

SRI VENKATESWARA COLLEGE OF PHARMACY

[AUTONOMOUS] Approved by AICTE & PCI, New Delhi, Permanently Affiliated to JNTUA, Ananthapuramu Accredited by NBA, New Delhi for UG Programme under Tier-II & NAAC, Bengaluru Recognized under section 2(f) & 12 (B) of UGC Act, 1956 Recognized Research Centre for Pharmaceutical Sciences by JNTUA RVS Nagar, Tirupati Road, Chittoor - 517127, Andhra Pradesh

M.PHARM IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

SEMESTER – I

S.	Course	Course Name	H	lours	per	Credits
No.	codes		L	Т	Р	
1.	21S01101	Modern Pharmaceutical Analytical Techniques	4	-	-	4
2.	21S03101	Advanced Physical Pharmaceutics	4	-	-	4
3.	21S03102	Modern Pharmaceutics-I	4	-	-	4
4.	21S03103	Advanced Biopharmaceutics & Pharmacokinetics	4	-	-	4
5.	21S01105	Modern Pharmaceutical Analytical Techniques lab	-	-	6	3
6.	21S03104	Modern Pharmaceutics -I lab	-	-	6	3
7.	21DAC101a 21DAC101b 21DAC101c	Audit Course – I English for Research paper writingDisaster Management Sanskrit for Technical Knowledge	2	-	-	0
8.	21S03105	Seminar/Assignment	-	1	6	4
		Total	18	1	18	26

SEMESTER – II

S.No.	Course	Course Name	H	ours	per	Credita
	codes		L	Т	P	Creuits
1.	21S03201	Modern Pharmaceutics-II	4	-	-	4
2.	21S03202	Advanced Drug Delivery system	4	-	-	4
3.	21S03203	Industrial Pharmacy	4	-	-	4
4.	21S03204	Nano Drug Delivery system	4	-	-	4
5.	21S03205	Modern Pharmaceutics-II Lab	-	-	6	3
6.	21S03206	Advanced Drug Delivery System Lab	-	-	6	3
7.	21DAC201a 21DAC201b 21DAC201c	Audit Course – II Pedagogy Studies Stress Management for Yoga Personality Development through Life Enlightenment Skills	2	-	-	0
8.	21S03207	Seminar/Assignment	-	1	6	4
		Total	18	1	18	26



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COURSE STRUCTURE & SYLLABI

SEMSTER - III

S.No.	Course	Course Name	Ho	urs po	er	Credits
	codes			Т	P	
1.	21DRM101	Research Methodology and Intellectual Property Right	4	-	-	4
2.	21SOE301d 21SOE301a 21SOE301c	Open Elective Biological Screening methods Pharmaceutical Validation Entrepreneurship Management	3	-	-	3
3.	21S03301	Teaching Practice/Assignment	-	-	4	2
4.	21803302	Comprehensive viva voce	-	-	-	2
5.	21\$03303	Research Work - I	-		24	12
		Total	7	-	32	23

SEMESTER - IV

S.No.	Course	Course Name	Hours per week			Credits
	codes		L	Т	P	
1.	21S03401	Co-Curricular Activities	2			2
2.	21S03402	Research Work - II	3		30	18
		Total	5		30	20



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Course Code	MODERN PHARMACEUTICAL ANALYTICAL	LT	Р	С
21S01101	TECHNIQUES	4 0	0	4
	Semester		Ι	
Course Objectives:				
This subject deals w	ith various advanced analytical instrumental techniques for idea	ntificatio	on,	
characterization and	quantification of drugs. Instruments dealt are NMR, Mass s	pectrom	eter,	IR,
HPLC, GC etc.		_		
Course Outcomes (CO): Student will be able to			
After completion o	f course student is able to know about chemicals and excip	ients.		
• The analysis	of various drugs in single and combination dosage forms			
Theoretical	and practical skills of the instruments			
UNIT - I				
UV-Visible spectros	copy: Introduction, Theory, Laws, Instrumentation associated	with UV	/-Visi	ble
spectroscopy. Choic	e of solvents and solvent effect and Applications of UV-Visil	ole spect	rosco	pv.
Difference/ Derivati	ve spectroscopy.	r		r <i>J</i> ,
UNIT - II				
IR spectroscopy: Th	eory. Modes of Molecular vibrations. Sample handling. Instru	mentatio	on of	
Dispersive and Four	ier -Transform IR Spectrometer. Factors affecting vibrational	frequer	cies a	and
Applications of IR si	pectroscopy. Data Interpretation.			
UNIT - III				
NMR spectroscopy:	Ouantum numbers and their role in NMR. Principle. Instrum	entation	. Solv	vent
requirement in NMR	Relaxation process. NMR signals in various compounds. Chem	ical shif	t. Faci	tors
influencing chemica	l shift. Spin-Spin coupling. Coupling constant. Nuclear magneti	ic double	Э	
resonance. Brief out	line of principles of FT-NMR and 13C NMR. Applications of N	MR spec	troscc	opy.
		1		1.
UNIT - IV				
Mass Spectroscopy:	Principle, Theory, Instrumentation of Mass Spectroscopy, D	ifferent	types	s of
ionization like elect	ron impact, chemical, field, FAB and MALDI, APCI, ESI, Al	PPI Ana	lyzers	s of
Quadrupole and Tin	ne of Flight, Mass fragmentation and its rules, Meta stable ion	s, Isotop	ic pea	aks
and Applications of	Mass spectroscopy.			
UNIT - V				
Chromatography				
Introduction to chron	natography and classification of chromatographic methods base	d on the		
mechanism of sepa	aration, Principle, instrumentation, selection of solvents; chro	omatogr	aphic	
parameters, factors a	ffecting resolution, applications of the following:			
a) Thin Layer chrom	atography; b) High Performance Thin Layer Chro	matogra	phy	
c) Paper Chromatogr	caphy; d) Column chromatography			
e) Gas chromatograp	bhy; f) High Performance Liquid chromatog	raphy		
g) Affinity chromato	graphy; h) Gel Chromatography			
i)Hyphenated techni	ques :			
Ultra High F	Performance Liquid chromatography- Mass spectroscopy			
Gas Chroma	tography-Mass Spectroscopy			
Reference Books:				
1. Instrumental M	ethods of Chemical Analysis by B.K Sharma			
2. Vogel's Text be	bok of Quantitative Chemical Analysis by A.I. Vogel			
3. Spectrometric I	dentification of Organic compounds - Robert M Silverstein, Si	xth editi	on, Jo	ohn
Wiley & Sons,	2004.			
	2			



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- 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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21503101 ADVARCEDTIFISTICAL HIMMACENTICS 4 0 0 4 Semester I Course Objectives: The students shall know about particle science, polymer science and its use in pharmaceutical dosage forms. They also know the compression and consolidation parameters for powders and granules. Students also know about the rheology, disperse systems, dissolution and solubility parameters for dosage forms. Course Outcomes (CO): Student will be able to The students will know particle size analysis method, solid dispersion, physics of tablets, polymer classification and its applications, student will also know the stability calculations, shelf life calculations and accelerated stability studies. They also know the factors affecting the dissolution and solubility in related to invitro/invivo correlations. UNIT - I Polymer science: Classification, properties and characterization of polymers in pharmaceutical formulations. Mechanism of biodegradation of biodegradable polymers in cluding controlled drug delivery systems, Mucoadhesive, Hydrodynamically balanced and Transdermal Systems. UNIT - II Physics of tablet compression: Basic principles of interactions, compression and consolidation, compression and consolidation, under high loads, effect of friction, distribution of forces in compaction, force volume relationships, Heckel plots, compaction profiles, energy involved in compaction, Measurement of compression with strain gauges, compression pressure-QA parameters. UNI	Course Code	ADVANCED PHYSICAL PHAPMACEUTICS	L	Т	P	C
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formulations. Mechanism of biodegradation of biodegradable polymers including controlled drug delivery systems, Mucoadhesive, Hydrodynamically balanced and Transdermal Systems. UNIT - II Physics of tablet compression: Basic principles of interactions, compression and consolidation, compression and consolidation under high loads, effect of friction, distribution of forces in compaction, force volume relationships, Heckel plots, compaction profiles, energy involved in compaction, Measurement of compression with strain gauges, compression pressure-QA parameters. UNIT - III Kinetics and drug stability: Stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications.	polymers in solid si	tate, preparation of polymer solution, application of polymers i	n pł	narma	aceuti	ical
UNIT - II Physics of tablet compression: Basic principles of interactions, compression and consolidation, compression and consolidation under high loads, effect of friction, distribution of forces in compaction, force volume relationships, Heckel plots, compaction profiles, energy involved in compaction, Measurement of compression with strain gauges, compression pressure-QA parameters. UNIT - III Image: Compression with strain gauges, compression pressure-QA parameters. UNIT - III Image: Compression with strain gauges, compression pressure-QA parameters. UNIT - III Image: Compression with strain gauges, compression pressure-QA parameters. UNIT - III Image: Compression with strain gauges, compression pressure-QA parameters. UNIT - III Image: Compression with strain gauges, complex order kinetics, Factors influencing stability, strategy of stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Image: Compression of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications. UNIT - V	formulations. Mech	anism of biodegradation of biodegradable polymers including		ntroll	ed di	rug
Physics of tablet compression: Basic principles of interactions, compression and consolidation, compression and consolidation under high loads, effect of friction, distribution of forces in compaction, force volume relationships, Heckel plots, compaction profiles, energy involved in compaction, Measurement of compression with strain gauges, compression pressure-QA parameters. UNIT - III Kinetics and drug stability: Stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentations.	UNIT - II			•		
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compaction, force volume relationships, Heckel plots, compaction profiles, energy involved in compaction, Measurement of compression with strain gauges, compression pressure-QA parameters. UNIT - III Kinetics and drug stability: Stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications and interpretations.	compression and co	onsolidation under high loads, effect of friction, distribution of	for	ces in	1	,
compaction, Measurement of compression with strain gauges, compression pressure-QA parameters. UNIT - III Kinetics and drug stability: Stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentations.	compaction, force	volume relationships, Heckel plots, compaction profiles, energy	inv	olvec	l in	
UNIT - III Kinetics and drug stability: Stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: UNIT - V Interpretations	compaction, Measu	rement of compression with strain gauges, compression pressure	-QA	para	meter	rs.
Kinetics and drug stability: Stability calculations, rate equations, complex order kinetics, Factors influencing stability, strategy of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV	UNIT - III		1.		-	
Initial energy stability, surgey of stability testing, method of stabilization, method of accelerated stability testing in dosage forms, temperature and humidity control, physical stability testing of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: UNIT - V	Kinetics and drug	stability: Stability calculations, rate equations, complex order	kin	etics,	Fact	tors
statistic in dosage forms, temperature and numberly control, physical statistic esting of pharmaceutical products. Photodecomposition, Method, solid state decomposition. UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentations. UNIT - V Instrumentations	stability testing in	dosage forms temperature and humidity control physical sta	ahili	i acc	sting	of
UNIT - IV Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications and interpretations. UNIT - V UNIT - V	pharmaceutical proc	lucts. Photodecomposition, Method, solid state decomposition.	uom	ly le	sting	01
Theoretical consideration, instrumentation, rheological properties of disperse systems and semisolids. Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications and interpretations.	UNIT - IV					
Oscillatory testing, Creep measurement. Characterization of API and excipients: Differential Scanning Calorimetry: Principle, thermal transitions, advantages, disadvantages, instrumentation, applications and interpretations X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications and interpretations.	Theoretical conside	tration instrumentation rheological properties of disperse system	s an	d sen	nisoli	ds
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X Ray Diffraction methods: Origin of x-rays, principle, advantages, disadvantages, instrumentation, applications and interpretations.	transitions, advantag	ges, disadvantages, instrumentation, applications and interpretati	ons			
Instrumentation, applications and interpretations.	X Ray Diffract	ion methods: Origin of x-rays, principle, advantages	, d	isadv	antag	ges,
	Instrumentation, apj					
			ļ	1	.1	
Dissolution and solubility: Solubility and solubilization of nonelectrolytes, solubilization by the use of surfactants, cosplyants, complexation, drug derivatization and solid state manipulation. Machanisma	Dissolution and sol	ubility: Solubility and solubilization of nonelectrolytes, solubility vents, complexation, drug derivatization and solid state manipula	zati	on by Mac	the hand	use
of Drug release - dissolution diffusion (Matrix and Reservoir) and swelling controlled	of Drug release - di	ssolution diffusion (Matrix and Reservoir) and swelling control	lled	11100	114111	51115
(Porpos Model) and dissolution againment		contraction (intraction and resortion) and swelling control				



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Textbooks:

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

Course Code	MODERN PHARMACEUTICS – I		1	P	
21503102	Somostar	4	U	U	4
	Semester				
Course Objectiv	0.04				
Students will kno	us the preformulation studies, methodology, different excinients us	ad ir		id	
dosage forms an	d their evaluation with references to production technologies. T	be c	i sui tude	iu nte a	ادم
know the optimiz	ation techniques and their applications in pharmaceutical industries	ne s	iuuc.	into a	150
Course Outcome	s (CO): Student will be able to				
Students shall ex	plain the preformulation parameters apply ICH guidelines and ex	alua	te dr	na q	riio
excipients compa	tibility Students also explain about formulation and development	use	of ex	ag, a cinie	ents
in tablets, powder	s, capsules, micro-encapsules and coating techniques. They also le	arn a	and a	pply	the
statistical design i	in different formulations.			FF-J	
UNIT - I					
Preformulation	studies: Goals of Preformulation preformulation parameters	Poly	mor	nhs	and
Amorphous form	s selection of drugs- solubility partition coefficient salt forms hu	nidi	tv s	olids	tate
properties. Partic	le Size Analysis (Laser Diffraction and Dynamic Light Scattering)	dru	ig - e	xcin	ien
compatibility. flo	w properties, format and content of reports of preformulation.		0	P	4
preformulation sta	ability studies (ICH)				
UNIT - II	•				
Formulation dev	relopment of solid dosage forms – I: New materials, excipients s	cien	ce - c	lilue	nts,
disintegrants, sup	er disintegrants, etc, evaluation of functional properties of excipient	nts, o	co-pr	oces	sed
materials, method	ls of preparation and evaluation.		•		
UNIT - III					
Formulation dev	elopment of solid dosage forms-II: Coating, coating machines, co	oatir	ng teo	chniq	ues
in tablet techno	logy for product development, computerization, inprocess co	ntro	l of	tabl	ets.
formulation devel	opment and manufacture of powder dosage forms for internal use.				
Microencapsulat	ion- types, methodology, problems encountered.				
UNIT - IV					
Formulation dev	relopment of soft and hard gelatin capsules: Introduction, prod	uctio	on ar	nd	
methods of manu	ifacture, filling equipment and filling operations, formulations, f	inisł	ning,	spec	cial
techniques, advar	ices in capsule manufacture, machines, processing and control incl	udın	ıg		
pharmaceutical as	spects, physical stability and packaging.				
$\frac{\text{UNII - V}}{\text{O}(1) + \frac{1}{2}}$			T /	1 (
Optimization to	echniques in pharmaceutical formulation and processin	g:	Intro	duct	ion.
design nortial for	atorial design, simpley methods, mixture designs, Diseket Purh	agra	ms,		
Benken method	upplications in pharmaceutical formulation		neun	Ju, L	OX
Textbooks.	ipplications in pharmaceutear formulation.				
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	Sch	wartz	
3. Pharmaceutical	Dosage forms - Capsules (Vol I, II and III) by Avis, Lieberman and	d La	ichm	an	-•
4. Pharmaceutica	1 Dosage forms – Disperse systems (Vol I. II and III) by Avis. Lief	erm	an ai	nd	
Lachman.					
5. Modern Pharm	aceutics by Gilbert S. Banker and Christopher T. Rhodes.				
6. Pharmaceutica	l statistics by Bolton				



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



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COURSE STRUCTURE & SYLLABI

Course Code	ADVANCED BIOPHARMACEUTICS &	L	Т	Р	С
21S03103	PHARMACOKINETICS	4	0	0	4
	C			Г	

Semester

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.



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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.

Clinical Pharmacokinetics: Altered kinetics in pregnancy, child birth, infants and geriatrics. Kinetics in GI disease, mal absorption 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	MODERN PHARMACEUTICAL ANALYTICAL	L	Т	Р	C
Code	TECHNIQUES LAB		0		
21S01105		0	0	6	3
	Semester]	[
List of Experiments	8				
1. Analysis of Pharm	nacopoeial compounds and their formulations by UV Vis Spectro	opho	tome	ter.	
2. Simultaneous esti	mation of multi component containing formulations by UV Spec	trop	hotoi	netry	7
3. Effect of pH and s	solvent on UV –Spectrum				
4. Determination of	Molar absorption coefficient				
5. Estimation of ribo	flavin/ quinine sulphate by fluorimetry				
6. Study of quenchin	g effect by fluorimetry				
7. Estimation of sodi	um or potassium by flame photometry				
8. Colorimetric deter	mination of drugs by using different reagents				
9. Quantitative deter	mination of functional groups				
10. Experiments bas	ed on Column chromatography				
11. Experiments bas	ed on HPLC				
12. Experiments bas	ed on Gas Chromatography				



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Course Code		L	Т	P	C
21S03104	MODEKN PHARMACEUTICS – I LAB	0	0	6	3
	Semester]	I	
List of Experiments	8				
1. To carry out the p	reformulation studies of solid dosage forms.				
2. To study the effect	et of compressional force on tablet disintegration time				
3. To study the micr	omeritic properties of powders and granules				
4. To study the effect	et of particle size on dissolution of tablets				
5. To study the effect	et of binders on dissolution of tablets				
6. To study pharmac	cokinetic models, to determine similarity factors				
7. Accelerated stabil	lity testing of different tablets				
8. Determination of	first order, second order rate constants by acid and alkaline hydr	olys	is		
9. Preparation and e	valuation of beta cyclodextrin complexes of new drugs				
10.Preparation of par	racetamol tablets and comparison with marketed products				



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COURSE STRUCTURE & SYLLABI

Course Code 21S03201

MODERN PHARMACEUTICS - II

Semester

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Course Objectives:

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

Course Outcomes (CO): Student will be able to

Students will understand the planning of pilot plant techniques used for all pharmaceutical dosage forms such as tablets, capsules, parenterals, aerosols, cosmetics and neutraceuticals

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.

UNIT - 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 labelling & O.C. of anti-ageing products, sun screen lotion and fairness creams.

b. Nutraceuticals:

1. Introduction, source, manufacture and analysis of glucosamine & 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.



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COURSE STRUCTURE & SYLLABI

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.
- 6. Industrial Pharmacy Selected Topics, CVS Subramanyam and J Thimmasetty, Vallabha Prakashan Delhi 2013



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Course Code	ADVANCED DRUG DELIVERY SYSTEMS	L	T	P	C
21803202	ADVANCED DRUG DEELVERT STSTEMS	4	0	0	4
	Semester		Ι	Ι	
Course Objectives:					
The students shall ar	pply the pharmacokinetic and pharmacodynamic principles in the	e de	sign (of	
CDDS. They also a	pply the design, evaluation and applications related to oral, p	aren	teral.		
Transdermal, implan	its, bio adhesives 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 s	yste	ms o	f abo	ove
drug delivery system	ns with relevant applications.	2			
01111 - 1					
Fundamentals of con	ntrolled drug delivery systems, pharmacokinetic and pharmaco	dyna	amic	basis	s of
controlled drug deliv	very. Design, fabrication, evaluation and applications of the foll	owii	ng cc	ontrol	lled
releasing systems					
a. Controlled release	oral drug delivery systems				
b. Parenteral control	led release drug delivery systems				
UNIT - II					
Design, fabrication,	evaluation and applications of the following				
a. Implantable Thera	peutic systems				
b. Transdermal deliv	ery systems				
c. Ocular and Intraut	erine delivery systems				
d. Vaccine delivery	7: Delivery systems used to promote uptake, absorption enhancement	ncer	s, or	al	
immunization, control	olled release microparticles form vaccine development				
UNIT - III					
Biochemical and mo	lecular biology approaches to controlled drug delivery of				
a. Bioadhesive drug	delivery systems				
b. Nasal drug deliver	rv systems				
c. Drug delivery to C	Colon				
UNIT – IV					
Biochemical and mo	lecular biology approaches to control drug delivery of				
a. Liposomes					
b. Niosomes					
c. Microspheres					
d. Nanoparticles					
e. Resealed erythroc	vtes				
UNIT – V					
Drug targeting to par	rticular organs				
a. Delivery to lungs					
b. Delivery to the bra	ain and problems involved				
c. Drug targeting in I	neoplasams				



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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.
- 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 21S03203	INDUSTRIAL PHARMACY	L 4	T 0	P 0	C 4
	Semester	-	Ī	[
Correct Objections					
Course Objectives:					
The students shall I	earn the theory of unit operations, machinery, materials of c	bout		ons,	
quantication of equ	ciples of CMP. TOM and offluent analysis and specification	bout	hov.		
understand the requ	latory basis for the validation of analytical methods related	15. 1 to se	ney a alida	1150	
sterile and liquid do	sage forms	10 50	Jus)	
Course Outcomes (CO: Student will be able to				
The students will	explain the machinery involved in milling, mixing, filtrat	ion,	dryi	ng a	ind
packing material co	onstructions used in the production of pharmaceutical mate	erials	s. Ťh	ey a	lso
learn salient feature	els of GMP, TQM applicable in industry. They also underst	tand	the e	efflu	ent
treatments and pre	vent the pollution. They also should evaluate the validation	on o	f		
analytical methods	and processes				
UNIT - I					
Pharmaceutical up	nit operations: A detailed study involving machinery and the	heor	y of		
Pharmaceutical uni	t operations like milling, mixing, filtration, and drying.				
UNIT - II					
a. Materials of con	struction of pharmaceutical equipment and packaging materia	als: S	Study	of	the
principles, produc	tion techniques in the large scale production of tablets, capsules, su	usper	nsion	s, l1q	uid
b Qualification of e	opitnalmic products and sterile products. (IO, OO, PO)				
UNIT - III					
Production mana	agement: Production organization, objectives and po	licie	s of	go	bod
manufacturing pra-	ctices, layout of buildings, services, equipments and the	ir m	nainte	enan	ce,
material manageme	ent, handling and transportation, inventory management and	d cor	ntrol,		
production and pla	nning control, Sales forecasting, budget and cost control	, inc	dustr	ial a	ınd
personal relationshi	ip. Total Quality Management (TQM)				
UNIT - IV					
Effluent Testing an	nd Treatment: Effluent analysis, specifications and preven	tive	mea	sure	S
water of pollution,	solid pollution, air pollution and sound pollution.				
UNIT - V			• • •		
Validation: Regula	atory basis, validation of analytical methods, and process, in	n sol	id do	osag	e
Torms, sterile produ	icts, and liquid dosage forms.				
1 The Theory and	Practice of industrial Pharmacy by Leon Lachman, Herber	tΔI	iehe	rma	n
2. Good Manufactu	ring Practice for Dharmacouticals by Sidnay H. willig	ι Λ.Ι		/ina	,11.
2. Olou Manufactu	Ining Flactice for Finalitaceuticals by Stuney II. while.				
5. Pharmaceutical I	Tocess valuation by Robert A. Nash, Alfred H. wachter.				
4. Modern Pharmac	ceutics by Gilbert S. Banker and Christopher T. Rhodes.				
5. Pharmaceutical p	production management, C.V.S. Subrahmanyam, Vallabh Pr	akas	sh.		



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Reference Books:

- 1. Unit operations of Chemical Engineering by Warren L. McCabe, Julian C. Smith, Peter Harriott.
- 2. Remington's Science and Practice of Pharmacy by A. Gennaro.
- 3. Bentley's Text book of Pharmaceutics by EA Rawlins.
- CGMP, H.P.P. Sharma



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Course Code	Course Code NANO DRUG DELIVERY SYSTEMS	L	T	P	C
21803204	Somoston	4	0	0 T	4
	Semester		1	1	
Course Objectives:					
To develop expertise properties to the fail availability for targe	e regarding suitability and evaluation of nanomaterials, able to brication of nanopharmaceuticals, evaluate the intensity of d ting and controlled delivery.	appl osag	y the	e rms a	ınd
Course Outcomes (CO): Student will be able to				
The students should with appropriate tech	be able to select the right kind of materials, able to develop n nologies, evaluate the product related test and for identified dis	ano ease	form s	ulatio	ons
UNIT - I					
Introduction to Na	notechnology				
a. Definition of nano	technology				
b. History of nanoted	chnology				
c. Unique properties	and classification of nanomaterials				
d. Role of size and si	ze distribution of nanoparticles properties.				
e. Marketed formula	tions based on nanotechnology and science behind them				
UNIT - II					
Synthesis of Nanon	naterials				
Physical, chemical a	nd biological Methods				
Methods for synthes	is of				
Gold nanopa	urticles				
• Magnetic na	noparticles				
Polymeric n	anonarticles				
• Folymene na	mbly structures such as linesomes. Niesomes, transferasome	a m	icoll	20	
	and nanoomulsions	s, m	ICCII	-5,	
aquasomes a	ind nanoemulsions				
UNIT - III					
Biomedical applicat	tions of Nanotechnology				
a. Nanotechnology p	roducts used for in vitro diagnostics				
b. Improvements to 1	medical or molecular imaging using nanotechnology				
c. Targeted nanomat	erials for diagnostic and therapeutic purpose				
d.					
UNIT - IV					
Design of nanomate cancer therapy and c	erials for drug delivery, pulmonary and nasal drug delivery, r ardiovascular diseases. Localized drug delivery systems.	nano	mate	rials	for
UNIT - V					
Characterization inc separation, stability,	luding the principles, size reduction, analysis of nanoparticles methods of analysis regarding integrity and release of drugs	s, siz	ze, P	DI, s	ize
Defenence Deeler					



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- 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 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.



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Course Code		L	Т	Р	С
21803205	MUDEKN PHARMACEUTICS – II LAB	0	0	6	3
	Semester				
List of Experiments	S:				
1. Preparation of mo	uth washes				
2. Preparation and ev	valuation of cold creams and vanishing creams				
3. Preparation and ev	valuation of calamine lotion				
4. Preparation and ev	valuation of foundation creams and cleansing creams				
5. Preparation of ant	iseptic cream (turmeric)				
6. Preparation and ev	valuation Film coated tablets				
7. Preparation and ev	valuation Floating tablets				
8. Preparation and ev	valuation Fast dissolving tablets				
9. Preparation and ev	valuation Chewable tablets				
10. Effect of surfacta	ant in <i>in-vitro</i> drug release				
11. Preparation of or	al rehydration solution (ORS)				
12. Preparation and e	evaluation of calcium carbonate tablets				



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COURSE STRUCTURE & SYLLABI

Course Code	ADVANCED DRUG DELIVERY SYSTEMS LAB		L	Т	Р	С
21S03206	ADVANCED DRUG DELIVERY	SISIENIS LAD	0	0	6	3
Pre-requisite		Semester				
List of Experiments	5:					
1. Study on diffusion	of drugs through various polymeric memb	ranes (2 experiments	5)			
2. Formulation and e	valuation of sustained release oral matrix ta	ablet (2 experiments)				
3. Formulation and e	valuation of sustained release oral reservoir	system (2 experime	nts)			
	1					

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)



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Course Code	RESEARCH METHODOLOGY AND	L	Т	Р	С
21DRM101	INTELLECTUAL PROPERTY RIGHTS	4	0	0	4
	Semester		I	Π	
Course Objectives:					
To understar	nd the research problem				
• To know the	literature studies, plagiarism and ethics				
• To get the ki	nowledge about technical writing				
• To analyze t	he nature of intellectual property rights and new developments				
 To know the 	patent rights				
Course Outcomes (CO): Student will be able to				
At the end of this co	urse, students will be able to				
Understand	research problem formulation.				
Analyze rese	earch related information				
Follow research	rch ethics				
• Understand	that today's world is controlled by Computer, Information	Tec	hnolo	ogy,	but
tomorrow w	orld will be ruled by ideas, concept, and creativity.				
Understandi	ng that when IPR would take such important place in growth	of i	ndivi	duals	s &
nation, it is i	needless to emphasis the need of information about Intellectual	Prop	berty	Righ	t to
be promoted	among students in general & engineering in particular.				
• Understand	that IPR protection provides an incentive to inventors for furth	er r	esear	ch w	ork
and investm	ent in R & D, which leads to creation of new and better prod	lucts	, and	in ti	urn
brings about	, economic growth and social benefits.				
UNIT - I					
Meaning of research	problem, Sources of research problem, Criteria Characteristics of	of a	good	resea	rch
problem, Errors in	selecting a research problem, Scope and objectives of re-	esear	ch p	roble	em.
Approaches of invest	tigation of solutions for research problem, data collection, anal	ysis	,		
interpretation, Neces	sary instrumentations				
UNIT - II					
Effective literature st	tudies approaches, analysis, Plagiarism, Research ethics				
UNIT - III					
Effective technical w	riting, how to write report, Paper Developing a Research Propos	sal, l	Form	at of	
research proposal, a	presentation and assessment by a review committee				
UNIT - IV					
Nature of Intellectua	al Property: Patents, Designs, Trade and Copyright. Process of F	'ater	nting	and	
Development: techno	ological research, innovation, patenting, development. Internation	onal	Scen	ario:	
International coopera	ation on Intellectual Property. Procedure for grants of patents, Pa	itent	ing u	nder	
INIT V					
Dotont Dichter Corre	of Detent Diahte, Licensing and transfer of technology Deter	. :f	-	tion	I
databasas Casaran	or ratent kignts. Licensing and transfer of technology. Patent	f IIII f D-	orma	Cuot	and
Now dovelopments	in IDD, IDD of Diological Systems, Computer Software at 7	n ra Fradi	ition	Syste	JIII.
knowledge Case Stu	dies IPR and IITs	[I au	niona	.1	
Reference Rooks	uios, 11 17 uitu 11 1 5.				



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

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



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COURSE STRUCTURE & SYLLABI

AUDIT COURSE-I



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Course Code	ENGLISH FOR RESEARCH PAPER WRITING	L	Т	P	С
21DAC101a		2	0	0	0
	Semester			Ι	
Course Objectiv	ves: This course will enable students:				
• Understa	nd the essentials of writing skills and their level of readability				
 Learn ab 	out what to write in each section				
• Ensure q	ualitative presentation with linguistic accuracy				
Course Outcom	es (CO): Student will be able to				
• Understa	nd 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		ectur	e Hrs	s:10	
10verview of a up Long Sentence	Research Paper- Planning and Preparation- Word Order- Useful I es-Structuring Paragraphs and Sentences-Being Concise and Remo	Phrase oving	es - E Redu	Break unda	ing ncy
-Avoiding Ambig	guity				
UNIT - II		ectur	e Hrs	s:10	
Essential Compo Highlight Finding	onents of a Research Paper- Abstracts- Building Hypothesis-Res gs- Hedging and Criticizing, Paraphrasing and Plagiarism, Cauteri	earch zatio	n Pro n	blem	i -
UNIT - III		ectur	e Hrs	s:10	
Introducing Revi Conclusions-Rec	ew of the Literature – Methodology - Analysis of the Data-Find ommendations.	ings -	Dis	cussi	on-
UNIT - IV		Le	cture	Hrs:	9
Key skills needed	for writing a Title, Abstract, and Introduction				
UNIT - V		Le	cture	Hrs:	9
Appropriate lang Conclusions	uage to formulate Methodology, incorporate Results, put forth Ar	gume	nts a	nd dı	aw
Suggested Read	ing				
 Goldbort Model C Day R (2 Highman Uichman 	R (2006) Writing for Science, Yale University Press (available or urriculum of Engineering & Technology PG Courses [Volume-I] 2006) How to Write and Publish a Scientific Paper, Cambridge Uni n N (1998), Handbook of Writing for the Mathematical Sciences, S	n Goo versi IAM	gle E ty Pr	Books ess	;)
4. Adrian V Heidelbe	Vallwork, English for Writing Research Papers, Springer New Yo rg London, 2011	rk Do	rdrec	cht	



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COURSE STRUCTURE & SYLLABI

Course Code			L	Т	Р	С
21DAC101b	DISASTER MANAC	EMENT	2	0	0	0
		Semester]	[
Course Object	ves: This course will enable students					
• Learn to and hur	demonstrate critical understand anitarian response.	ing of key concepts in	n disast	ter risk	reducti	on
Critical Multipl	y evaluatedisasterriskreduction and h perspectives.	umanitarian response po	olicy and	d practic	e from	
• Develoy of disas	anunderstandingofstandardsofhuman ers and conflict situations	itarianresponseandpract	icalrele	vanceins	specific	types
Critical	yunderstandthestrengthsandweaknes	sesofdisastermanagemen	tapproa	ches,pla	nninga	nd
program	iming in different countries, particula	riy their nome country o	r the co	untries t	ney wo	rk in
Introduction:						
Disaster Defin	tion FactorsandSignificance:Difference	eBetweenHazardandDis	aster·Na	turaland	1	
Manmade Dise	sters: Difference Nature Types and	Magnitude	15101,144	luraiano	L	
Disaster Pron	A reas in India.	Wagintude.				
Study of Seisn	ic Zones: Areas Prone to Floods and	Droughts Landslides a	nd Aval	anches	Areas	Prone
to Cyclonic a	d Coastal Hazards with Special R	eference to Tsunami. P	lost- Di	saster I	Diseases	s and
Enidemics	te coastal mazares with Special K	cicicice to Tsunann, I	031- D1	saster 1	J15Cases	s and
Benercussion	of Disastars and Hazards					
Economic Da	aga Loss of Human and Animal	Life Destruction of Fe	osystan	a Notur	ol Dice	ostara
Economic Da	lage, Loss of Human and Amman	DroughtsondEngings I	ondelid	n. Natur	A volor	isters.
Man mada dis	ster: Nuclear Deaster Moltdown Ind	ustrial Assidants Oil Sl	anusnu	es allu	Avaiai	when of
Disease and E	idemica, War and Conflicts	usulai Accidentis, Oli Sh	icks and	i spins,	Outblea	1K5 01
$\frac{\text{UNII} - \text{III}}{\text{D}^2}$						
Disaster Prep	redness and Management:			1		
Preparedness:	Monitoring of Phenomena Irigge	ering ADIsasteror Haz	ard; E	valuatio	n of I	KISK:
Application of	Remote Sensing, Data from Mete	orological and Other	Agenci	es, Mec	na Re	ports:
Governmental	and Community Preparedness.					
Risk Assessme	nt Disaster Risk:	~			. ~.	
Concept and	Elements, Disaster Risk Reduction	n, Global and Nationa	al Disa	ster Ris	sk Situ	ation.
TechniquesofF	iskAssessment,GlobalCo-Operationir	RiskAssessmentand Wa	rning, F	eople's	Particip	pation
in Risk Assess	nent. Strategies for Survival.					
UNIT - V						
Disaster Mitig	ation:					
Meaning,Conc	ptandStrategiesofDisasterMitigation,	EmergingTrendsInMitig	ation.St	ructural		
Mitigationand	Non-Structural Mitigation, Programs	of Disaster Mitigation in	India.			

Suggested Reading



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- 1. R.Nishith, SinghAK, "DisasterManagementinIndia:Perspectives, issues and strategies
- "New Royal book Company..Sahni,PardeepEt.Al.(Eds.),"DisasterMitigationExperiencesAndReflections",PrenticeHa Il OfIndia, New Delhi.
- 3. GoelS.L.,DisasterAdministrationAndManagementTextAndCaseStudies",Deep&Deep Publication Pvt. Ltd., New Delhi



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Course Code	SANSKI	RITFOR TECHNICAL KNOWLEDG	E	L	Т	P	С
21DAC101c				2	0	0	0
		Sem	nester		ـــــــــــــــــــــــــــــــــــــ	I	
Course Objecti	ves: This cour	se will enable students:					
• To get a	working know	vledge in illustrious Sanskrit, the scientif	ic lang	uage in	the wo	rld	
Learnin	g of Sanskrit t	o improve brain functioning					
• Learnin	gofSanskrittod	evelopthelogicinmathematics,science&o	thersub	ojects en	nhancin	g the	
memory	/ power						
• The eng	gineering schol	ars equipped with Sanskrit will be able to	o explo	re the h	uge		
Knowle	dge from anci	entliterature					
Course Outcom	nes (CO): Stud	lent will be able to					
Underst	anding basic S	anskrit language					
Ancient	Sanskrit litera	ture about science &technology can be u	ndersto	bod			
Being a	logical langua	ge will help to develop logic in students					
UNIT - I							
Alphabets in Sa	anskrit,						
UNIT - II							
Past/Present/Fut	ure Tense, Sin	ple Sentences					
UNIT - III							
Order, Introduct	tion of roots						
UNIT - IV							
Technical infor	rmation about S	Sanskrit Literature					
UNIT - V							
Technical conc	epts of Engine	ering-Electrical, Mechanical, Architectur	e, Matł	nematic	S		
Suggested Read	ling						
1."Abhyaspust	akam" –Dr.V	ishwas, Sanskrit-Bharti Publication, I	New D	elhi			
2."Teach Your	rself Sanski	it" Prathama Deeksha- VempatiK	utumb	shastri	, Rasht	triyaSa	nskrit
Sansthanam, N	lew Delhi Pul	blication				-	
3."India's Glor	rious Scientif	cTradition" Suresh Soni, Ocean book	ts (P) I	Ltd.,Ne	w Dell	ni	



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AUDIT COURSE-II



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Course Code	PFI	DAGOGY STUDIES		L	Т	P	С
21DAC201a	1 121			2	0	0	0
		S	Semester]	I	
Course Objecti	ves: This course will e	enable students:					
Review	existingevidenceonthe	reviewtopictoinformprogramm	nedesignan	dpolicy	/ makin	g	
undertal	ten by the DfID, other	agencies and researchers.					
 Identify 	critical evidence gaps	to guide the development.					
Course Outcon	nes (CO): Student will	be able to					
Students will be	able to understand:						
• Whatpe countrie	lagogicalpracticesareb s?	eingusedbyteachersinformala	ndinforma	lclassro	oms in	develop	ing
• What is	the evidence on the ef	fectiveness of these pedagogic	cal practic	es, in w	hat		
conditio	ns, and with what pop	ulation of learners?		1	1		
Howcar material	teachereducation(curr s best support effective	culumandpracticum)andthescl	hoolcurric	uluman	d guida	nce	
UNIT - I							
terminology questions. Ove	Theories oflear rview of methodology	and Searching.	ation.Con	ceptual	framew	ork,Res	earch
UNIT - II							
Thematic ove classrooms in c	rview: Pedagogical leveloping countries.	practices are being used by Curriculum, Teacher education	teachers n.	in for	mal an	d inf	ormal
UNIT - III							
Evidence on th of included stu guidance mater evidence for es attitudes and be	eeffectivenessofpedag dies. How can teache ials best support effec fective pedagogical p liefs and Pedagogic s	ogicalpractices,Methodologyf er education (curriculumandpr tive pedagogy? Theory of cha ractices. Pedagogic theory an trategies.	Fortheinde racticum) inge. Stren id pedagog	pthstage andthes agth and gical ap	e:quality scho cu l nature oproach	y assess rriculur of th bo es. Tead	men t n and ody of chers'
UNIT - IV							
Professional d Support from t teacherandthec	evelopment: alignment ne head community.Curriculum	nt with classroom practices an andassessment,Barrierstolearn	d follow-u	ıp supp dresour	ort, Pee cesand I	er suppo large cla	rt, ass
SIZES							
			т	1 1	<i>.</i> .		
Kesearchgaps Curriculum and	andituredirections: assessment, Dissemin	Researchdesign,Contexts,Peda	agogy,Tea	cheredi	ication,		



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Suggested Reading

- $1. \quad Ackers J, Hardman F(2001) Classroom interaction in Kenyan primary schools, Compare,\\$
- 31 (2): 245-261.
- $2. \quad A grawal M (2004) Curricular reformins chools: The importance of evaluation, Journal of Marco 1999 (2004) Curricular reformins chools: The importance of evaluation of the second second$



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Course Code	ST	RESSMANAGEMEN'	Г ВҮ ҮОСА		T	P 0	C
21DAC2010			Semester	<u>_</u>	1	U T	U
			Semester				
Course Objecti	ives: This cour	se will enable students:					
To achi	eve overall hea	th of body and mind					
• To over	come stres	2					
Course Outcon	nes (CO): Stud	ent will be able to					
Develor	b healthy mind	in a healthy body thus	improving social health	also			
Improve	e efficiency						
IINIT - I	5						
Definitions of l	Fight parts of x	ag (Ashtanga)					
	Light parts of y	Jg.(Ashtanga)					
Vam and Nivay	m						
LINIT III	111.						
De`eend Den't							
Do sand Don t	sin me.	1	•\				
1) Aninsa, satya	astneya, orann	acharyaand aparigrana	1)				
LINIT - IV	sii,tapa,swauiiy	ay,isiiwaipiailiuliali					
A con and Pron	avom						
INIT - V	ayam						
i)Variousvogn	osecond theirb	nefitsformind & body					
i) Variousyogp	osesand then of	abrigues and its offects	Types of monoyiem				
	ding	chinques and its effects	s-Types orpranayani				
1 'Vogic Asana	s forGroupTar	ning_Part_I". Janardan	SwamiVogabhyasiMar	dal Na	mur		
2 "Rajavogaor c	conquering the	internal Nature" by Sw	ami Vivekananda Adva	ita Ashr	ama		
(Publication De	partment). Kol	kata		100 / 10111	umu		
(1 done don De							



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Course Code	PERSONALITY	DEVELOPMENT THRO	UGHLIFE	L	Т	Р	С
21DAC201c	ENI	LIGHTENMENTSKILLS	0000000	2	0	0	0
			Semester		Ι	I	
				1			
Course Objecti	ves: This course wi	ll enable students:					
To learn	to achieve the high	nest goal happily					
To beco	me a person with st	able mind, pleasing personal	ity and detern	nination			
To awal	ten wisdom in stude	ents					
Course Outcon	nes (CO): Student v	vill be able to					
Studyof	Shrimad-Bhagwad-	Geetawillhelpthestudentindev	velopinghispe	rsonality	yand ac	hieve	
the high	est goal in life						
• The per	son who has studied	l Geetawilllead the nation and	d mankind to	peace a	nd pros	perity	
Study of	Neetishatakam wi	ll help in developing versatile	e personality of	of stude	nts		
UNIT - I							
Neetisatakam-	Holistic developme	nt of personality					
Verses-19,	20,21,22(wisdom)						
Verses-29,	31,32(pride &herois	sm)					
Verses-26,	28,63,65(virtue)						
UNIT - II							
Neetisatakam-	Holistic developme	nt of personality					
Verses-52,	53,59(dont's)						
Verses-71,	73,75,78(do's)						
UNIT - III							
Approach to da	y to day work and c	luties.					
ShrimadBh	agwadGeeta:Chapt	er2-Verses41,47,48,					
Chapter 3-V	Verses13,21,27,35,C	hapter6-Verses5,13,17,23,35,	,				
Chapter18-	Verses45,46,48.						
UNIT - IV							
Statements of b	asic knowledge.			•			
ShrimadBh	agwadGeeta:Chapt	er2-Verses 56,62,68					
Chapter12	-Verses13,14,15,16	,17,18					
Personality	of Rolemodel. Shr	imad Bhagwad Geeta:					
UNIT - V							
Chapter2-V	Verses 17, Chapter 3-	Verses36,37,42,					
Chapter4-V	/erses18,38,39						
Chapter18-	- Verses37,38,63						
Suggested Read	ling						
1."SrimadBhaga	wadGita"bySwamiS	SwarupanandaAdvaitaAshran	n(Publication	Departm	ent),		
Kolkata							
2.Bhartrihari'sT	hree Satakam (Niti	-sringar-vairagya) by P.Gop	oinath, Rashti	riyaSans	skrit		
Sansthanam,	New Delhi.						



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OPEN ELECTIVE



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Course Code	BIOLOGICAL SCREENING METHODS	L	Т	Р	C
21SOE301d	(Elective)	3	0	0	3
	Semester		L	u	
Course Objectives:					
The students are goin	ng to study about various techniques for screening of drugs				
for various pharmac	ological activities and guide lines for handling animals and huma	an ai	nd an	imal	
ethics for screening	of drugs.				
Course Outcomes (CO): Student will be able to				
The expected outcor	nes are students will know how to handle animals and know				
about various techni	ques for screening of drugs for different pharmacological activit	ies, g	guide	lines	
and regulations for s	creening new drug molecules on animals.				
UNII - I	<u></u>				
Drug discovery proc	ess: Principles, techniques and strategies used in new drug disco	over	y. Hig	gh	
throughput screening	g, numan genomics, robotics and economics of drug discovery, F	kegu	latio	ns.	
molecular biology te	chniques	uoi	node	ls,	
UNIT - II					
Bioassavs: Basic prin	nciples of bioassays, official bioassays, experimental models and	1 sta	tistic	al	
designs employed in	biological standardization.				
UNIT - III					
Principles of toxicity	v evaluations, ED50, LD50 and TD values. International guideling	nes (ICH		
recommendations).			-		
Preclinical studies: C	General principles and procedures involved in acute, sub-acute, c	hror	nic,		
teratogenicity, mutag	genicity and carcinogenicity				
UNIT - IV					
Screening of differen	nt classes of drugs using micro-organisms. Vitamin and antibioti	c as	says.		
Screening methods 1	nvolved in toxins and pathogens.				
UNIT - V					
Enzymatic screening	ng methods: α-glucosidase, α- amylase, DNA polyme	rase	, nu	cleas	ses,
Lasparginase, lipase	s and peptidases.				
Reference Books:					
1. Basic and clinical	pharmacology by Bertram G. Katzung (International edition) la	nge	nedio	cal	
book / Mc Graw Hill	l, USA 2001 8th edition	1	4/-		
2. Pharmacology by	Rang H.P., Dale MM and Ritter JM., Churchill Livingston, Lond	ion,	4/e	10	
Graw Hill USA 200	1 10th edition	sam) IV	ic	
4 General and appli	d toxicology by B Ballantyne. T Marrs, P Turner (Eds) The Mc	Mill	an nr	ess	
Ltd. London.	a toxicology by Dibanantyne, Tittanis, Tittanier (Eas) The Ne		in pr	000	
5. Drug Discovery b	y Vogel's				
6. Drug Discovery a	nd evaluation – Pharmacological assays by H.Gerhard.Vogel, 2r	d ed	ition	,	
Springer verlag, Ber	lin, Heidelberg.				
7. Tutorial Pharmacy	(Vol I and II) by Cooper and Gunns.				



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	COURSE STRUCTURE & SYLLABI				
Course Code	PHARMACEUTICAL VALIDATION	L	Т	Р	C
21SOE301a	(Elective)	3	0	0	3
	Semester		I	Ι	
Course Objective	es:				
The main purpose	e of the subject is to understand about validation and how it can be a	appli	ed to)	
industry and thus	to improve the quality of the products. The subject covers the comp	olete	info	rmati	ion
about validation,	types, methodology and application				
Course Outcome	es (CO): Student will be able to				
Course Outcome	: Upon completion of the subject student shall be able to				
 Explain the second secon	ne aspect of validation				
Carryout	validation of manufacturing processes				
Apply the	knowledge of validation to instruments and equipments				
Validate t	he manufacturing facilities				
UNIT - I	Č.				
Introduction: Def	inition of Qualification and Validation. Advantage of Validation.	Str	eaml	ining	of
Oualification &	Validation process and Validation Master Plan. Qualification: U	ser	Reau	lirem	ient
Specification. De	sign Qualification. Factory Acceptance Test (FAT)/ Site Accepta	ince	Test	(SA	T).
Installation Quali	fication. Operational Qualification. Performance Qualification.	Re- (Duali	ficat	ion
(Maintaining stat	us -Calibration Preventive Maintenance. Change management).	Oua	lifica	ation	of
Manufacturing Ec	uipment, Qualification of Analytical Instruments and Laboratory e	quip	ment	ts.	
UNIT - II					
Qualification o	f analytical instruments: Electronic balance, pH met	er,	UV	-Visi	ble
spectrophotomete	r, FTIR, GC, HPLC, HPTLC	,			
Qualification of C	lassware: Volumetric flask, pipette, Measuring cylinder, beakers a	nd b	urette	e.	
UNIT - III					
Qualification of 1	aboratory equipments: Hardness tester, Friability test apparatus, ta	ap de	ensity	y test	ter,
Disintegration tes	ter, Dissolution test apparatus.	1	-		
Validation of Util	ity systems: Pharmaceutical water system & pure steam, HVAC systems	stem	•		
Compressed air an	nd nitrogen.		,		
UNIT - IV					
Cleaning Validati	on: Cleaning Validation - Cleaning Method development, Validation	on ai	nd va	lidat	ion
of analytical met	hod used in cleaning. Cleaning of Equipment. Cleaning of Facili	ties.	Clea	ning	in
place (CIP).				6	
UNIT - V					
Analytical metho	d validation: General principles, Validation of analytical meth	od a	as pe	er IC	H
guidelines and US	SP.		1		
Reference Books	•				



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- 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 Dekker).
- 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 Code	ENTREPRENEURSHIP MANA(GEMENT		<u>r</u>	<u>P</u>	C
21SOE301c	(Elective)	~	3 (0	0	3
		Semester		Π	I	
Course Objectiv						
This course is de	es:	eary to train the stu	donta	<u></u>		
entrepreneurship	management	sary to train the stu	uents (Л		
	$\frac{1}{2} \left(\frac{CO}{CO} \right) = \frac{1}{2} \left(\frac{1}{2} + \frac{1}{2} \right) = \frac{1}{2} \left(\frac$					
Course Outcome	es (CO): Student will be able to	-1-1				
On completion of	this course it is expected that students will be	able to:				
• The Role of e	nterprise in national and global economy					
• Dynamics of	abellanges of Crowth Strategies and Network	ng				
		ng				
UNII - I						
enterprise in nati policies and sche management.	onal and global economy. Types of enterprise mes for enterprise development. Institutional s	upport in enterprise	develo	pm	ent a	en inc
UNIT - II						
Entrepreneur: En	trepreneurial motivation - dynamics of motiva	tion. Entrepreneurial	comp	eter	ıcy –	-
Concepts. Develo	pping Entrepreneurial competencies - requirem	ents and understandi	ng the	pro	cess	o
entrepreneurship	development, self-awareness, interpersonal sl	kills, creativity, asse	rtivene	ess,		
achievement, fac	tors affecting entrepreneur role.					
UNIT - III						
Launching and C	rganizing an Enterprise: Environment scannin	ıg – Information, sou	irces, s	sche	emes	0
assistance, probl	ems. Enterprise selection, market assessment	, enterprise feasibili	ty stuc	ly,	SW	ΓO
Analysis. Resour	ce mobilization -finance, technology, raw mate	erial, site and manpo	wer. C	osti	ing a	ind
marketing manag	ement and quality control. Feedback, monitoring	ng and evaluation.	1			
UNIT - IV						
Growth Strategie	s and Networking: Performance appraisal and	assessment. Profita	bility a	and	cont	tro
measures, demai	nds and challenges. Need for diversification	n. Future Growth -	- Tecl	hnic	jues	0
expansion and d	iversification, vision strategies. Concept and	dynamics. Method	s, Joir	it v	entu	re,
coordination and						
		1 5 1114		DI	<u> </u>	
Preparing Projec	t Proposal to Start on New Enterprise Project	work – Feasibility	report;	Pla	annii	ng,
Defenence Rock						
1 Althour	M M D (1000): Entropyonourship for Warran	in India MIECDUD	Nor		hi	
1. Aknauri,	M. M. P. (1990): Entrepreneursnip for women	In India, NIESBUD	, new \sim		nı.	
2. HISTICH, J	N. D & DIUSH, C.G. (1990) The Women Entrep D and Datars M.D. (1995). Entrepresentitie	Storting David	$\alpha = 0$, 10 4	nant	υ.
5. HISTICH, J	A.D. and Feters, W.F. (1993). Enurepreneursmin a a New Enterprise Richard D. Inwin INC 1	J – Starting Develop	ing and	T		
Maradith	g a new Emerginse, Nichalu D., mwin, mv., U G.G. etal (1982): Practice of Entrepreneurshi	n II O Geneva				
5 Datel V	, (1987) : Women Entrepreneurshin – Develop	p, ILO, Oclieva.	ure A	hm₄	adah	he
EDII	c. (1767). Women Entrepreneutsmp – Develop		uis, A		Juan	uu
6. Arya kur Organiza	nar.(2012): Entrepreneurship- Creating and Leation Pearson	ading an Entrepreneu	rial			