II Semester

3207	Modern Analytical Techniques Theory Common for all	2		100	5 out of 7
	specializations				
3208	Quality Assurance and Drug Regulatory Affairs Theory Common for all specializations	rv.	1	100	5 out of 7
3209	Novel Drug Delivery Systems Theory	22	1	100	5 out of 7
3210	Product Formulation and Development Theory	2	1	100	5 out of 7
3211	Product Formulation and Development Practical		10	100	
3212	3212 Comprehensive Viva RAIII	AJO.		50	
	(I) Sold (A) Principal M. Pharm. Ph.	M, Pharm., Ph.			

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