

M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

$\boldsymbol{SEMESTER-I}$

S.	Course	Course Name	Hours	per v	week	Credits
No.	codes		L	T	P	
1.	21S01101	Modern Pharmaceutical Analytical Techniques	4	-	-	4
2.	21S03101	Advanced Physical Pharmaceutics	4	-	-	4
3.	21S08101	Pharmaceutical formulation Development	4	-	-	4
4.	21S03103	Advanced Biopharmaceutics & Pharmacokinetics	4	-	-	4
5.	21S01105	Modern Pharmaceutical Analytical Techniques Lab	-	-	6	3
6.	21S08102	Advanced Physical Pharmaceutics Lab	-	-	6	3
7.	21DAC101b	Audit Course – I English for Research paper writing Disaster Management Sanskrit for Technical Knowledge	2	1	-	0
8.	21S10101	Seminar/Assignment	-	1	6	4
		Total	18	1	18	26

SEMESTER - II

S.No.	Course	Course Name	He	ours p	oer	Credits
	codes		L	T	P	
1.	21S08201	Pharmaceutical Production Technology	4	-	-	4
2.	21S08202	Advanced Drug Delivery systems	4	-	-	4
3.	21S08203	Pharmaceutical Industrial Management	4	-	-	4
4.	21S03204	Nano Drug Delivery systems	4	-	-	4
5.	21S08204	Pharmaceutical Production Technology Lab	-	-	6	3
6.	21S03206	Advanced Drug Delivery systems Lab	-	-	6	3
7.	21DAC201b	Audit Course – II Pedagogy Studies Stress Management for Yoga Personality Development through Life Enlightenment Skills	2	-	1	0
8.	21S10201	Seminar/Assignment	-	1	6	4
		Total	18	1	18	26



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SEMSTER - III

S.No.	Course	Course Name	Hours per week		Credits	
	codes		L	T	P	
1.	21DRM101	Research Methodology and Intellectual Property Right	4	1	-	4
2.	21SOE301d 21SOE301a	Open Electives Biological Screening methods Pharmaceutical Validation Entrepreneurship Management	3	1	ı	3
3.	21S10301	Teaching Practice/Assignment	1	-	4	2
4.	21S10302	Comprehensive viva voce	-	-	-	2
5.	21S10303	Research Work - I	-		24	12
		Total	7	_	32	23

SEMESTER - IV

S.No.	Course	Course Name	Hours per week		Credits	
	codes		\mathbf{L}	T	P	
1.	21S10401	Co-Curricular Activities	2			2
2.	21S10402	Research Work - II	3		30	18
		Total	5		30	20



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Course Code	MODEDN DILADMA CEUTICAT	ANIAT STOLETAT	Т	T	n	
Course Code 21S01101	MODERN PHARMACEUTICAL TECHNIQUES	ANALYTICAL	L 4	$\frac{T}{0}$	P 0	<u>C</u>
Pre-requisite	TECHNIQUES	Semester	4		<u> </u>	4
Pre-requisite		Semester				
Course Objectives:						
· ·	with various advanced analytical instrum	nantal taahniguas t	Con i	donti	ficati	ion
	quantification of drugs. Instruments deal					
HPLC, GC etc.	quantification of drugs. Instruments dear	t are minik, mass	spec	HOIII	cici,	щ,
	CO): Student will be able to					
	letion of course student is able to know	about chemicals a	nd e	xcini	ents	
_	s of various drugs in single and combination			т-г	• • • • • • • • • • • • • • • • • • • •	•
_	and practical skills of the instruments	ation dosage forms	,			
UNIT - I	and practical skins of the instruments					
		· · · · · ·	<u> </u>			
_	copy: Introduction, Theory, Laws, Instrum					
	e of solvents and solvent effect and Appli	cations of UV-Visi	ble s	specti	rosco	ру,
Difference/ Derivativ	ve spectroscopy.		1			
UNIT - II		1 1 11 Y	1		. •	
	heory, Modes of Molecular vibrations, S					
	rier -Transform IR Spectrometer, Factors	affecting vibrationa	I fre	quen	cies	and
	pectroscopy, Data Interpretation.					
UNIT - III	0 1 1 1 1 1 1 ND (D	D : 1 T :	<u> </u>	.•	G 1	
	Quantum numbers and their role in NMR					
	R, Relaxation process, NMR signals in					
	chemical shift, Spin-Spin coupling, Coupling					
	utline of principles of FT-NMR and	"C NMR. Applic	catioi	is o	I INI	MK
spectroscopy.						
UNIT – IV	Disciple Theory Instrumentation of M	C T	>: cc -	4	4	C
	Principle, Theory, Instrumentation of Ma					
	ron impact, chemical, field, FAB and MA					
	ne of Flight, Mass fragmentation and its ru	nes, meta stable 101	18, 18	orob	ic pe	aks
and Applications of I	wiass specificacopy.					
Chromotography			1			

Chromatography

Introduction to chromatography and classification of chromatographic methods based on the mechanism of separation, Principle, instrumentation, selection of solvents; chromatographic parameters, factors affecting resolution, applications of the following:

a) Thin Layer chromatography; b) High Performance Thin Layer Chromatography

c) Paper Chromatography; d) Column chromatography

e) Gas chromatography; f) High Performance Liquid chromatography

g) Affinity chromatography; h) Gel Chromatography

i)Hyphenated techniques:

- Ultra High Performance Liquid chromatography- Mass spectroscopy
- Gas Chromatography-Mass Spectroscopy

Reference Books:

- 1. Instrumental Methods of Chemical Analysis by B.K Sharma
- 2. Vogel's Text book of Quantitative Chemical Analysis by A.I. Vogel
- 3. Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John



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- Wiley & Sons, 2004.
- 4. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5th edition, Eastern press, Bangalore, 1998.
- 5. Instrumental methods of analysis Willards, 7th edition, CBS publishers.
- 6. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4thedition, CBS Publishers, New Delhi, 1997.
- 7. Organic Spectroscopy William Kemp, 3rd edition, ELBS, 1991.
- 8. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi,3rd Edition, CBS Publishers, New Delhi, 1997.
- 9. Pharmaceutical Analysis Modern Methods Part B J W Munson, Vol11, Marcel. Dekker Series
- 10. Spectroscopy of Organic Compounds, 2nd edn., P.S/Kalsi, Wiley esternLtd., Delhi.
- 11. Textbook of Pharmaceutical Analysis, KA.Connors, 3rd Edition, John Wiley& Sons, 1982.
- 12. Organic Chemistry by I. L. Finar
- 13. Quantitative Analysis of Drugs by D. C. Garrett
- 14. HPTLC by P.D. Seth
- 15. Indian Pharmacopoeia 2007
- 16. High Performance thin layer chromatography for the analysis of medicinal plants by Eike
- 17. Reich, Anne Schibli
- 18. Introduction to instrumental analysis by Robert. D. Braun



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COURSE STRUCTURE & SYLLABI

Course Code	ADVANCED DIVERGAT DITADMA CETTEROS	L	T	P	C
21S03101	ADVANCED PHYSICAL PHARMACEUTICS	4	0	0	4
	Semester			I	
Course Objectives:					
The students shall kn	now about particle science, polymer science and its use in pharm	nace	utica	l dos	age
	now the compression and consolidation parameters for powder				
Students also know	about the rheology, disperse systems, dissolution and solubilit	y pa	aram	eters	for
dosage forms.					
	CO): Student will be able to				
The students will kn	ow particle size analysis method, solid dispersion, physics of	tab	lets,	polyı	mer
classification and it	s applications, student will also know the stability calcula	ation	ıs, sl	nelf	life
calculations and acc	elerated stability studies. They also know the rheology, abso	orpti	on re	lated	l to
liquids and semi-so	lid dosage forms. They also know the factors affecting the	dis	solut	ion	and
	o invitro/invivo correlations.				
UNIT - I					
Polymer science: (Classification, properties and characterization of polymers, p	ohas	e set	arati	on,
	ate, preparation of polymer solution, application of polymers i				
formulations. Mecha	unism of biodegradation of biodegradable polymers including	co	ntroll	ed d	rug
	acoadhesive, Hydrodynamically balanced and Transdermal Syst				Ü
UNIT - II					
Physics of tablet co	ompression: Basic principles of interactions, compression ar	nd c	onso	lidati	on,
	onsolidation under high loads, effect of friction, distributi				
	volume relationships, Heckel plots, compaction profiles, end				
compaction, Measure	ement of compression with strain gauges, compression pressure	-QA	para	mete	rs.
UNIT - III	* * * * * * * * * * * * * * * * * * * *				
Kinetics and drug	stability: Stability calculations, rate equations, complex order	kin	etics,	Fact	tors
	, strategy of stability testing, method of stabilization, method				
stability testing in o	losage forms, temperature and humidity control, physical sta	abili	ty te	sting	of
pharmaceutical produ	acts. Photodecomposition Method, solid state decomposition.				
UNIT - IV					
Theoretical considera	ation, instrumentation, rheological properties of disperse system	s an	d ser	nisoli	ids.
Oscillatory testing, C	Creep measurement.				
Characterization of	API and excipients: Differential Scanning Calorimetry: I	rinc	iple,	ther	mal
transitions, advantage	es, disadvantages, instrumentation, applications and interpretation	ons			
X Ray Diffraction	on methods: Origin of x-rays, principle, advantages,	d	isadv	antag	ges,
	lications and interpretations.				
UNIT - V					
	ability: Solubility and solubilization of nonelectrolytes, solubili		•		
	olvents, complexation, drug derivatization and solid sta-				
M 1 ' CD	g release - dissolution, diffusion (Matrix and Reservoir) and sv	zelli:	na ca	ntrol	led

Textbooks:

1. Physical Pharmacy, 4th Edition by Alfred Martin.

(Peppas Model) and dissolution equipment

- 2. Theory and Practice of Tablets Lachman, Vol.4
 3. Pharmaceutical Dosage forms Disperse systems Vol. I & II
- 4. Cartenson "Drug Stability, Marcel Decker Solid state properties, Marcel Dekker.



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5. Industrial Pharmacy - Selected Topics, CVS Subramanyam and J Thimmasetty, Vallabh Prakashan Delhi – 2013

Reference Books:

- 1. Dispersive systems I, II, and III
- 2. Robinson. Controlled Drug Delivery Systems



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Course Code	PHARMACEUTICAL FORMULATION		T	P	C
21S08101	DEVELOPMENT		0	0	4
Pre-requisite	Semes	ter	Í		

Course Objectives:

This subject is to make the student achieve different parameters and factors that influence the dosage form design. This subject also impart the knowledge about unit operations, solid dosage forms and powders.

Course Outcomes (CO): Student will be able to

- Different machinery used for various steps in manufacture of various dosage forms.
- Formulation and evaluation of hard and soft gelatin capsules and their advantages over other dosage forms.

UNIT - I

- a. **Preformulation studies:** Goals of preformulation, preformulation parameters, methodology, solid state manipulation and characterization, solubility and partition coefficient, drug excipients compatibility, intrinsic dissolution.
- b. Advances in Pharmaceutical excipients. Excipients selection for capsules, tablets, suspensions and emulsions.
- c. Packaging development selection of primary and secondary packaging materials and testing

UNIT - II

Pharmaceutical unit operations: A detail study involving machinery and theory of pharmaceutical unit operations like solid orals: Wet granulation- Rapid mixer granulator and Top spray granulation, Dry granulation- Slugging and roller compaction, drying, milling, blending, filtration and sterilization.

UNIT - III

Formulation development of solid and powder dosage forms: Improved production techniques for tablets, new materials, processes, equipment improvements, high shear mixers, compression machines, coating machines, coating techniques in tablet technology for product development and computerization for in process quality control of tablets. Formulation development and manufacture of powder dosage form for internal and external use including inhalation dosage forms.

UNIT - IV

Formulation development of soft and hard gelatin capsules: Introduction, production and methods of manufacture, filling equipment and filling operations, formulations, finishing, special techniques, the nature of the capsule shell and capsule, advances in capsule manufacture, processing and control including pharmaceutical aspects, physical stability and packaging.

UNIT - V

Optimization techniques in pharmaceutical formulation and processing: Quality by Design: Concept and application to formulation development. Design of experiments (DOE): Formula and process optimization statistical design, response surface method, contour diagrams, factorial design, partial factorial design, simplex methods, mixture designs, Plackett Burman method, Box Benken method, applications in pharmaceutical formulation.

Textbooks:

- 1. Pharmaceutics The Science of Dosage form design by ME Aulton.
- 2. Pharmaceutical Dosage forms Tablets (Vol I, II and III) by Lieberman, Lachman and Schwartz.
- 3. Pharmaceutical Dosage forms Capsules (Vol I, II and III) by Avis, Lieberman and Lachman.
- 4. Pharmaceutical Dosage forms Disperse systems (Vol I, II and III) by Avis, Lieberman and



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Lachman.

- 5. Modern Pharmaceutics by Gilbert S. Banker and Christopher T. Rhodes.
- 6. Hand book of Pharmaceutical excipients
- 7. CVS Subhramanyam & J Thimmasethy, Industrial Pharmacy, Vallabh Prakasham, Delhi, 2014

Reference Books:

- 1. The Theory and Practice of industrial Pharmacy by Leon Lachman, Herbert A. Lieberman.
- 2. Remington's Science and Practice of Pharmacy by A. Gennaro.
- 3. Ansel's Pharmaceutical Dosage form and Drug delivery system by Loyd V. Allen, Jr. Nicholas G. Popovich, Howard C. Ansel.
- 4. Generic Drug Product Development by Leon Shargel and Isadore Kanfer.
- 5. Dispensing for Pharmaceutical Students by SJ Carter.
- 6. Pharmaceutical Packaging Technology by UK Jain, DC Goupale S Nayak.



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Course Code	ADVANCED BIOPHARMACEUTICS &	L	T	P	C
21S03103	PHARMACOKINETICS	4	0	0	4
	Semester	I			

Course Objectives:

The student shall know about bioavailability, bioequivalence and factor affecting bioavailability. They also know the pharmacokinetic parameter like drug disposition, absorption, nonlinear and time dependant pharmacokinetics. They also know about the drug interactions & problems associated in pharmacokinetic parameters calculations.

Course Outcomes (CO): Student will be able to

Students will be able to tell factors affecting the bioavailability and stability of dosage form; they also know the bioequivalence studies and protocols for bioequivalent studies. They also know the parameters for the disposition, absorption and Michaelis-Menton constants for nonlinear kinetics.

UNIT - I

- a. Biological and metabolic factors affecting bioavailability, complexation, dissolution techniques of enhancing dissolution.
- b. Formulation factors affecting bioavailability of drugs in dosage forms of tablets, capsules, Parenterals, liquid orals and topical dosage forms.
- c. **Bioavailability:** Importance, dose dependency, AUC, rate and extent, assessment, blood and urine samples, single dose and multiple dose studies, *Invitro- Invivo* Correlation analysis and Levels of Correlations.
- d. **Bioequivalence:** Importance equivalency concepts, biowaivers, study designs, protocol, transformation of data, Statistical Criteria as per the Regulations.

UNIT - II

Pharmacokinetics – **Drug Disposition:** compartment models: One, two and non-compartmental approaches to pharmacokinetics. Recent trends, merits and limitations of these approaches.

Application of these models to determine the various pharmacokinetic parameters pertaining to:

- a. Distribution: Apparent volume of distribution and its determination, factors affecting.
- b. Metabolism: Metabolic rate constant, Factors affecting Metabolism
- c. Elimination: Over all apparent elimination rate constant, and half life.

All the above under the following conditions:

- 1. Intravenous infusion
- 2. Multiple dose injections
- d. Non-invasive methods of estimating pharmacokinetics parameters with emphasis on salivary and urinary samples.
- e. Concept of clearance: organ, total clearance, hepatic clearance, lung clearance and renal clearance.

UNIT - III

Pharmacokinetics – **Absorption:** Rate constants – Zero order, first order, Models of experimental study of absorption (in silico, in vitro, in situ and in vivo) – Absorption half lives, method of residuals, Wagner – Nelson method, Loo - Reigleman method, Analysis of kinetics from urine samples. Pharmacokinetic parameters determination pertaining to: Multiple dosage oral administration.

UNIT - IV

Non-linear pharmacokinetics: Concepts of linear and non-linear pharmacokinetics, Michaelis-Menton kinetics characteristics. Basic kinetic parameters, possible causes of non-induction, nonlinear binding, and non-linearity of pharmacological responses.



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COURSE STRUCTURE & SYLLABI

Clinical Pharmacokinetics: Altered kinetics in pregnancy, child birth, infants and geriatrics. Kinetics in GI disease, malabsorption syndrome, liver, cardiac, renal and pulmonary disease states.

UNIT - V

Time dependent pharmacokinetics: Introduction, classification, physiologically induced time dependency: Chronopharmacokinetics - principles, drugs— (amino glycosides, NSAIDS, antihypertensive drug) chemically induced dependency.

Drug Interactions: Kinetics of drug interaction, study of drug-drug interaction mediated through absorption, distribution, metabolism and elimination, mechanisms of interaction and consequence. Numerical problems associated with all units, if any.

Textbooks:

- 1. Biopharmaceutics and Clinical Pharmacokinetics by Milo Gibaldi.
- 2. Learn Shargel and ABC yu, Applied Biopharmacokinetics and Pharmacokinetics
- 3. Biopharmaceutics and Pharmacokinetics by C.V.S. Subrahmanyam, Vallabh Prakashan. 2010.
- 4. Basic biopharmaceutics, Sunil S. Jambhekar and Philip J Brean.
- 5. Text book of Biopharmaceutics and Clinical Pharmacokinetics by NiaziSarfaraz

Reference Books:

- 1. Bio-Pharmaceutics and Pharmacokinetics by V. Venkateshwarlu.
- 2. Pharmacokinetics, Biopharmaceutics and Clinical pharmacy by Robert E. Notari.
- 3. Biopharmaceutics and Clinical Pharmacokinetics An Introduction by Robert E. Notari.
- 4. Drug drug interactions, scientific and regulatory perspectives by Albert P. G



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Course Code	MODERN PHARMACEUTICAL ANALYTICAL	L	T	P	C
21S01105 TECHNIQUES LAB		0	0	6	3
Semester]	[

List of Experiments

- 1. Analysis of Pharmacopoeial compounds and their formulations by UV Vis Spectrophotometer.
- 2. Simultaneous estimation of multi component containing formulations by UV Spectrophotometry
- 3. Effect of pH and solvent on UV –Spectrum
- 4. Determination of Molar absorption coefficient
- 5. Estimation of riboflavin/ quinine sulphate by fluorimetry
- 6. Study of quenching effect by fluorimetry
- 7. Estimation of sodium or potassium by flame photometry
- 8. Colorimetric determination of drugs by using different reagents
- 9. Quantitative determination of functional groups
- 10. Experiments based on Column chromatography
- 11. Experiments based on HPLC
- 12. Experiments based on Gas Chromatography



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Course Code	ADVANCED PHYSICAL PHARMACEUTICS LA	L	T	P	C
21S08102	ADVANCED PHISICAL PHARMACEUTICS LAB 0		0	6	3
Semester]	[

List of Experiments

- 1. Determinates of molecular weight of some selected polymers.
- 2. Preparation and evaluation of solid dispersions (Immediate release and sustained release)
- 3. Accelerated stability testing of Aspirin Tablets
- 4. Stability evaluation of Aspirin at various pH and temperature conditions
- 5. Determination of Ist order and 2nd order rate constant. Half life by Acid / Alkali hydrolysis
- 6. Preparation and evaluation of multiple emulsions
- 7. Preparation and evaluation of β -cyclodextrin complexes of some drugs.
- 8. Generation of dissolution profiles of few dosage forms and application of the data into various kinetic equations. Calculation of Hixon-crowell dissolution rate constant
- 9. Preparation and dissolution study of paracetmol tablets and comparison with the marketed product.
- 10. Study of solubility and dissolution for few drugs and their respective salts.
- 11. Study of drug release from commercial suspension and emulsion dosage forms
- 12. Viscosity measurement of Newtonian and Non-Newtonian liquids



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Course Code	PHARMACEUTICAL PRODUCTION		L	T	P	C
21S08201	TECHNOLOGY		4	0	0	4
Pre-requisite		Semester		II		

Course Objectives:

The students shall know about the pilot plant scale up techniques for manufacturing of tablets, capsules, suspensions, emulsions and semisolids. The students also know about the filling of capsules, compression machines, sterilizers for formulation of parenterals and also know about the propellants, DPI, MDI and their quality control. The students also know about the cosmetics and nutraceuticals.

Course Outcomes (CO): Student will be able to

Students will know about the scale up and pilot plant techniques used for all pharmaceutical dosage forms like tablets, capsules, parenterals, aerosols, cosmetics and nutraceuticals.

UNIT - I

Pilot plant scale-up techniques used in pharmaceutical manufacturing

- **a. Pilot plant:** Technology transfer from R&D to pilot plant to pilot scale considerations of steps involved with manufacture, layout design, facility, equipment selection of tablets, capsules, suspensions, emulsions & semisolids.
- **b. Scale up:** Importance, Scale up process-size reduction, mixing, blending, granulation, compression, coating involved in tablets, capsules & liquid-liquid mixing.

UNIT - II

Formulation development of parenteral dosage forms: Advances in materials and production techniques, filling machines, sterilizers, product layout.

IINIT - III

Pharmaceutical Aerosols: Advances in propellants, metered dose inhaler designs, dry powder inhalers, selection of containers and formulation aspects in aerosols formulation, manufacture and quality control.

UNIT - IV

a. Cosmetics: Formulation approaches, preparation & method of manufacturing labeling& Q.C. of anti ageing products, sun screen lotion and fairness creams.

b. Nutraceuticals:

- 1. Introduction, source, manufacture and analysis of glucosamine and cartinine.
- 2. Monographs: General and specific properties of glucosamine &cartinine.
- 3. A brief overview of role of nutraceuticals in cancer prevention & cardio vascular disorders.

UNIT - V

Aseptic processing operation

- **a.** Introduction, contamination control, microbial environmental monitoring, microbiological testing of water, microbiological air testing, characterization of aseptic process, media and incubation condition, theoretical evaluation of aseptic operations.
- **b.** Air handling systems: Study of AHUs, humidity & temperature control.

Textbooks:

- 1. Pharmaceutics The Science of Dosage form design by ME Aulton.
- 2. The Theory and Practice of industrial Pharmacy by Leon Lachman, Herbert A. Lieberman.
- 3. Remington's Science and Practice of Pharmacy by A. Gennaro.
- 4. Ansel's Pharmaceutical Dosage form and Drug delivery system by Loyd V. Allen, Jr. Nicholas G. Popovich, Howard C. Ansel.



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- 5. Pharmaceutical Dosage forms Parenterals (Vol I, II and III) by Avis, Lieberman and Lachman.
- 6. Scale up techniques Pharmaceutical process by Michael Levin, Marcel Dekker

Reference Books:

- 1. Bentley's Text Book of Pharmaceutics by EA Rawlins.
- 2. Generic Drug Product Development by Leon Shargel.
- 3. Dispensing for Pharmaceutical Students by SJ Carter.
- 4. Modern Pharmaceutics by Gilbert S. Banker and Christopher T. Rhodes.
- 5. Nutraceuticals, 2nd edition by Brian lock wood



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Course Code	Course Code 21S08202 ADVANCED DRUG DELIVERY SYSTEMS		L	T	P	C	
21S08202			4	0	0	4	
Pre-requisite		Semester	II				
Course Objectives:							
The students shall a	apply the pharmacokinetic and pharmaco	dynamic principles	in tl	he de	esign	of	
CDDS. They also	apply the design, evaluation and appli	cations related to	oral	, pa	rente	ral,	
Transdermal, implan	its, bioadhesives and targeted drug delivery	systems.					
Course Outcomes (CO): Student will be able to	_		•		•	
Students will select the drugs for CDDS design of the formulation fabrication of systems of above							

UNIT - I

Fundamentals of controlled drug delivery systems, pharmacokinetic and pharmacodynamic basis of controlled drug delivery. Design, fabrication, evaluation and applications of the following controlled releasing systems

a. Controlled release oral drug delivery systems

drug delivery systems with relevant applications.

b. Parenteral controlled release drug delivery systems

UNIT - II

Design, fabrication, evaluation and applications of the following

- a. Implantable Therapeutic systems
- b. Transdermal delivery systems
- c. Ocular and Intrauterine delivery systems
- d. Vaccine delivery: Delivery systems used to promote uptake, absorption enhancers, oral immunization, controlled release microparticles form vaccine development

UNIT - III

Biochemical and molecular biology approaches to controlled drug delivery of

- a. Bioadhesive drug delivery systems
- b. Nasal drug delivery systems
- c. Drug delivery to Colon

UNIT - IV

Biochemical and molecular biology approaches to control drug delivery of

- a. Liposomes
- b. Niosomes
- c. Microspheres
- d. Nanoparticles
- e. Resealed erythrocytes

UNIT - V

Drug targeting to particular organs

- a. Delivery to lungs
- b. Delivery to the brain and problems involved
- c. Drug targeting in neoplasms

Textbooks:

- 1. Novel Drug Delivery System by Yie W. Chien.
- 2. Controlled Drug Delivery by Joseph R. Robinson and Vincent H. L. Lee.
- 3. Controlled and Novel Drug Delivery Systems by N. K. Jain.
- 4. Targeted and Controlled Drug Delivery (Novel carrier systems) by S. P. Vyas and Khar.
- 5. Modern Pharmaceutics by Gilbert S. Banker and Christopher T. Rhodes.



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- 6. Advances in Drug Delivery, Vol 1, 2, 3 by Y. Madhusudan Rao, A.V. Jithan
- 7. Oral Drug Delivery Technology, 2nd ed, by Aukunuru Jithan



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Course Code 21S08203	PHARMACEUTICAL INDUSTRIAL	L 4	T 0	P 0	C
21500205	MANAGEMENT Semester	4	U I		4
	Schrester				
Course Objectiv	es:				
	dy of the course aimed at achieving, enabling the student effective	ly m	anag	e a	
	n in planning, hiring, personnel, selection training and other infrast	•	_		
0	t from design, lay-out and handling of the equipment				
	es (CO): Student will be able to				
This subject aims waste materials.	at validation of different process, equipment methods and effective	e ma	nage	ment	of
UNIT - I					
Human Resourc	e management: Human resource planning, job analysis and design	, rec	ruitn	nent,	
Personnel selection appraisal key resu	on, orientation and placement, training and development, supervisional tarea and key performance area remuneration and salaries, Comprial relations, motivation.	n, p	erfor	manc	e
UNIT - II	·				
design, layout, co control contamina installation and m	ganization and personnel, responsibilities, training, hygiene Premise instruction, maintenance, and sanitations, environmental control, station, Equipments procedure and documentation for selection, purchaintenance, clean in place, sterilization in place.	erile	areas	8,	on,
UNIT - III					
practices, layout of handling and tran selection of vendo	agement: Production organization, objectives and policies of good of buildings, services, equipments and their maintenance, materials sportation, inventory management and control, production planning ors, purchase cycle, sales forecasting, budget and cost control.	man	agen	nent,	
UNIT - IV					
	on: General Principles of Validation, Regulatory basis, validation of ocesses, validation of analytical methods.	f pha	ırma	ceuti	cal
UNIT - V					
Industrial Hazar hazards, safety m Pollution, air Poll Noise Abatement Plants, Effluent T	rds and Pollution Management: Chemical hazards, gas hazards, franagement. Water pollution, water Pollution abatement and effluent ution Control Devices. Solid waste, Solid Waste Management, Noi, Effluent Analysis and Treatment-Methods, Effluent Treatment in reatment in Synthetic Drugs Industry, Effluent Treatment in Ferme cho Parmacovigilance.	t trea se P Forn	atme olluti nulat	nt, A ion, ion	ir
Textbooks:					
1. Unit operations	of Chemical Engineering by Warren L. McCabe, Julian C. Smith, d Practice of industrial Pharmacy by Leon Lachman, Herbert A. Lie			rriott	•

Pharmaceutical Process validation by Robert A. Nash, Alfred H. Wachter.
 Modern Pharmaceutics by Gilbert S. Banker and Christopher T. Rhodes.

Online Learning Resources:

5. Pharmaceutical production management, C.V.S. Subrahmanyam, Vallabh Prakash.



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- 1. Remington's Science and Practice of Pharmacy by A. Gennaro.
- 2. Bentley's Text book of Pharmaceutics by EA Rawlins.



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

Course Code	NANO DRUG DELIVERY SYSTEMS	L	T	P	C
21S03204	NANO DRUG DELIVERT STSTEMS	4	0	0	4
	Semester		I	I	
Course Objective					
	ertise regarding suitability and evaluation of nanomaterials, at				
	fabrication of nanopharmaceutical, evaluate the intensity of de	osag	e for	ms	and
	rgeting and controlled delivery.				
	es (CO): Student will be able to				
	ald be able to select the right kind of materials, able to develop n			ulati	ons
with appropriate t	echnologies, evaluate the product related test and for identified disc	eases	S		
UNIT - I					
Introduction to I	Nanotechnology				
a. Definition of na					
b. History of nano	otechnology				
	ies and classification of nanomaterials				
d. Role of size an	d size distribution of nanoparticles properties.				
	ulations based on nanotechnology and science behind them				
UNIT - II					
Synthesis of Nan	omaterials				
Physical, chemica	al and biological Methods				
Methods for syntl	nesis of				
 Gold nan 	oparticles				
 Magnetic 	nanoparticles				
 Polymeric 	c nanoparticles				
• Self – a	assembly structures such as liposomes, Niosomes, transferas	some	es, n	nicel	les,
	es and nanoemulsions				
UNIT - III					
Biomedical appli	ications of Nanotechnology				
	y products used for in vitro diagnostics				
b. Improvements	to medical or molecular imaging using nanotechnology				
c. Targeted nanor	naterials for diagnostic and therapeutic purpose				
UNIT - IV					
Design of nanom	naterials for drug delivery, pulmonary and nasal drug delivery, n	ano	mate	rials	for
	d cardiovascular diseases. Localized drug delivery systems.				
UNIT - V					
	including the principles size reduction, analysis of panoparticles	015	70 D	DI (170

Characterization including the principles, size reduction, analysis of nanoparticles, size, PDI, size separation, stability, methods of analysis regarding integrity and release of drugs

Reference Books:

- 1. Nanomedicine and Nanoproducts: Applications, Disposition and Toxicology in the Humanbody, Eiki Igarashi, CRC press. 2015
- 2. Nanotechnology and Drug Delivery Volume one and two: Nanoplatforms in Drug Delivery, Jose L. Arias, CRC press
- 3. Nano: The Essentials: Understanding Nanoscience and Nanotechnology, T. Pradeep, Tata McGraw-Hill Publishing Company Limited, New Delhi, 2008.
- 4. Nanocrystals: Synthesis, Properties and Applications, C. N. R. Rao, P. J. Thomas and



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- G.U.Kulkarni, Springer (2007)
- 5. Nanostructures and Nanomaterials: Synthesis, Properties and Application, Guozhong Gao, Imperial College Press (2004)
- 6. Nano chemistry: A Classical Approach to Nanomaterials Royal Society for Chemistry, Cambridge, UK (2005)
- 7. Nanocomposite science and technology, pulickel M. Ajayan, Linda S. Schadler, paul V.Braun, Wiley VCH Verlag, Weiheim (2003)
- 8. Nanoscale materials in chemistry, Edited by Kenneth J. Klabunde, John Wiley & Sons, 2009
- 9. Nanoparticles as Drug carriers, Vladimir P Torchiling, Imperial College Press, USA, 2006
- 10.Introduction to Nano Science and Technologies, Ankaneyulu Yerramilli, BS Publications. 2016



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

Course Code	PHARMACEUTICAL PRODUCTION TECHNOLOGY	L	T	P	C
21S08204	LAB	0	0	6	3
	Semester				

List of Experiments

- 1. Preparation of four different types of semisolid forms and evaluation of their performance using in vitro diffusion method
- 2. Evaluation of test sterility for commercial preparations including sterile water for injection and
 - Antibiotic injection.
- 3. Collecting samples of environment of aseptic room and counting the colonies
- 4. Validation of one-unit operation (eg. Mixing) and development of protocol.
- 5. Comparative evaluation of different marketed products (tablets) of the same API
- 6. Dissolution studies of drug in three different bio relevant dissolution media
- 7. Stability study testing of tablet dosage forms (Any two products)



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COURSE STRUCTURE & SYLLABI

Course Code	ADVANCED DRUG DELIVERY SYSTEMS LAB	L	T	P	C
21S03206		0	0	6	3
	Semester		Ι	Ι	

List of Experiments:

- 1. Study on diffusion of drugs through various polymeric membranes (2 experiments)
- 2. Formulation and evaluation of sustained release oral matrix tablet (2 experiments)
- 3. Formulation and evaluation of sustained release oral reservoir system (2 experiments)
- 4. Formulation and evaluation of microspheres / microen capsules (2 experiments)
- 5. Study of in-vitro dissolution of various SR products in market (2 experiments)
- 6. Formulation and evaluation of transdermal films (2 experiments)
- 7. Formulation and evaluation mucoadhesive system (2 experiments)
- 8. Preparation and evaluation enteric coated pellets / tablets (2 experiments)



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

	RESEARCH METHODOLOGY AND	L	T	P	C
21DRM101	INTELLECTUAL PROPERTY RIGHTS	4	0	0	4
	Semester		I	II	
Course Objectives:					
	he research problem				
	erature studies, plagiarism and ethics				
 To get the know 	rledge about technical writing				
 To analyze the r 	nature of intellectual property rights and new developments				
 To know the pat 	tent rights				
Course Outcomes (CO): Student will be able to				
At the end of this course	e, students will be able to				
 Understand rese 	arch problem formulation.				
 Analyze researc 	h related information				
 Follow research 	ethics				
• Understand that	t today's world is controlled by Computer, Information	Tecl	hnol	ogy,	bu
tomorrow world	I will be ruled by ideas, concept, and creativity.				
 Understanding t 	that when IPR would take such important place in growth	of in	ndiv	idual	ls 8
nation, it is need	dless to emphasis the need of information about Intellectual	Prop	erty	Righ	nt to
be promoted am	ong students in general & engineering in particular.				
	IPR protection provides an incentive to inventors for furth				
	in R & D, which leads to creation of new and better production	lucts	, and	d in	turi
	onomic growth and social benefits.				
UNIT - I					
Meaning of research p	roblem, Sources of research problem, Criteria, Character	istic	s of	a g	3000
research problem, Error	s in selecting a research problem, Scope and objectives of r	esea	rch	probl	lem
Approaches of invest	igation of solutions for research problem, data coll-	ectio	n,	analy	ysis
interpretation, Necessary	y instrumentations				
UNIT - II					
Effective literature studi	es approaches, analysis, Plagiarism, Research ethics				
UNIT - III					
Effective technical writi	ng, how to write report, Paper Developing a Research Propo	sal,	Forn	nat o	f
	ntation and assessment by a review committee				
UNIT - IV					
	operty: Patents, Designs, Trade and Copyright. Process of Pa	atent	ing	and	
	gical research, innovation, patenting, development. Internation				

PCT.

UNIT - V

Patent Rights: Scope of Patent Rights. Licensing and transfer of technology. Patent information and databases. Geographical Indications. New Developments in IPR: Administration of Patent System. New developments in IPR; IPR of Biological Systems, Computer Software etc. Traditional knowledge Case Studies, IPR and IITs.

Reference Books:

1. Stuart Melville and Wayne Goddard, "Research methodology: an introduction for science & engineering students"



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COURSE STRUCTURE & SYLLABI

2. Wayne Goddard and Stuart Melville, "Research Methodology: An Introduction"



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

AUDIT COURSE-I



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

Course Objectives: This course will enable students: Understand the essentials of writing skills and their level of readability	Course Code	ENGLISH FOR RESEARCH PAPER WRITING	L	T	P	C
Course Objectives: This course will enable students: • Understand the essentials of writing skills and their level of readability • Learn about what to write in each section • Ensure qualitative presentation with linguistic accuracy Course Outcomes (CO): Student will be able to • Understand the significance of writing skills and the level of readability • Analyze and write title, abstract, different sections in research paper • Develop the skills needed while writing a research paper UNIT - I Lecture Hrs:10 1Overview of a Research Paper- Planning and Preparation- Word Order- Useful Phrases - Breaking up Long Sentences-Structuring Paragraphs and Sentences-Being Concise and Removing Redundancy -Avoiding Ambiguity UNIT - II Lecture Hrs:10 Essential Components of a Research Paper- Abstracts- Building Hypothesis-Research Problem - Highlight Findings- Hedging and Criticizing, Paraphrasing and Plagiarism, Cauterization UNIT - II Lecture Hrs:10 Introducing Review of the Literature – Methodology - Analysis of the Data-Findings - Discussion-Conclusions-Recommendations. UNIT - IV Lecture Hrs:9 Key skills needed for writing a Title, Abstract, and Introduction UNIT - V Lecture Hrs:9 Appropriate language to formulate Methodology, incorporate Results, put forth Arguments and draw Conclusions Suggested Reading 1. Goldbort R (2006) Writing for Science, Yale University Press (available on Google Books) Model Curriculum of Engineering & Technology PG Courses [Volume-I] 2. Day R (2006) How to Write and Publish a Scientific Paper, Cambridge University Press 3. Highman N (1998), Handbook of Writing for the Mathematical Sciences, SIAM. Highman'sbook 4. Adrian Wallwork , English for Writing Research Papers, Springer New York Dordrecht	21DAC101a		2	0	0	0
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M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

		DICACCEED NAARIACCERACERUS	L	T	P	C
21DAC101b		DISASTER MANAGEMENT	2	0	0	0
		Semester]	[
Course Objecti	ves: This cour	se will enable students:				
• Learn to	demonstrate	e critical understanding of key concepts in	n disas	ter risk	reducti	ion
	nanitarian resp					
 Critical 	y evaluatedisa	sterriskreduction and humanitarian response po	olicy and	d practic	e from	
Multiple	e perspectives.					
		ingofstandardsofhumanitarianresponseandpracti	icalrele	vanceins	specific	types
	ters and confli					
		estrengthsandweaknessesofdisastermanagemen				
	ımıng ın difter	ent countries, particularly their home country o	r the co	untries t	hey wo	rk in
UNIT - I Introduction:						
	4: E4	1C:;C:	N		.1	
		dSignificance;DifferenceBetweenHazardandDis	saster;N	aturaian	ıa	
		ce, Nature, Types and Magnitude.				
	A • T 1	•				
	e Areas in Ind		1 4			D
Study of Seism	nic Zones; Are	as Prone to Floods and Droughts, Landslides a				
Study of Seism to Cyclonic and	nic Zones; Are					
Study of Seism to Cyclonic as Epidemics	nic Zones; Are	as Prone to Floods and Droughts, Landslides a				
Study of Seism to Cyclonic at Epidemics UNIT - II	nic Zones; Are and Coastal Ha	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; F				
Study of Seism to Cyclonic an Epidemics UNIT - II Repercussions	of Disasters	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; Fand Hazards:	Post- D	isaster l	Disease	s and
Study of Seism to Cyclonic and Epidemics UNIT - II Repercussions Economic Dar	of Disasters anage, Loss of	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; Fand Hazards: Human and Animal Life, Destruction of Economics	Post- Di	isaster l	Disease	s and
Study of Seism to Cyclonic an Epidemics UNIT - II Repercussions Economic Dar Earthquakes, Vo	of Disasters anage, Loss of Dicanisms, Cyc	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; Fand Hazards: Human and Animal Life, Destruction of Ecolones, Tsunamis, Floods, Droughts and Famines, Landslides and Property and Proper	Post- D	n. Natur	Disease ral Disa Avalan	asters:
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Study of Seism to Cyclonic and Epidemics UNIT - II Repercussions Economic Dar Earthquakes, Von Man-made disardisease and Epunit - III Disaster Preparation Cyclonic American Cyclonic Cycloni	of Disasters and Loss of Disasters of Disasters of Disasters of Disasters. Nuclear idemics, War	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; Fand Hazards: Human and Animal Life, Destruction of Ecolones, Tsunamis, Floods, Droughts and Famines, La Reactor Meltdown, Industrial Accidents, Oil Sliand Conflicts.	Post- Di cosystem andslide icks and	n. Natur s and l Spills,	Tal Disa Avalar Outbrea	asters:
Study of Seism to Cyclonic at Epidemics UNIT - II Repercussions Economic Dar Earthquakes, Vo Man-made disa Disease and Epunit - III Disaster Preparedness:	of Disasters and Loss of Disasters of Disasters of Disasters of Disasters. Nuclear didemics, War	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; Fand Hazards: Human and Animal Life, Destruction of Ecolones, Tsunamis, Floods, Droughts and Famines, La Reactor Meltdown, Industrial Accidents, Oil Sli and Conflicts. Management:	cosystem andslide acks and	n. Natures and Spills,	ral Disa Avalar Outbrea	asters: nches, aks of
Study of Seism to Cyclonic at Epidemics UNIT - II Repercussions Economic Dar Earthquakes, Vo Man-made disa Disease and Epunit - III Disaster Preparedness: Application of	of Disasters and Loss of Dicanisms, Cyclester: Nuclear idemics, War Monitoring of Remote Sense.	as Prone to Floods and Droughts, Landslides as azards with Special Reference to Tsunami; Fand Hazards: Human and Animal Life, Destruction of Ecolones, Tsunamis, Floods, Droughts and Famines, La Reactor Meltdown, Industrial Accidents, Oil Sliand Conflicts. Management: of Phenomena Triggering ADisasteror Hazards	cosystem andslide acks and	n. Natures and Spills,	ral Disa Avalar Outbrea	asters: nches, aks of

Risk Assessment Disaster Risk:

Concept and Elements, Disaster Risk Reduction, Global and National Disaster Risk Situation. TechniquesofRiskAssessment,GlobalCo-OperationinRiskAssessmentand Warning, People's Participation in Risk Assessment. Strategies for Survival.

UNIT - V

Disaster Mitigation:

Meaning, Conceptand Strategies of Disaster Mitigation, Emerging Trends In Mitigation. Structural Mitigation and Non-Structural Mitigation, Programs of Disaster Mitigation in India.

Suggested Reading



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- 1. R.Nishith, Singh AK, "Disaster Management in India: Perspectives, issues and strategies
- 2. "'New Royal book Company..Sahni,PardeepEt.Al.(Eds.),"DisasterMitigationExperiencesAndReflections",PrenticeHa ll OfIndia, New Delhi.
- 3. GoelS.L.,DisasterAdministrationAndManagementTextAndCaseStudies",Deep&Deep Publication Pvt. Ltd., New Delhi



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

Course Code	SANSKI	RITFOR TECHNICAL KNOWLEDGE	L	T	P	C
21DAC101c			2	0	0	0
		Semester	•		I	
G 01: 4:	771:	11 11 . 1 .				
Course Objecti	ves: This cours	se will enable students:				
•	•	ledge in illustrious Sanskrit, the scientific lar	iguage ii	n the wo	orld	
· ·	_	improve brain functioning				
	-	evelopthelogicinmathematics, science & others	ıbjects e	nhancin	g the	
memory						
_		ars equipped with Sanskrit will be able to exp	lore the	huge		
	edge from ancie					
		ent will be able to				
	•	anskrit language				
		ture about science &technology can be unders	tood			
• Being a UNIT - I	logical langua	ge will help to develop logic in students				
	1					
Alphabets in Sa	anskrit,					
UNIT - II		1.0				
Past/Present/Fut UNIT - III	ure Tense, Sim	ple Sentences				
Order, Introduct	tion of mosts					
-	.1011 01 10018		1			
UNIT - IV						
	rmation about S	Sanskrit Literature	1			
UNIT - V						
Technical conc	epts of Engine	ering-Electrical, Mechanical, Architecture, Ma	thematic	es		
Suggested Read						
1."Abhyaspust	akam" –Dr.V	ishwas, Sanskrit-Bharti Publication, New	Delhi			
2."Teach You	rself Sanski	rit" Prathama Deeksha- VempatiKutur	nbshastı	ri, Rash	triyaSa	ınskrit
Sansthanam, N						
3."India's Glo	rious Scientifi	cTradition" Suresh Soni, Ocean books (P	Ltd.,N	ew Del	hi	



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COURSE STRUCTURE & SYLLABI

AUDIT COURSE-II



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

Course Code		PEDAGOGY STUDIES	I		T	P	С
21DAC201a		TEDAGOGI STODIES	2	2	0	0	0
		Semeste	r		I	T T	
		Demeste	1			<u>.</u>	
Course Objecti	ves: This cours	se will enable students:					
Reviewe	existingevidence	ceonthereviewtopictoinformprogrammedesig	nandpo	olicy n	nakin	ng	
undertal	ken by the DfII	O, other agencies and researchers.					
 Identify 	critical eviden	ce gaps to guide the development.					
Course Outcom	nes (CO): Stud	ent will be able to					
Students will be							
Whatped countries		icesarebeingusedbyteachersinformalandinfor	malcla	ssroon	ns in	develo	ping
• What is	the evidence o	n the effectiveness of these pedagogical practices of the pedagogical prac	tices, i	n wha	t		
		hat population of learners?					
		on(curriculumandpracticum)andtheschoolcu	rıculuı	mand g	guida	ance	
	s best support	effective pedagogy?					
UNIT - I							
terminology	Theories	ogy: Aims and rationale, Policy back groun oflearning, Curriculum, Teachereducation. Cology and Searching.					
UNIT - II							
		ogical practices are being used by teacher. Curriculum, Teacher education.	ers in	forma	ıl an	d inf	ormal
UNIT - III							
of included stu guidance mater evidence for e	idies. How car ials best suppo ffective pedago	ofpedagogicalpractices, Methodology for the in a teacher education (curriculum and practiculum rt effective pedagogy? Theory of change. Strogical practices. Pedagogic theory and pedagogic strategies.	n) and ength	thesch and na	o cui	rriculur of th bo	n and ody of
UNIT - IV							
Support from the teacher and the considerations sizes	ne head	lignment with classroom practices and follow riculumandassessment, Barrierstolearning: lim					
UNIT - V							

Researchgapsandfuturedirections: Researchdesign, Contexts, Pedagogy, Teachereducation, Curriculum and assessment, Dissemination and research impact.

Suggested Reading

- 1. AckersJ, HardmanF(2001) Classroom interaction in Kenyan primary schools, Compare, 31 (2): 245-261.
- 2. AgrawalM(2004)Curricularreforminschools:Theimportanceofevaluation,Journalof



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- 3. Curriculum Studies, 36 (3): 361-379.
- 4. AkyeampongK(2003) Teacher training in Ghana does it count? Multi-site teachereducation research project (MUSTER) country report 1. London: DFID.
- 5. Akyeampong K, LussierK, PryorJ, Westbrook J (2013)Improving teaching and learning of basic maths and reading in Africa: Does teacherpreparation count?International Journal Educational Development, 33 (3): 272–282.
- 6. Alexander RJ(2001) Culture and pedagogy: International comparisons in primary education. Oxford and Boston: Blackwell.
 - Chavan M (2003)ReadIndia: A mass scale, rapid, 'learning to read'campaign.
- 7. www.pratham.org/images/resource%20working%20paper%202.pdf.



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Course Code	CEDI			L	T	P	C
21DAC201b	SIKI	ESSMANAGEMENT BY YOGA		2	0	0	0
		Se	emester		I	I	
Course Objecti	ves: This course	will enable students:					
To achie	eve overall healt	h of body and mind					
To over	come stres						
Course Outcom	nes (CO): Studer	nt will be able to					
_	healthy mind ir efficiency	a healthy body thus improving soci	al health a	also			
UNIT - I							
Definitions of I	Eight parts of yo	g.(Ashtanga)					
UNIT - II							
Yam and Niyar	n.						
UNIT - III							
Do`sand Don't	sin life.						
	•	charyaand aparigrahaii) ,ishwarpranidhan					
UNIT - IV		•					
Asan and Prana	ıyam						
UNIT - V							
i)Variousyogpo	sesand theirben	efitsformind &body					
ii)Regularizatio	onofbreathingtecl	hniques and its effects-Types ofprana	ayam				
Suggested Read							
		ng-Part-I": Janardan SwamiYogabh					
		Internal Nature" by Swami Viv	vekananda	a, Adv	aita		
Ashrama (Public	cation Department	nt), Kolkata					



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

Course Code	PERSONALI	TY DEVELOPMENT THRO	OUGHLIFE	L	T	P	С
21DAC201c		NLIGHTENMENTSKILLS	_	2	0	0	0
			Semester		I	Ι	
Course Objectiv	ves: This course	will enable students:					
		ghest goal happily					
	•	stable mind, pleasing persona	ality and determ	nination	l		
	ken wisdom in stu						
	nes (CO): Studen						
		d-Geetawillhelpthestudentind	evelopinghispe	ersonali	yand ac	chieve	
_	est goal in life						
•		ed Geetawilllead the nation a		•	•	perity	
	f Neetishatakam v	vill help in developing versati	le personality	of stude	nts		
UNIT - I							
	-	nent of personality					
	20,21,22(wisdom)						
Verses-29,3	31,32(pride &hero	oism)					
	28,63,65(virtue)						
UNIT - II							
Neetisatakam- l	Holistic developn	nent of personality					
Verses-52,5	53,59(dont's)						
Verses-71,7	73,75,78(do's)						
UNIT - III							
Approach to da	y to day work and	d duties.					
ShrimadBh	agwadGeeta:Cha	pter2-Verses41,47,48,					
Chapter3-V	Verses 13, 21, 27, 35	,Chapter6-Verses5,13,17,23,3	5,				
Chapter 18-	Verses45,46,48.						
UNIT - IV							
Statements of b	asic knowledge.						
ShrimadBh	agwadGeeta:Cha	pter2-Verses 56,62,68					
	-Verses 13, 14, 15, 1						
•		hrimad Bhagwad Geeta:					
UNIT - V		<u> </u>					
Chapter 2-V	Verses 17, Chapter	3-Verses36,37,42,					
*	Verses 18, 38, 39	•					
*	- Verses37,38,63						
Suggested Read							
		niSwarupanandaAdvaitaAshra	am(Publication	Departr	nent),		
Kolkata	-	-		_			
	,	iti-sringar-vairagya) by P.Go	opinath, Rasht	riyaSan	skrit		
Sansthanam,	New Delhi.						



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COURSE STRUCTURE & SYLLABI

OPEN ELECTIVE



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

Course Code	BIOLOGICAL SCREENING METHODS	L	T	P	C
21SOE301d	(Elective)	3	0	0	3
	Semester		IJ	Ι	
Course Objectives:					
	ng to study about various techniques for screening of drugs				
	ological activities and guide lines for handling animals and huma	an an	d an	imal	
ethics for screening of					
-	CO): Student will be able to				
	nes are students will know how to handle animals and know				
	ques for screening of drugs for different pharmacological activit	ies, g	guide	lines	,
	creening new drug molecules on animals.				
UNIT - I					
Drug discovery proc	ess: Principles, techniques and strategies used in new drug disco	very.	. Hig	<u>h</u>	
throughput screening	g, human genomics, robotics and economics of drug discovery, I	Regul	latio	ns.	
Alternatives to anima	al screening procedures, cell-line, patch -clamp technique, In-vi	tro n	node	ls,	
molecular biology te	chniques				
UNIT - II					
Bioassays: Basic prin	nciples of bioassays, official bioassays, experimental models and	d stat	istica	al	
designs employed in	biological standardization.				
UNIT - III					
Principles of toxicity	evaluations, ED50, LD50 and TD values, International guideling	nes (I	СН		
recommendations).		`			
Preclinical studies: C	General principles and procedures involved in acute, sub-acute, of	chron	ic,		
	genicity and carcinogenicity				
UNIT - IV					
Screening of differen	at classes of drugs using micro-organisms. Vitamin and antibioti	c ass	ays.		
Screening methods is	nvolved in toxins and pathogens.		·		
_		1			
UNIT - V					
	ng methods: α-glucosidase, α- amylase, DNA polyme	rase,	nu	cleas	ses,
Lasparginase, lipases	s and peptidases.				
Reference Books:					
	pharmacology by Bertram G. Katzung (International edition) la	nge n	nedio	cal	
book / Mc Graw Hill	, USA 2001 8th edition				

- 2. Pharmacology by Rang H.P, Dale MM and Ritter JM., Churchill Livingston, London, 4/e
- 3. Goodman and Gilman's The pharmacological basis of therapeutics (International edition) Mc Graw Hill, USA 2001 10th edition.
- 4. General and applid toxicology by B.Ballantyne, T.Marrs, P.Turner (Eds) The Mc Millan press Ltd, London.
- 5. Drug Discovery by Vogel's
- 6. Drug Discovery and evaluation Pharmacological assays by H.Gerhard. Vogel, 2nd edition, Springer verlag, Berlin, Heidelberg.
- 7. Tutorial Pharmacy (Vol I and II) by Cooper and Gunns.



M.PHARM. IN PHARMACEUTICAL TECHNOLOGY

COURSE STRUCTURE & SYLLABI

Course Code	PHARMACEUTICAL VALIDATION	L	T	P	C
21SOE301a	(Elective)	3	0	0	3
	Semester		I	I	
Course Objectives:					
	f the subject is to understand about validation and how it can be				
	improve the quality of the products. The subject covers the comp	plete	info	rmat	ion
about validation, typ	pes, methodology and application				
Course Outcomes	(CO): Student will be able to				
Course Outcome:	Upon completion of the subject student shall be able to				
 Explain the 	aspect of validation				
 Carryout va 	lidation of manufacturing processes				
• Apply the k	nowledge of validation to instruments and equipments				
* * *	e manufacturing facilities				
UNIT - I	<i>g</i>				
Introduction: Defin	ition of Qualification and Validation, Advantage of Validation	Str	eaml	inino	of
	lidation process and Validation Master Plan. Qualification: U				
	gn Qualification, Factory Acceptance Test (FAT)/ Site Accepta				
	cation, Operational Qualification, Performance Qualification, I			-	
	-Calibration Preventive Maintenance, Change management),				
	pment, Qualification of Analytical Instruments and Laboratory	_			
UNIT - II					
Qualification of	analytical instruments: Electronic balance, pH met	er,	UV	-Visi	ble
spectrophotometer,	FTIR, GC, HPLC, HPTLC				
Qualification of Gla	ssware: Volumetric flask, pipette, Measuring cylinder, beakers a	nd b	urett	e.	
UNIT - III					
Qualification of lab	oratory equipments: Hardness tester, Friability test apparatus, t	ap d	ensit	y tes	ter,
	, Dissolution test apparatus.	-			
Validation of Utility	systems: Pharmaceutical water system & pure steam, HVAC sy	sten	1,		
Compressed air and	nitrogen.				
UNIT - IV					
Cleaning Validation	: Cleaning Validation - Cleaning Method development, Validation	on a	nd va	lidat	ion
of analytical metho	d used in cleaning. Cleaning of Equipment. Cleaning of Facili	ties.	Clea	ning	; in
place (CIP).	· · · · · · · · · · · · · · · · · · ·				
TINITE X7					

Analytical method validation: General principles, Validation of analytical method as per ICH guidelines and USP.

Reference Books:

- 1. T. Loftus & R. A. Nash, "Pharmaceutical Process Validation", Drugs and Pharm Sci. Series, Vol.129, 3rd Ed., Marcel Dekker Inc., N.Y.
- 2. The Theory & Practice of Industrial Pharmacy, 3rd edition, Leon Lachman, Herbert A. Lieberman, Joseph. L. Karig, Varghese Publishing House, Bombay.
- 3. Validation Master plan by Terveeks or Deeks, Davis Harwood International publishing.
- 4. Validation of Aseptic Pharmaceutical Processes, 2nd Edition, by Carleton & Agalloco, (Marcel
- 5. Michael Levin, Pharmaceutical Process Scale-Upl, Drugs and Pharm. Sci. Series, Vol. 157, 2nd Ed., Marcel Dekker Inc., N.Y.



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COURSE STRUCTURE & SYLLABI

	1 3	T	P	\mathbf{C}
	J	0	0	3
Schrester	III			
Course Objectives:				
This course is designed to impart knowledge and skills necessary to train t	he	stude	ents	on
entrepreneurship management.				
Course Outcomes (CO): Student will be able to				
On completion of this course it is expected that students will be able to:				
• The Role of enterprise in national and global economy				
Dynamics of motivation and concepts of entrepreneurship				
Demands and challenges of Growth Strategies and Networking				
UNIT - I				
Conceptual Frame Work: Concept need and process in entrepreneurship develop	nme	ent.	Role	of
enterprise in national and global economy. Types of enterprise – Merits and Demer				
policies and schemes for enterprise development. Institutional support in enterprise d				
management.		10p11		
UNIT - II				
Entrepreneur: Entrepreneurial motivation – dynamics of motivation. Entrepreneurial of	com	pete	ncv -	
Concepts. Developing Entrepreneurial competencies - requirements and understanding				
entrepreneurship development, self-awareness, interpersonal skills, creativity,				
achievement, factors affecting entrepreneur role.				,
UNIT - III				
Launching and Organizing an Enterprise: Environment scanning – Information, sour	rces	, sch	emes	s of
assistance, problems. Enterprise selection, market assessment, enterprise feasibility	y st	udy,	SW	TO
Analysis. Resource mobilization -finance, technology, raw material, site and manpov	ver.	Cost	ting a	and
marketing management and quality control. Feedback, monitoring and evaluation.				
UNIT - IV				
Growth Strategies and Networking: Performance appraisal and assessment. Profitable	ility	and	con	trol
	Te	chni	ques	of
measures, demands and challenges. Need for diversification. Future Growth -				
measures, demands and challenges. Need for diversification. Future Growth – expansion and diversification, vision strategies. Concept and dynamics. Methods		oint '	ventı	ure,
		oint '	ventı	ure,
expansion and diversification, vision strategies. Concept and dynamics. Methods		oint '	ventı	ure,

Reference Books:

resource mobilization and implementation.

- 1. Akhauri, M. M. P.(1990): Entrepreneurship for Women in India, NIESBUD, New Delhi.
- 2. Hisrich, R. D & Brush, C.G. (1996) The Women Entrepreneurs, D.C. Health& Co., Toranto.
- 3. Hisrich, R.D. and Peters, M.P. (1995): Entrepreneurship Starting Developing and Managing a New Enterprise, Richard D., Inwin, INC, USA.
- 4. Meredith, G.G. et al (1982): Practice of Entrepreneurship, ILO, Geneva.
- 5. Patel, V.C. (1987): Women Entrepreneurship Developing New Entrepreneurs, Ahmedabad EDII
- 6. Arya kumar.(2012): Entrepreneurship- Creating and Leading an Entrepreneurial Organization, Pearson