

M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

SEMESTER - I

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

SEMESTER - II

S.No.	Course	Course Name	Н	Hours per		Hours per		Credits
	codes			T	P			
1.	21S03201	Modern Pharmaceutics-II	4	ı	-	4		
2.	21S03202	Advanced Drug Delivery system	4	1	-	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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SEMSTER - III

S.No.	Course	Course Name	Ho	Hours per		Credits
	codes			T	P	
1.	21DRM101	Research Methodology and Intellectual Property Right	4	-	-	4
2.	21SOE301a	Open Elective Biological Screening methods Pharmaceutical Validation Entrepreneurship Management	3	1	1	3
3.	21S03301	Teaching Practice/Assignment	-	1	4	2
4.	21S03302	Comprehensive viva voce	-	-	-	2
5.	21S03303	Research Work - I	-		24	12
		Total	7	-	32	23

SEMESTER - IV

S.No.	Course	Course Name	Hours per week			Hours per week		Credits
	codes		L	T	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	L	T	P	C	
21S01101	TECHNIQUES	4	0	0	4	
	Semester			Ī		
Course Objectives:						
This subject deals	with various advanced analytical instrumental techniques f	or i	denti	ficati	ion,	
characterization and	quantification of drugs. Instruments dealt are NMR, Mass s	spect	rom	eter,	IR,	
HPLC, GC etc.						
Course Outcomes (CO): Student will be able to					
After completion of	f course student is able to know about chemicals and excip	ient	S.			
 The analysis 	of various drugs in single and combination dosage forms					
Theoretical a	and practical skills of the instruments					
UNIT - I						
UV-Visible spectros	copy: Introduction, Theory, Laws, Instrumentation associated	with	ı UV	-Visi	ible	
spectroscopy, Choic	e of solvents and solvent effect and Applications of UV-Visil	ole s	pect	rosco	ру,	
Difference/ Derivativ	ve spectroscopy.					
UNIT - II						
1	heory, Modes of Molecular vibrations, Sample handling, In					
	rier -Transform IR Spectrometer, Factors affecting vibrational	free	quen	cies a	and	
	pectroscopy, Data Interpretation.					
UNIT - III						
	Quantum numbers and their role in NMR, Principle, Instrum					
	R, Relaxation process, NMR signals in various compounds,					
Factors influencing chemical shift, Spin-Spin coupling, Coupling constant, Nuclear magnetic double						
resonance, Brief outline of principles of FT-NMR and 13C NMR. Applications of NM						
spectroscopy.						

UNIT - IV

Mass Spectroscopy: Principle, Theory, Instrumentation of Mass Spectroscopy, Different types of ionization like electron impact, chemical, field, FAB and MALDI, APCI, ESI, APPI Analyzers of Quadrupole and Time of Flight, Mass fragmentation and its rules, Meta stable ions, Isotopic peaks and Applications of Mass spectroscopy.

UNIT - V

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 F

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

- 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 Wiley & Sons, 2004.



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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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Course Code	ADVANCED DIVERCAL DILADMA CEUTICE	L	T	P	C
21S03101	ADVANCED PHYSICAL PHARMACEUTICS	4	0	0	4
	Semester			I	
Course Objectives:					
The students shall kr	now about particle science, polymer science and its use in pharm	nace	utica	ıl dos	age
forms. They also kr	now the compression and consolidation parameters for powder	ers a	and g	granu	les.
Students also know	about the rheology, disperse systems, dissolution and solubilit	у ра	aram	eters	for
dosage forms.					
	CO): Student will be able to				
	ow particle size analysis method, solid dispersion, physics of			_	
	s applications, student will also know the stability calcula				
	elerated stability studies. They also know the rheology, absorb				
	lid dosage forms. They also know the factors affecting the	dis	solu	tion	and
-	o invitro/invivo correlations.				
UNIT - I					
	Classification, properties and characterization of polymers, I				
	ate, preparation of polymer solution, application of polymers is				
	nism of biodegradation of biodegradable polymers including			led d	rug
	acoadhesive, Hydrodynamically balanced and Transdermal Syst	ems			
UNIT - II					
	ompression: Basic principles of interactions, compression and				
	onsolidation under high loads, effect of friction, distributi				
	volume relationships, Heckel plots, compaction profiles, ene				
	ement of compression with strain gauges, compression pressure	-QA	para	ımete	rs.
UNIT - III					
	stability: Stability calculations, rate equations, complex order				
_	strategy of stability testing, method of stabilization, method				
	losage forms, temperature and humidity control, physical sta	abili	ty te	sting	; of
	acts. Photodecomposition, Method, solid state decomposition.				
UNIT - IV					
	ation, instrumentation, rheological properties of disperse system	s an	d sei	nisol	ids.
Oscillatory testing, C				. •	
	API and excipients: Differential Scanning Calorimetry: F		aple,	ther	mal
	es, disadvantages, instrumentation, applications and interpretation		1		
	on methods: Origin of x-rays, principle, advantages,	ď	isadv	/anta	ges,
	lications and interpretations.				
UNIT - V	shilter, Colubility and colubilization of non-leatening and 11111	-c+:	on 1-	, 41	
	ability: Solubility and solubilization of nonelectrolytes, solubility		•	•	
of surfactants, cos	olvents, complexation, drug derivatization and solid state	ie i	namj	pulat	ion,

(Peppas Model) and dissolution equipment **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

Mechanisms of Drug release - dissolution, diffusion (Matrix and Reservoir) and swelling controlled



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Delhi – 2013	
Reference Books:	
1. Dispersive systems I, II, and III	
2. Robinson. Controlled Drug Delivery Systems	



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Course Code		L	T	P	С
21S03102	MODERN PHARMACEUTICS – I	4	0	0	4
	Semester			Ī	
Course Objectiv	es:				
	ow the preformulation studies, methodology, different excipien				
	d their evaluation with references to production technologies. T		stude	nts a	ılso
	ation techniques and their applications in pharmaceutical industries				
	es (CO): Student will be able to				
	plain the preformulation parameters, apply ICH guidelines and eva-				
	tibility. Students also explain about formulation and development,				
	rs, capsules, micro-encapsules and coating techniques. They also le	arn a	and a	pply	the
	in different formulations.				
UNIT - I					
Preformulation	studies: Goals of Preformulation, preformulation parameters,	Poly	mor	phs a	and
Amorphous form	as, selection of drugs- solubility, partition coefficient, salt forms	, hu	midi	ty, so	olid
state properties,	Particle Size Analysis (Laser Diffraction and Dynamic Light S	catte	ering) dru	ıg -
	tibility, flow properties, format and content of reports of	pre	eforn	nulati	on,
preformulation st	ability studies (ICH)				
UNIT - II					
	velopment of solid dosage forms – I: New materials, excipients s				
	er disintegrants, etc, evaluation of functional properties of excipies	nts,	co-p	roces	sed
	ls of preparation and evaluation.				
UNIT - III					
	evelopment of solid dosage forms- II: Coating, coating m				
	blet technology for product development, computerization, inpr			ntrol	of
-	on development and manufacture of powder dosage forms for international	nal u	ise.		
	tion- types, methodology, problems encountered.				
UNIT - IV					
	velopment of soft and hard gelatin capsules: Introduction,				
	ufacture, filling equipment and filling operations, formulations, t				
	ances in capsule manufacture, machines, processing and co	ontr	ol i	nclud	ing
	spects, physical stability and packaging.				
UNIT - V					
	echniques in pharmaceutical formulation and processing				
	ameters, statistical design, response surface method, contour dia				
	actorial design, simplex methods, mixture designs, Placket Burh	an 1	meth	od, E	3ox
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 Lachman.
- 5. Modern Pharmaceutics by Gilbert S. Banker and Christopher T. Rhodes.
- 6. Pharmaceutical statistics by Bolton

Reference Books:

1. The Theory and Practice of industrial Pharmacy by Leon Lachman, Herbert A. Lieberman.



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- 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 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.

Clinical Pharmacokinetics: Altered kinetics in pregnancy, child birth, infants and geriatrics.



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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

- 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	I			

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		L	T	P	C
21S03104		0	0	6	3
	Semester			I	

List of Experiments

- 1. To carry out the preformulation studies of solid dosage forms.
- 2. To study the effect of compressional force on tablet disintegration time
- 3. To study the micromeritic properties of powders and granules
- 4. To study the effect of particle size on dissolution of tablets
- 5. To study the effect of binders on dissolution of tablets
- 6. To study pharmacokinetic models, to determine similarity factors
- 7. Accelerated stability testing of different tablets
- 8. Determination of first order, second order rate constants by acid and alkaline hydrolysis
- 9. Preparation and evaluation of beta cyclodextrin complexes of new drugs
- 10. Preparation of paracetamol tablets and comparison with marketed products



M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	MODERN BULL BALL CELUTICS II	L	T	P	C			
21S03201	MODERN PHARMACEUTICS - II	4	0	0	4			
	Semester	er II						
Course Objectiv	es:							
The students shal	l understand about the pilot plant and their scale up techniques for	man	ufact	uring	g of			
tablets capsules,	suspensions, emulsions and semisolids. The students also lear	rn tl	he fi	lling	of			
capsules, compre	ssion machines, sterilizers for formulation of parenterals and als	o un	ıders	tand	the			
	pellants, DPI, MDI and their quality control. The students also und	ersta	ınd a	bout	the			
cosmetics and nu	traceuticals.							
Course Outcome	es (CO): Student will be able to							
	lerstand the planning of pilot plant techniques used for all pharm	ıaceı	ıtical	dosa	age			
forms such as tab	lets, capsules, parenterals, aerosols, cosmetics and neutraceuticals							
UNIT - I								
Pilot plant scale-	up techniques used in pharmaceutical manufacturing							
a. Pilot plant: T	echnology transfer from R&D to pilot plant to pilot scale consider	erat	ions	of st	eps			
	nanufacture, layout design, facility, equipment selection of t	able	ts, c	apsu	les,			
•	lsions & semisolids.							
	Importance, Scale up process-size reduction, mixing, blendi	ng,	grai	nulati	on,			
	ting involved in tablets, capsules & liquid-liquid mixing.							
UNIT - II								
	velopment of parenteral dosage forms: Advances in materials	ano	d pro	oduct	ion			
techniques, filling	g machines, sterilizers, product layout.							
UNIT - III								
	Aerosols: Advances in propellants, metered dose inhaler designation	_	•	•				
inhalers, selection	inhalers, selection of containers and formulation aspects in aerosols formulation, manufacture and							
quality control.								

quality control.

UNIT - IV

a. Cosmetics: Formulation approaches, preparation & method of manufacturing labelling & Q.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.

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



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- 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	(
21S03202	ADVANCED DRUG DELIVERT STSTEMS	4	0	0	4
	Semester		I	Ι	
Course Objectives:					
	apply the pharmacokinetic and pharmacodynamic principles				
	apply the design, evaluation and applications related to	oral	, pa	rente	ra
	nts, bio adhesives and targeted drug delivery systems.				
	CO): Student will be able to				
	the drugs for CDDS design of the formulation fabrication of s	syste	ms c	of ab	ov
drug delivery systen	ns with relevant applications.				
UNIT - I					
	ntrolled drug delivery systems, pharmacokinetic and pharmaco	dyna	mic	hacie	
	very. Design, fabrication, evaluation and applications of the foll				
releasing systems	very. Design, faorication, evaluation and applications of the for	OWII	ig co	iiuoi	.IC
	e oral drug delivery systems				
	led release drug delivery systems				
UNIT - II	led release drug derivery systems				_
	evaluation and applications of the following				
a. Implantable Thera					
b. Transdermal deliv					
	terine delivery systems				
	ry: Delivery systems used to promote uptake, absorption	enh	ance	re (ar.
	olled release microparticles form vaccine development	CIIII	ance	15, (л
UNIT - III	oned release interopartieres form vacetile de veropinent				_
	blecular biology approaches to controlled drug delivery of				_
a. Bioadhesive drug					
b. Nasal drug delive	• •				
c. Drug delivery to (
UNIT – IV	Colon				-
	blecular biology approaches to control drug delivery of				-
a. Liposomes	recular biology approaches to control drug derivery of				
b. Niosomes					
c. Microspheres					
d. Nanoparticles					
e. Resealed erythroc	vtes				
UNIT – V	<i>yees</i>				_
Drug targeting to pa	rticular organs				
a. Delivery to lungs	iticular organis				
a. Denvery to fullgs					

- b. Delivery to the brain and problems involved
- c. Drug targeting in neoplasams

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



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7. Oral Drug Delivery Technology, 2nd ed, by Aukunuru Jithan



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Course Code	INDUSTRIAL PHARMACY	L	T	P	C
21S03203		4	0	0	4
	Semester		I	1	
Course Objectives:					
	learn the theory of unit operations, machinery, materials of	cor	strii	ction	10
	sipments and its utility. The students shall also understand				13,
	ciples of GMP, TQM and effluent analysis and specification				0
	ulatory basis for the validation of analytical methods relate			•	O
sterile and liquid de	· · · · · · · · · · · · · · · · · · ·	u to	SOII	40,	
	CO): Student will be able to				
	explain the machinery involved in milling, mixing, filtrat	ion.	drv	ing a	and
	onstructions used in the production of pharmaceutical mate				
	re1s of GMP, TQM applicable in industry. They also			-	
	and prevent the pollution. They also should evaluate the				
analytical methods	· · · · · · · · · · · · · · · · · · ·				
UNIT - I	•				
	init operations: A detailed study involving machinery it operations like milling, mixing, filtration, and drying.	and	d the	eory	of
UNIT - II					
a. Materials of cons	struction of pharmaceutical equipment and packaging materia	als:	Study	y of	the
	ction techniques in the large scale production of tablets, capsu	ıles,	susp	ensio	ons,
	ticals, ophthalmic products and sterile products.				
	quipment (IQ, OQ, PQ)				
UNIT - III		1			
	agement: Production organization, objectives and po			_	
	ctices, layout of buildings, services, equipments and the				
_	nent, handling and transportation, inventory management				
1.	anning control, Sales forecasting, budget and cost control	l, ın	dusti	nal a	and
1	ip. Total Quality Management (TQM)				
UNIT - IV	nd Treatment, Effluent analysis anaifications and ana	t:-	70	000	#0 °
_	nd Treatment: Effluent analysis, specifications and preve	entiv	e m	easu	res
water of pollution,	solid pollution, air pollution and sound pollution.				
UNIT - V					

Validation: Regulatory basis, validation of analytical methods, and process, in solid dosage forms, sterile products, and liquid dosage forms.

Textbooks:

- 1. The Theory and Practice of industrial Pharmacy by Leon Lachman, Herbert A. Lieberman.
- 2. Good Manufacturing Practice for Pharmaceuticals by Sidney H. willig.
- 3. Pharmaceutical Process validation by Robert A. Nash, Alfred H. Wachter.
- 4. Modern Pharmaceutics by Gilbert S. Banker and Christopher T. Rhodes.
- 5. Pharmaceutical production management, C.V.S. Subrahmanyam, Vallabh Prakash.



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M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

- 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



M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	NAMO DRUG DEL MIDDA GAGRIDAG	LI	' I	•
21S03204	NANO DRUG DELIVERY SYSTEMS	4 (0)
	Semester	,	II	,
Caura Objectives				
Course Objectives:	se regarding suitability and evaluation of nanomaterials, al	ala to	onnly	, +1
	brication of nanopharmaceuticals, evaluate the intensity of d			
	ting and controlled delivery.	osage	OTTI	, ai
	CO): Student will be able to			
	be able to select the right kind of materials, able to develop n	ano fo	mula	ntio
	nnologies, evaluate the product related test and for identified dis		111010	
** *	morogres, evaluate the product related test and for identified dis	ı		
UNIT - I				
Introduction to Na				
a. Definition of nanc	••			
b. History of nanoted				
	and classification of nanomaterials			
	ze distribution of nanoparticles properties.			
	tions based on nanotechnology and science behind them	I		
UNIT - II				
Synthesis of Nanom				
	nd biological Methods			
Methods for synthes				
Gold nanopa				
Magnetic na	•			
Polymeric na				
• Self – asso	embly structures such as liposomes, Niosomes, transfera	somes,	mic	elle
•	and nanoemulsions	1		
UNIT - III				
	tions of Nanotechnology			
	roducts used for in vitro diagnostics			
_	medical or molecular imaging using nanotechnology			
	erials for diagnostic and therapeutic purpose			
UNIT - IV			4	I - C
	erials for drug delivery, pulmonary and nasal drug delivery, i	nanoma	teria	is f
cancer therapy and c	ardiovascular diseases. Localized drug delivery systems.			
IINIT - V				

UNIT - V

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

- 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)



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M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

- 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



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M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	MODEDN DITADMA CEUTICE ILLAD	L	T	P	C
21S03205	MODERN PHARMACEUTICS – II LAB 0	0	0	6	3
		I	Ι		

List of Experiments:

- 1. Preparation of mouth washes
- 2. Preparation and evaluation of cold creams and vanishing creams
- 3. Preparation and evaluation of calamine lotion
- 4. Preparation and evaluation of foundation creams and cleansing creams
- 5. Preparation of antiseptic cream (turmeric)
- 6. Preparation and evaluation Film coated tablets
- 7. Preparation and evaluation Floating tablets
- 8. Preparation and evaluation Fast dissolving tablets
- 9. Preparation and evaluation Chewable tablets
- 10. Effect of surfactant in in-vitro drug release
- 11. Preparation of oral rehydration solution (ORS)
- 12. Preparation and evaluation of calcium carbonate tablets



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M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

Course Code	ADVANCED DDIC DELIVEDY	CVCTEMCIAD	L	T	P	C
21S03206	ADVANCED DRUG DELIVERY SYSTEMS LAB			0	6	3
Pre-requisite]	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 PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	Course Code RESEARCH METHODOLOGY AND				C	
21DRM101	RM101 INTELLECTUAL PROPERTY RIGHTS					
	Semester					

Course Objectives:

- To understand the research problem
- To know the literature studies, plagiarism and ethics
- To get the knowledge about technical writing
- To analyze the 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 course, students will be able to

- Understand research problem formulation.
- Analyze research related information
- Follow research ethics
- Understand that today's world is controlled by Computer, Information Technology, but tomorrow world will be ruled by ideas, concept, and creativity.
- Understanding that when IPR would take such important place in growth of individuals & nation, it is needless to emphasis the need of information about Intellectual Property Right to be promoted among students in general & engineering in particular.
- Understand that IPR protection provides an incentive to inventors for further research work and investment in R & D, which leads to creation of new and better products, and in turn brings about, economic growth and social benefits.

UNIT - I

Meaning of research problem, Sources of research problem, Criteria Characteristics of a good research problem, Errors in selecting a research problem, Scope and objectives of research problem. Approaches of investigation of solutions for research problem, data collection, analysis, interpretation, Necessary instrumentations

UNIT - II

Effective literature studies approaches, analysis, Plagiarism, Research ethics

UNIT - III

Effective technical writing, how to write report, Paper Developing a Research Proposal, Format of research proposal, a presentation and assessment by a review committee

UNIT - IV

Nature of Intellectual Property: Patents, Designs, Trade and Copyright. Process of Patenting and Development: technological research, innovation, patenting, development. International Scenario: International cooperation on Intellectual Property. Procedure for grants of patents, Patenting under 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.

- 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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M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

AUDIT COURSE-I



M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	ENGLISH FOR RESEARCH PAPER WRITING	L	T	P	C
21DAC101a		2	0	0	0
	Semester			I	
Course Objectiv	es: 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				
	ualitative presentation with linguistic accuracy				
Course Outcome	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			e Hrs		
	Research Paper- Planning and Preparation- Word Order- Useful Pes-Structuring Paragraphs and Sentences-Being Concise and Remoguity				
UNIT - II	Le	ectur	e Hrs	::10	
	nents of a Research Paper- Abstracts- Building Hypothesis-Regs- Hedging and Criticizing, Paraphrasing and Plagiarism, Cauteriz			oble	n -
UNIT - III	Le	ectur	e Hrs	:10	
Introducing Revi Conclusions-Rec	ew of the Literature – Methodology - Analysis of the Data-Findi ommendations.	ngs	- Dis	cussi	on-
UNIT - IV		Le	cture	Hrs:	9
Key skills needed	for writing a Title, Abstract, and Introduction				
UNIT - V			cture		
Appropriate lang Conclusions	uage to formulate Methodology, incorporate Results, put forth Arg	gume	ents a	nd di	aw
Suggested Read	ing				
	R (2006) Writing for Science, Yale University Press (available on	Goo	gle F	Books	;)
	urriculum of Engineering & Technology PG Courses [Volume-I]				
	006) How to Write and Publish a Scientific Paper, Cambridge Univ			ess	
	N (1998), Handbook of Writing for the Mathematical Sciences, S	IAM			
Highman		l. D.	andana :	-h+	
	Vallwork, English for Writing Research Papers, Springer New Yorrg London, 2011	к D(лаге	ا111ز	



M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

Course Code	DICACRED MANAGEMENT		L	T	P	С
21DAC101b	DISASTER MANAGEMENT	DISASTER MANAGEMENT	2	0	0	0
		Semester	I			

Course Objectives: This course will enable students:

- Learn to demonstrate critical understanding of key concepts in disaster risk reduction and humanitarian response.
- Critically evaluatedisasterriskreduction and humanitarian response policy and practice from Multiple perspectives.
- Developanunderstandingofstandardsofhumanitarianresponseandpracticalrelevanceinspecific types of disasters and conflict situations
- Criticallyunderstandthestrengthsandweaknessesofdisastermanagementapproaches, planning and programming in different countries, particularly their home country or the countries they work in

UNIT - I

Introduction:

Disaster:Definition,FactorsandSignificance;DifferenceBetweenHazardandDisaster;Naturaland Manmade Disasters: Difference, Nature, Types and Magnitude.

Disaster Prone Areas in India:

Study of Seismic Zones; Areas Prone to Floods and Droughts, Landslides and Avalanches; Areas Prone to Cyclonic and Coastal Hazards with Special Reference to Tsunami; Post- Disaster Diseases and Epidemics

UNIT - II

Repercussions of Disasters and Hazards:

Economic Damage, Loss of Human and Animal Life, Destruction of Ecosystem. Natural Disasters: Earthquakes, Volcanisms, Cyclones, Tsunamis, Floods, Droughtsand Famines, Landslides and Avalanches, Man-made disaster: Nuclear Reactor Meltdown, Industrial Accidents, Oil Slicks and Spills, Outbreaks of Disease and Epidemics, War and Conflicts.

UNIT - III

Disaster Preparedness and Management:

Preparedness: Monitoring of Phenomena Triggering ADisasteror Hazard; Evaluation of Risk: Application of Remote Sensing, Data from Meteorological and Other Agencies, Media Reports: Governmental and Community Preparedness.

UNIT - IV

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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M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

- 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 PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

Course Code	SANSKRI	TFOR TECHNICAL KNOWLE	DGE	L	T	P	C
21DAC101c				2	0	0	0
		S	Semester			I	L
Course Objecti	ves: This course	will enable students:					
To get a	working knowl	edge in illustrious Sanskrit, the scie	ntific lang	uage in	the wo	rld	
 Learning 	g of Sanskrit to	improve brain functioning					
 Learning 	gofSanskrittode	velopthelogicinmathematics, science	&othersub	ojects e	nhancin	g the	
memory	power						
• The eng	ineering scholar	s equipped with Sanskrit will be abl	le to explo	re the l	nuge		
• Knowle	dge from ancier	tliterature					
Course Outcon	nes (CO): Stude	nt will be able to					
 Underst 	anding basic Sa	nskrit language					
		re about science &technology can b		ood			
	logical language	e will help to develop logic in studer	nts				
UNIT - I							
Alphabets in Sa	anskrit,						
UNIT - II							
Past/Present/Fut	ure Tense, Simp	le Sentences					
UNIT - III							
Order, Introduct	ion of roots						
UNIT - IV							
Technical infor	mation about Sa	nskrit Literature					
UNIT - V							
Technical conc	epts of Engineer	ing-Electrical, Mechanical, Architec	cture, Matl	nematic	S		
Suggested Read							
		shwas, Sanskrit-Bharti Publicatio					
2."Teach You:	rself Sanskri	t" Prathama Deeksha- Vempa	ıtiKutuml	oshastr	i, Rash	triyaSa	nskrit
Sansthanam, N							
3."India's Glor	rious Scientific	Tradition" Suresh Soni, Ocean be	ooks (P)	Ltd.,No	ew Dell	hi	



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M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

AUDIT COURSE-II



M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

Course Code 21DAC201a	PEDAGOGY STUDIES	L 2	T 0	P 0	0				
	Semester II								
Course Objecti	ves: This course will enable students:								
	Reviewexistingevidenceonthereviewtopictoinformprogrammedesignandpolicy making undertaken by the DfID, other agencies and researchers.								

Course Outcomes (CO): Student will be able to

Students will be able to understand:

- Whatpedagogical practices are being used byteachers in formal and informal class rooms in developing countries?
- What is the evidence on the effectiveness of these pedagogical practices, in what conditions, and with what population of learners?
- Howcanteachereducation(curriculumandpracticum)andtheschoolcurriculumand guidance materials best support effective pedagogy?

UNIT - I

Introduction and Methodology: Aims and rationale, Policy back ground, Conceptual frame work and terminology Theories oflearning, Curriculum, Teachereducation. Conceptual framework, Research questions. Overview of methodology and Searching.

UNIT - II

Identify critical evidence gaps to guide the development.

Thematic overview: Pedagogical practices are being used by teachers in formal and informal classrooms in developing countries. Curriculum, Teacher education.

UNIT - III

Evidence on theeffectivenessofpedagogical practices, Methodology for the indepth stage: quality assessment of included studies. How can teacher education (curriculum and practicum) and the scho curriculum and guidance materials best support effective pedagogy? Theory of change. Strength and nature of the body of evidence for effective pedagogical practices. Pedagogic theory and pedagogical approaches. Teachers' attitudes and beliefs and Pedagogic strategies.

UNIT - IV

Professional development: alignment with classroom practices and follow-up support, Peer support, Support from the head

teacher and the community. Curriculum and assessment, Barrier stolearning: limited resources and large class sizes

UNIT - V

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

Suggested Reading

- 1. Ackers J, Hardman F (2001) Class room 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.



M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

Course Code	CED		.	L	T	P	C
21DAC201b	STR	ESSMANAGEMENT BY YOGA	A	2	0	0	0
			Semester		I	I	
Course Objecti	ves: This course	e will enable students:					
To achie	eve overall healt	h of body and mind					
• To over	come stres						
Course Outcom	nes (CO): Stude	nt will be able to					
•	•	n a healthy body thus improving so	ocial health	also			
 Improve 	efficiency						
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	j	•					
Asan and Prana	ıyam						
UNIT - V							
i)Variousyogpo	sesand theirben	efitsformind &body					
ii)Regularizatio	onofbreathingtec	hniques and its effects-Types ofpr	anayam				
Suggested Read							
		ing-Part-I": Janardan SwamiYoga					
		e Internal Nature" by Swami	Vivekananda	a, Adv	aıta		
Ashrama (Public	cation Departme	ent), Kolkata					



M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	PERSONALITY	DEVELOPMENT THRO	UGHLIFE	L	T	P	C
21DAC201c	ENLI	GHTENMENTSKILLS		2	0	0	0
			Semester		I	I	
Course Objecti	ves: This course will	enable students:					
 To learn 	to achieve the highe	st goal happily					
	_	ble mind, pleasing personal	lity and deterr	ninatior	1		
	en wisdom in studen						
	es (CO): Student wi						
•	•	eetawillhelpthestudentinde	evelopinghispe	ersonali	tyand ac	chieve	
•	est goal in life	G			,	•.	
_		Geetawilllead the nation an		_		perity	
	Neetishatakam will	help in developing versatil	e personality of	of stude	nts		
UNIT - I	(T. 1)	C 1'.					
	Holistic development	of personality					
	20,21,22(wisdom)						
	31,32(pride &heroisn	1)					
	28,63,65(virtue)		1				
UNIT - II							
	Holistic development	of personality					
Verses-52,	53,59(dont's)						
	73,75,78(do's)		1				
UNIT - III							
	y to day work and du						
	agwadGeeta:Chapter						
Chapter3-V	erses13,21,27,35,Ch	apter6-Verses5,13,17,23,35	5,				
	Verses45,46,48.						
UNIT - IV							
Statements of b	asic knowledge.						
ShrimadBh	agwadGeeta:Chapter	2-Verses 56,62,68					
Chapter 12	-Verses 13, 14, 15, 16, 1	7,18					
Personality	of Rolemodel. Shrin	nad Bhagwad Geeta:					
UNIT - V							
Chapter2-V	erses 17,Chapter3-V	erses36,37,42,					
Chapter4-V	erses18,38,39						
Chapter 18-	- Verses37,38,63						
Suggested Read							
	vadGita"bySwamiSv	warupananda Advaita Ashra	m(Publication	Departi	ment),		
Kolkata	0.1 077				1		
		sringar-vairagya) by P.Go	pınath, Rasht	rıyaSan	skrit		
Sansthanam,	New Deini.						



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M.PHARM. IN PHARMACEUTICS

COURSE STRUCTURE & SYLLABI

OPEN ELECTIVE



M.PHARM. IN PHARMACEUTICS **COURSE STRUCTURE & SYLLABI**

Course Code	BIOLOGICAL SCREENING METHODS	L	T	P	C
21SOE301d	(Elective)	3	0	0	3
	Semester		III		
Course Objectives:					
	ng to study about various techniques for screening of drugs				
for various pharmaco ethics for screening of	ological activities and guide lines for handling animals and huma	an ai	nd an	imal	
	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	iec	mide	lines	
	creening new drug molecules on animals.	ics, į	guiuc	iiiics	
UNIT - I	erecting new drug infriences on unimais.				
	ess: Principles, techniques and strategies used in new drug disco	Mort	, Ці,	rh	
	s, human genomics, robotics and economics of drug discovery, I				
0 1	al screening procedures, cell-line, patch—clamp technique, In-vi	_			
molecular biology te		1101	nouc	15,	
UNIT - II	cimiques				
	nciples of bioassays, official bioassays, experimental models and	d eta	tistic	91	
	biological standardization.	1 514	tistic	uı	
	0101081411 011111111111111111111				
UNIT - III	1 PD50 1 D50 1 TD 1 1 1 1		ICII		
	evaluations, ED50, LD50 and TD values, International guideling	ies (ICH		
recommendations).	Sanagal main ain less and muse advues investived in courte, sub-courte	haci			
	General principles and procedures involved in acute, sub-acute, or genicity and carcinogenicity	лиог	nc,		
UNIT - IV	genicity and careinogenicity				
	nt classes of drugs using micro-organisms. Vitamin and antibioti	0.00	COLIC		
•	nvolved in toxins and pathogens.	c as	says.		
Screening memous i	involved in toxins and pathogens.				
UNIT - V					
Enzymatic screening	ng methods: α-glucosidase, α- amylase, DNA polyme	rase	, nu	cleas	ses,
Lasparginase, lipases					
Reference Books:	• •				
1. Basic and clinical	pharmacology by Bertram G. Katzung (International edition) la	nge	medi	cal	
book / Mc Graw Hill	I, USA 2001 8th edition				
2. Pharmacology by	Rang H.P, Dale MM and Ritter JM., Churchill Livingston, Lond	lon,	4/e		
3. Goodman and Gil	man's The pharmacological basis of therapeutics (International	editi	on) N	Лс	
Graw Hill, USA 200	1 10th edition.				

- 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 PHARMACEUTICS

COURSE STRUCTURE & SYLLABI							
Course Code	PHARMACEUTICAL VALIDATION	L	T	P	C		
21SOE301a	(Elective)	3	0	0	3		
	Semester	III		•			
Course Objectiv	es:						
	e of the subject is to understand about validation and how it can be						
	to improve the quality of the products. The subject covers the comp	plete	info	rmat	ion		
about validation,	types, methodology and application						
Course Outcome	es (CO): Student will be able to						
Course Outcome	e: Upon completion of the subject student shall be able to						
 Explain t 	he aspect of validation						
Carryout	validation of manufacturing processes						
_	e knowledge of validation to instruments and equipments						
* * *	the manufacturing facilities						
UNIT - I							
Introduction: Det	Finition of Qualification and Validation, Advantage of Validation.	Str	eaml	ining	of		
	Validation process and Validation Master Plan. Qualification: U						
	esign Qualification, Factory Acceptance Test (FAT)/ Site Accepta						
	ification, Operational Qualification, Performance Qualification, I						
	tus -Calibration Preventive Maintenance, Change management),						
	quipment, Qualification of Analytical Instruments and Laboratory e	_					
UNIT - II							
Qualification o	f analytical instruments: Electronic balance, pH met	er,	UV	-Visi	ible		
spectrophotomete	er, FTIR, GC, HPLC, HPTLC						
	Glassware: Volumetric flask, pipette, Measuring cylinder, beakers a	nd b	urett	e.			
UNIT - III							
Qualification of	laboratory equipments: Hardness tester, Friability test apparatus, t	ap d	ensit	y tes	ter,		
_	ster, Dissolution test apparatus.	•		•	-		
Validation of Uti	lity systems: Pharmaceutical water system & pure steam, HVAC sy	sten	1,				
Compressed air a							
UNIT - IV							
Cleaning Validat	ion: Cleaning Validation - Cleaning Method development, Validation	on a	nd va	alidat	ion		
	hod used in cleaning. Cleaning of Equipment. Cleaning of Facili						
place (CIP)					•		

place (CIP).

UNIT - V

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

- 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.



M.PHARM. IN PHARMACEUTICS COURSE STRUCTURE & SYLLABI

Course Code	ENTREPRENEURSHIP MANAGEMENT		L	T	P	C
21SOE301c	(Elective)		3	0	0	3
		Semester	III			
Course Objective	s:					
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This course is designed to impart knowledge and skills necessary to train the students 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 development. Role of enterprise in national and global economy. Types of enterprise – Merits and Demerits. Government policies and schemes for enterprise development. Institutional support in enterprise development and management.

UNIT - II

Entrepreneur: Entrepreneurial motivation – dynamics of motivation. Entrepreneurial competency – Concepts. Developing Entrepreneurial competencies - requirements and understanding the process of entrepreneurship development, self-awareness, interpersonal skills, creativity, assertiveness, achievement, factors affecting entrepreneur role.

UNIT - III

Launching and Organizing an Enterprise: Environment scanning – Information, sources, schemes of assistance, problems. Enterprise selection, market assessment, enterprise feasibility study, SWOT Analysis. Resource mobilization -finance, technology, raw material, site and manpower. Costing and marketing management and quality control. Feedback, monitoring and evaluation.

UNIT - IV

Growth Strategies and Networking: Performance appraisal and assessment. Profitability and control measures, demands and challenges. Need for diversification. Future Growth — Techniques of expansion and diversification, vision strategies. Concept and dynamics. Methods, Joint venture, coordination and feasibility study.

UNIT - V

Preparing Project Proposal to Start on New Enterprise Project work – Feasibility report; Planning, 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