SAVITRIBAI PHULE PUNE UNIVERSITY

FACULTY OF SCIENCE AND TECHNOLOGY



COURSE STRUCTURE AND SYLLABUS

FINAL YEAR BACHELOR OF PHARMACY (B. Pharm.) 2019PATTERN (EFFECTIVE FROM ACADEMIC YEAR 2022 – 2023)

CHAPTER-I: REGULATIONS

1. Short Title and Commencement

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

2. Minimum qualification for admission

2.1 First year B. Pharm:

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

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

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

3. Duration of the program

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

4. Medium of instruction and examinations

Medium of instruction and examination shall be in English.

5. Working days in each semester:

Each semestershall consist of not less than 90 working days. The odd semesters shall be conducted from the month of June/July to November/December and the even semesters shall be conducted from December/January to May/June in every calendar year.

6. Attendance and progress

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

7. **Program/Course credit structure**

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

7.1. Credit assignment

7.1.1. Theory and Laboratory courses

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

7.2. Minimum credit requirements

The minimum credit points required for award of a B. Pharm. degree **is 211**. These credits are divided into Theory courses, Tutorials, Practical, Practice School and Projectover the duration of eight semesters. The credits are distributed semester-wise as shown in Table IX. Courses generally progress in sequences, building competencies and their positioning indicates certain academic maturity on the part of the learners. Learners are expected to follow the semester-wise schedule of courses given in the syllabus. The lateral entry students shall get 52 credit points transferred from their D. Pharm program. Such students shall take up additional remedial courses of 'Communication Skills' (Theory and Practical) and 'Computer Applications in Pharmacy' (Theory and Practical) equivalent to 3 and 4 credit points respectively, a total of 7 credit points to attain 59 credit points, the maximum of I and II semesters.

8. Academic work

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

9. Course of study

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

Course code	Name of the course	No.of Hours per week/Total no of hours	Tutorial	Credit points
BP101T	Human Anatomy and Physiology I– Theory	3/45	1	4
BP102T	Pharmaceutical Analysis I – Theory	3/45	1	4
BP103T	Pharmaceutics I – Theory	3/45	1	4
BP104T	Pharmaceutical Inorganic Chemistry – Theory	3/45	1	4
BP105T	Communication skills – Theory *	2/30	-	2
BP106RBT BP106RMT	Remedial Biology/ Remedial Mathematics – Theory*	2/30	-	D
BP107P	Human Anatomy and Physiology – Practical	4/60	-	2
BP108P	Pharmaceutical Analysis I – Practical	4/60	-	2
BP109P	Pharmaceutics I – Practical	4/60	-	2
BP110P	Pharmaceutical Inorganic Chemistry – Practical	4/60	-	2
BP111P	Communication skills – Practical*	2/30	-	1
BP112RBP	Remedial Biology – Practical*	2/30	-	D
	Total	32/34 ^{\$} /36 [#] /480/510 ^{\$} /540 [#]	4	27

Table-I: Course of study for semester I

Applicable ONLY for the students who have studied Mathematics / Physics / Chemistry at HSC and appearing for Remedial Biology (RB) course. However for Remedial biology and Mathematics no credits to be allotted only 50 % passing i.e D grade will be prerequisite.

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

* Non University Examination (NUE)

Course Code	Name of the course	No. of Hours per week/Total no of hours	Tutorial	Credit points			
BP201T	Human Anatomy and Physiology II – Theory	3/45	1	4			
BP202T	Pharmaceutical Organic Chemistry I – Theory	3/45	1	4			
BP203T	Biochemistry – Theory	3/45	1	4			
BP204T	Pathophysiology – Theory	3/45	1	4			
BP205T	Computer Applications in Pharmacy – Theory *	3/45	-	3			
BP206T	Environmental sciences – Theory *	3/45	-	3			
BP207P	Human Anatomy and Physiology II – Practical	4/60	-	2			
BP208P	Pharmaceutical Organic Chemistry I– Practical	4/60	-	2			
BP209P	Biochemistry – Practical	4/60	-	2			
BP210P	Computer Applications in Pharmacy – Practical*	4/60	-	1			
	Total 32/480 4 29						

Table-II: Course of study for semester II

* Non University Examination (NUE)

Table-III: Course of study for semester III

Course code	Name of the course	week/Total no of hours		Credit points
BP301T	Pharmaceutical Organic Chemistry II – Theory	3/45	1	4
BP302T	Physical Pharmaceutics I – Theory	3/45	1	4
BP303T	Pharmaceutical Microbiology – Theory	3/45	1	4
BP304T	Pharmaceutical Engineering – Theory	3/45	1	4
BP305P	Pharmaceutical Organic Chemistry II – Practical	4/60	-	2
BP306P	Physical Pharmaceutics I – Practical	4/60	-	2
BP307P	Pharmaceutical Microbiology – Practical	4/60	-	2
BP 308P	Pharmaceutical Engineering –Practical	4/60	-	2
	Total	28/420	4	24

Table-IV: Course of study for semester IV

Course code	Name of the course	No. of Hours per week/Total no of hours	Tutorial	Credit points
BP401T	Pharmaceutical Organic Chemistry III– Theory	3/45	1	4
BP402T	Medicinal Chemistry I – Theory	3/45	1	4
BP403T	Physical Pharmaceutics II – Theory	3/45	1	4
BP404T	Pharmacology I – Theory	3/45	1	4
BP405T	Pharmacognosy and Phytochemistry I– Theory	3/45	1	4
BP406P	Medicinal Chemistry I – Practical	4/60	-	2
BP407P	Physical Pharmaceutics II – Practical	4/60		2
BP408P	Pharmacology I – Practical	4/60	-	2
BP409P	Pharmacognosy and Phytochemistry I – Practical	4/60	-	2
	Total	31/465	5	28

Table-V: Course of study for semester V

Course code	Name of the courseNo. of Hours per week/Total no of hours		Tutorial	Credit points
BP501T	Medicinal Chemistry II – Theory	3/45	1	4
BP502T	Industrial Pharmacy-I– Theory	3/45	1	4
BP503T	Pharmacology II – Theory	3/45	1	4
BP504T	Pharmacognosy and Phytochemistry II– Theory	3/45	1	4
BP505T	Pharmaceutical Jurisprudence – Theory	3/45	1	4
BP506P	Industrial Pharmacy-I - Practical	4/60	-	2
BP507P	Pharmacology II – Practical	4/60	-	2
BP508P	Pharmacognosy and Phytochemistry II – Practical	4/60	-	2
	Total	27/405	5	26

Table-VI: Course of study for semester VI

Course code	Name of the course	No. of Hours per week/Total no of hours	Tutorial	Credit points
BP601T	Medicinal Chemistry III – Theory	3/45	1	4
BP602T	Pharmacology III – Theory	3/45	1	4
BP603T	Herbal Drug Technology – Theory	3/45	1	4
BP604T	Biopharmaceutics and Pharmacokinetics – Theory	3/45	1	4
BP605T	Pharmaceutical Biotechnology – Theory	3/45	1	4
BP606T	Quality Assurance – Theory	3/45	1	4
BP607P	Medicinal chemistry III – Practical	4/60	-	2
BP608P	Pharmacology III – Practical	4/60	-	2
BP609P	Herbal Drug Technology – Practical	4/60	-	2
	Total	30/450	6	30

Table - VII: Course of study for semester VII

Course code	Name of the course	No. of Hours per week/Total no of hours	Hours per week/Total Tutorial	
BP701T	Instrumental Methods of Analysis – Theory	3/45	1	4
BP702T	Industrial Pharmacy-II – Theory	ial Pharmacy-II – Theory 3/45 1		4
BP703T	Pharmacy Practice – Theory	3/45	1	4
BP704T	Novel Drug Delivery System – Theory	3/45	1	4
BP705P	Instrumental Methods of Analysis – Practical 4/60		-	2
BP706PS	Practice School* 12/180 -		6	
	Total	28/420	5	24

* Non University Examination (NUE)

Course code	Name of the course	No. of Hours per week/Total no of hours	Tutorial	Credi t points		
BP801T	Biostatistics and Research Methodology	3/45	1	4		
BP802T	Social and Preventive Pharmacy	ventive Pharmacy 3/45 1				
BP803ET	Pharma Marketing Management					
BP804ET	Pharmaceutical Regulatory Science					
BP805ET	Pharmacovigilance					
BP806ET	Quality Control and Standardizations of Herbals					
BP807ET	Computer Aided Drug Design					
BP808ET	Cell and Molecular Biology	3 + 3 =		4 + 4		
BP809ET	Cosmetic Science	6/90	1 + 1 = 2	8		
BP810ET	Experimental Pharmacology			0		
BP811ET	Advanced Instrumentation Techniques					
BP812ET	Dietary Suppliments and Nutraceuticals					
BP813PW	Project Work	12/180	-	6		
	Total	24/360	4	22		

Table - VIII:Course of study for semester VIII

Table-IX: Semester wise credits distribution

Semester	Credit Points
Ι	27
Ш	29
III	24
IV	28
V	26
VI	30
VII	24
VIII	22
Extracurricular/ Co curricular activities	01*
Total credit points for the program	211

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

10. Program Committee

- 1. The B. Pharm. program shall have a Program Committee constituted by the Head of the institution in consultation with all the Heads of the departments.
- 2. The composition of the Program Committee shall be as follows:

A senior teacher shall be the Chairperson; One Teacher from each department handling B.Pharm courses; and four student representatives of the program (one from each academic year), nominated by the Head of the institution.

- 3. Duties of the Program Committee:
 - i. Periodically reviewing the progress of the classes.
 - ii. Discussing the problems concerning curriculum, syllabus and the conduct of classes.
 - iii. Discussing with the course teachers on the nature and scope of assessment for the course and the same shall be announced to the students at the beginning of respective semesters.
 - iv. Communicating its recommendation to the Head of the institution on academic matters.
 - v. The Program Committee shall meet at least thrice in a semester preferably at the end of each Sessional exam (Internal Assessment) and before the end semester exam.

11. Examinations/Assessments

The scheme for internal assessment and end semester examinations is given in Table -X.

11.1 End semester examinations

The End Semester Examinations for each theory and practical course through semesters I to VIII shall be conducted by the university except for the subjects with asterix symbol (*) in table I and II for which examinations shall be conducted by the subject experts at college level and the marks/grades shall be submitted to the university.

G			Internal Assessment				semester xams	T ()
Course code	Name of the	Contin	Sessiona	l Exams				Total Marks
	course	uousM ode	Marks	Duration	Total	Marks	Duration	
BP101T	Human Anatomy and Physiology I– Theory	10	15	1 Hr	25	75	3 Hrs	100
BP102T	Pharmaceutical Analysis I – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP103T	Pharmaceutics I – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP104T	Pharmaceutical Inorganic Chemistry – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP105T	Communication skills – Theory *	5	10	1 Hr	15	35	1.5 Hrs	50
BP106R BT BP106R MT	Remedial Biology/ Mathematics – Theory*	5	10	1 Hr	15	35	1.5 Hrs	50
BP107P	Human Anatomy and Physiology – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP108P	Pharmaceutical Analysis I – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP109P	Pharmaceutics I – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP110P	Pharmaceutical Inorganic Chemistry – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP111P	Communication skills – Practical*	5	5	2 Hrs	10	15	2 Hrs	25
BP112R BP	Remedial Biology – Practical*	5	5	2 Hrs	10	15	2 Hrs	25
	Total	70/75 ^{\$} /8 0 [#]	115/125 ^{\$} / 130 [#]	23/24 ^{\$} /26 [#] Hrs	185/200 ^{\$} /210 [#]	490/525 ^{\$} / 540 [#]	31.5/33 ^{\$} / 35 [#] Hrs	675/72 5 ^{\$} / 750 [#]

Tables-X: Schemes for internal assessments and end sem exam semester wiseSem I

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

- ^{\$} Applicable ONLY for the students studied Physics / Chemistry / Botany / Zoology at HSC and appearing for Remedial Mathematics (RM)course.
- * Non University Examination(NUE)

Semester II

			Internal Assessment				End Semester Exams	
Course code	Name of the course	Continu	Sessional Exams		Tatal	Maalaa	Denting	Total Marks
		ous Mode	Marks	Duration	Total	Marks	Duration	
BP201T	Human Anatomy and Physiology II – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP202T	PharmaceuticalOr ganic Chemistry I – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP203T	Biochemistry – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP204T	Pathophysiology – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP205T	Computer Applications in Pharmacy – Theory*	10	15	1 Hr	25	50	2 Hrs	75
BP206T	Environmental sciences – Theory*	10	15	1 Hr	25	50	2 Hrs	75
BP207P	Human Anatomy and Physiology II –Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP208P	Pharmaceutical Organic Chemistry I– Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP209P	Biochemistry – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP210P	Computer Applications in Pharmacy – Practical*	5	5	2 Hrs	10	15	2 Hrs	25
	Total	80	125	20 Hrs	205	520	30 Hrs	725

* The subject experts at college level shall conductexaminations.

Semester III

		I	Internal Assessment				emester ams	
Course code	Name of the course	Continuous	Session	al Exams	T-4-1	Maalaa	D	Total Marks
		Mode	Marks	Duration	Total	Marks	Duration	
BP301T	PharmaceuticalOrganic Chemistry II – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP302T	PhysicalPharmaceuticsI —Theory	10	15	1 Hr	25	75	3 Hrs	100
BP303T	Pharmaceutical Microbiology – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP304T	Pharmaceutical Engineering – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP305P	PharmaceuticalOrganic Chemistry II – Practical	5	10	4 Hr	15	35	4 Hrs	50
BP306P	Physical Pharmaceutics I – Practical	5	10	4 Hr	15	35	4 Hrs	50
BP307P	Pharmaceutical Microbiology – Practical	5	10	4 Hr	15	35	4 Hrs	50
BP308P	Pharmaceutical Engineering – Practical	5	10	4 Hr	15	35	4 Hrs	50
	Total	60	100	20	160	440	28Hrs	600

Semester IV

		Internal Assessment				End S Ex		
Course code	Name of the course	Continuous	Session	al Exams	Total	Marks	Duration	Total Marks
		Mode	Marks	Duration	Totai	Marks		
BP401T	Pharmaceutical Organic Chemistry III– Theory	10	15	1 Hr	25	75	3 Hrs	100
BP402T	Medicinal Chemistry I – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP403T	Physical Pharmaceutics II – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP404T	Pharmacology I – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP405T	Pharmacognosy I – Theory	10	15	1 Hr	25	75	3 Hrs	100

BP406P	Medicinal Chemistry I – Practical	5	10	4 Hr	15	35	4 Hrs	50
BP407P	Physical Pharmaceutics II – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP408P	Pharmacology I – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP409P	Pharmacognosy I – Practical	5	10	4 Hrs	15	35	4 Hrs	50
	Total	70	115	21 Hrs	185	515	31 Hrs	700

Semester V

		Internal Assessment				End S Ex		
Course code	Name of the course	Continuous	Session	al Exams	T ()			Total Marks
		Mode	Marks	Duration	Total	Marks	Duration	
BP501T	Medicinal Chemistry II – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP502T	Industrial Pharmacy–I- Theory	10	15	1 Hr	25	75	3 Hrs	100
BP503T	Pharmacology II – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP504T	Pharmacognosy II – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP505T	Pharmaceutical Jurisprudence– Theory	10	15	1 Hr	25	75	3 Hrs	100
BP506P	Industrial Pharmacy–I – Practical	5	10	4 Hr	15	35	4 Hrs	50
BP507P	Pharmacology II – Practical	5	10	4 Hr	15	35	4 Hrs	50
BP508P	Pharmacognosy II – Practical	5	10	4 Hr	15	35	4 Hrs	50
	Total	65	105	17 Hr	170	480	27 Hrs	650

Semester VI

		Internal Assessment				End Semester Exams		
Course code	Name of the course	Continuous	Sessior	al Exams		Maalaa	Derestien	Total Marks
		Mode	Marks	Duration	Total	Marks	Duration	
BP601T	Medicinal Chemistry III – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP602T	Pharmacology III – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP603T	Herbal Drug Technology – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP604T	Biopharmaceutics and Pharmacokinetics – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP605T	Pharmaceutical Biotechnology– Theory	10	15	1 Hr	25	75	3 Hrs	100
BP606T	Quality Assurance– Theory	10	15	1 Hr	25	75	3 Hrs	100
BP607P	Medicinal chemistry III – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP608P	Pharmacology III – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP609P	Herbal Drug Technology – Practical	5	10	4 Hrs	15	35	4 Hrs	50
	Total	75	120	18 Hrs	195	555	30 Hrs	750

Semester VII

		Internal Assessment				End S Ex		
Course code	Name of the course	Continuous	Sessional Exams		Tatal			Total Marks
		Mode	Marks	Duration	Total	Marks	Duration	
BP701T	Instrumental Methods of Analysis – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP702T	Industrial Pharmacy -II– Theory	10	15	1 Hr	25	75	3 Hrs	100
BP703T	Pharmacy Practice – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP704T	Novel Drug Delivery System – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP705 P	Instrumental Methods of Analysis – Practical	5	10	4 Hrs	15	35	4 Hrs	50
BP706 PS	Practice School*	25	-	-	25	125	5 Hrs	150
	Total	70	70	8Hrs	140	460	21 Hrs	600

* The subject experts at college level shall conduct examinations

Semester VIII

		Internal Assessment				End Semester Exams		
Course code	Name of the course	Continuous	Session	al Exams		Maalaa		Total Marks
		Mode	Marks	Duration	Total	Marks	Duration	
BP801T	Biostatistics and Research Methodology – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP802T	Social and Preventive Pharmacy – Theory	10	15	1 Hr	25	75	3 Hrs	100
BP803ET	Pharma. Marketing Management–Theory							
BP804ET	Pharmaceutical Regulatory Science – Theory							
BP805ET	Pharmacovigilance – Theory							
BP806ET	Quality Control and Standardizations of Herbals –Theory							
BP807ET	Computer Aided Drug Design –Theory							
BP808ET	Cell and Molecular Biology –Theory							100 +
BP809ET	Cosmetic Science – Theory	10 + 10 = 20	15 + 15 = 30	1 + 1 = 2 Hrs	25 + 25 = 50	75 + 75 = 150	$\begin{vmatrix} 3+3=6\\ Hrs \end{vmatrix}$	100 = 200
BP810ET	Experimental Pharmacology							
BP811ET	Advanced Instrumentation Techniques – Theory							
BP812ET	Dietary Suppliments and Nutraceuticals							
BP813PW	Project Work	-	-	-	-	150	4 Hrs	150
	Total	40	60	4 Hrs	100	450	16 Hrs	550

11.2 Internal assessment: Continuous mode

The marks allocated for Continuous mode of Internal Assessment shall be awarded as per the scheme given below.

Table-XI: Scheme for awarding internal a	assessment: Continuous mode
--	-----------------------------

Theory				
Criteria	Maximum Marks			
Attendance (Refer Table – XII)	4	2		
Academic activities (Average of any 2 activities e.g. quiz, assignment,4open book test, field work, group discussion and seminar)4				
Student – Teacher interaction	2			
Total	10	5		
Practical				
Attendance (Refer Table – XII)	2	2		
Based on Practical Records, Regular viva voce, etc. 3				
Total	5	;		

Table- XII: Guidelines for the allotment of marks for attendance

Percentage of Attendance	Theory	Practical
95 - 100	4	2
90 - 94	3	1.5
85 - 89	2	1
80 - 84	1	0.5
Less than 80	0	0

11.2.1. Sessional Exams

Two Sessional exams shall be conducted for each theory / practical course as per the schedule fixed by the college(s). The scheme of question paper for theory and

practical Sessional examinations is given below. The average marks of two Sessional exams shall be computed for internal assessment as per the requirements given in tables - X.

Sessional exam shall be conducted for 30 marks for theory and shall be computed for 15 marks. Similarly Sessional exam for practical shall be conducted for 40 marks and shall be computed for 10 marks. The duration for the conduct of the exam is as below

Exam Type	Marks allotted	Duration
Theory	30	1.5 Hr
Practical	40	04 Hr

Question paper pattern for theory Sessional

For subjects having University exams

I. Objective Type Questions (Answer 05 out of 7)	=5 x 2 = 10
II. Long Answers (Answer 1 out of 2)	=1 x 10 = 10
III. Short Answers (Answer 2 out of 3)	$=2 \times 5 = 10$
Total	30 marks

For subjects having Non University Examination

I. Long Answers (Answer 1 outof2)	=1 x 10 = 10		
II.Short Answers (Answer 4 outof 6)	$=4 \times 5 = 20$		
Total	30 marks		

Question paper pattern for practical sessional examinations

I. Synopsis	= 10
II. Experiments	= 25
III. Viva voce	= 05
Total	40 marks

12. Promotion and award of grades

A student shall be declared PASS and eligible for getting gradein a course of B.Pharm.program if he/she secures at least 50% marks in that particular course

including internal assessment .For example, to be declared as PASS and to get grade, the student has to secure a minimum of 50 marks for the total of 100 including continuous mode of assessment and end semester theory examination and has to secure a minimum of 25 marks for the total 50 including internal assessment and end semester practical examination.

13. Carry forward of marks

In case a student fails to secure the minimum 50% in any Theory or Practical course as specified in 12, then he/she shall reappear for the end semester examination of that course. However his/her marks of the Internal Assessment shall be carried over and he/she shall be entitled for grade obtained by him/her on passing.

14. Improvement of internal assessment

A student shall have the opportunity to improve his/her performance only oncein the Sessional exam component of the internal assessment. The re-conduct of the Sessional exam shall be completed before the commencement of next end semester theory examinations.

15. Re-examination of end semester examinations

Reexamination of end semester examination shall be conducted as per the schedule given in table XIII. The exact dates of examinations shall be notified from time to time.

Table-XIII: Tentative schedule of end semester examinations

Semester	For Regular Candidates	For Failed Candidates
I, III, V and VII	November / December	May / June
II, IV, VI and VIII	May / June	November / December

Question paper pattern for end semester theory examinations

For 75 marks paper

I. Objective Type Questions (Answer 5 out of 7)	=5x 3= 15
II. Long Answers (Answer 2 out of 4)	$= 2 \times 10 = 20$
III. Short Answers (Answer 8 out of 10)	$= 8 \ge 5 = 40$
Total	= 75marks

For 50 marks paper

I. Long Answers (Answer 2 out of 3)	$= 2 \times 10 = 20$
II. Short Answers (Answer 6 out of 8)	$= 6 \ge 5 = 30$
Total	= 50 marks

For 35 marks paper

I. Long Answers (Answer 1out of 2)	$= 1 \times 10 = 10$
II. Short Answers (Answer 5 out of 7)	$= 5 \times 5 = 25$
Total	= 25marks

Question paper pattern for end semester practical examinations

I. Synopsis	= 5
II. Experiments	= 25
III. Viva voce	= 05
Total	= 35marks

16. Academic Progression:

No student shall be admitted to any examination unless he/she fulfills the norms givenin6. Academic progression rules are applicable as follows:

A student shall be eligible to carry forward all the courses of I, II and III semesters till the IV semester examinations. However, he/she shall not be eligible to attend the courses of V semester until all the courses of I and II semesters are successfullycompleted.

A student shall be eligible to carry forward all the courses of III, IV and V semesters till the VI semester examinations. However, he/she shall not be eligible to attend the courses of VII semester until all the courses of I, II, III and IV semesters are successfully completed.

A student shall be eligible to carry forward all the courses of V, VI and VII semesters till the VIII semester examinations. However, he/she shall not be eligible to get the course completion certificate until all the courses of I, II, III, IV, V and VI semesters are successfully completed.

A student shall be eligible to get his/her CGPA upon successful completion of the courses of I to VIII semesters within the stipulated time period as per the norms specified in 26.

A lateral entry student shall be eligible to carry forward all the courses of III, IV and V semesters till the VI semester examinations. However, he/she shall not be eligible to attend the courses of VII semester until all the courses of III and IV semesters are successfullycompleted.

A lateral entry student shall be eligible to carry forward all the courses of V, VI and VII semesters till the VIII semester examinations. However, he/she shall not be eligible to get the course completion certificate until all the courses of III, IV, V and VI semesters are successfullycompleted.

A lateral entry student shall be eligible to get his/her CGPA upon successful completion of the courses of III to VIII semesters within the stipulated time period as per the norms specified in 26.

Any student who has given more than 4 chances for successful completion of I / III semester courses and more than 3 chances for successful completion of II / IV semester courses shall be permitted to attend V / VII semester classes

ONLY during the subsequent academic year as the case may be. In simpler terms there shall NOT be any ODD BATCH for anysemester.

Note: Grade ABshould be considered as failed and treated as one head for deciding academic progression. Such rules are also applicable for those students who fail to register for examination(s) of any course in anysemester.

Rules for Carry Forward:

The curriculum (including regulations, structure and syllabi) is in force from academic year 2018-19 and onwards for First Year B. Pharm, for academic

year 2019- 20 onwards for Second Year B. Pharm., for academic year 2020-21 and onwards for Third Year B. Pharm., and for academic year 2021-22 and onwards for Final Year B. Pharm.

The learners who were admitted to First Year B. Pharm. of 2015 pattern during the academic year 2017-18 or before & failed in the First Year B.Pharm. of 2015 pattern examination will have to take admission to Semester-III of Second Year B. Pharm. of 2018 pattern in academic year 2019-20 or onwards, provided that

a. Their result of F. Y. B. Pharm of 2015 pattern is either pass or fails with A. T. K. T.

The said students will have to take up additional remedial courses as follows.

b. The learners who were admitted to S.Y B. Pharm. of 2015 pattern during the academic year 2018-19 or before and fail in the S.Y B.Pharm. of 2015 pattern examination will have to take admission to Semester-V of Third Year B. Pharm. of 2018 pattern in academic year 2020-21 or onwards, provided thatTheir result of S. Y. B. Pharm of 2015 pattern is either pass or fails with A. T. K. T. The said students will have to take up additional remedial course as follows.

17. Grading of performances:

Letter grades and grade points allocations:

Based on the performances, each student shall be awarded a final letter grade at the end of the semester for each course. The letter grades and their corresponding grade points are given in Table –XII.

Percentage of Marks Obtained	Letter Grade	Grade Point	Performance
90.00 - 100	0	10	Outstanding
80.00 - 89.99	А	9	Excellent
70.00 - 79.99	В	8	Good
60.00 - 69.99	С	7	Fair
50.00 - 59.99	D	6	Average
Less than 50	F	0	Fail
Absent	AB	0	Fail

Table – XIV: Letter grades and grade points equivalent to Percentage of marks and performances

A learner who remains absent for any end semester examination shall be assigned a letter grade of AB and a corresponding grade point of zero. He/she should reappear for the said evaluation/examination in due course.

18. The Semester grade point average(SGPA)

The performance of a student in a semester is indicated by a number called 'Semester Grade Point Average' (SGPA). The SGPA is the weighted average of the grade points obtained in all the courses by the student during the semester. For example, if a student takes five courses(Theory/Practical) in a semester with credits C1, C2, C3, C4 and C5 and the student's grade points in these courses are G1, G2, G3, G4 and G5, respectively, and then students' SGPA is equal to:

SGPA=
$$\begin{array}{c} C_1G_1 + C_2G_2 + C_3G_3 + C_4G_4 + C_5G_5 \\ \hline C1 + C2 + C3 + C4 + C5 \end{array}$$

The SGPA is calculated to two decimal points. It should be noted that, the SGPA for any semester shall take into consideration the F and ABS grade awarded in that semester. For example if a learner has a F or ABS grade in course 4, the SGPA shall then be computed as:

SGPA=
$$C1G1 + C2G2 + C3G3 + C4* ZERO + C5G5$$

 $C1 + C2 + C3 + C4+ C5$

19. Cumulative Grade Point Average(CGPA)

The CGPA is calculated with the SGPA of all the VIII semesters to two decimal points and is indicated in final grade report card/final transcript showing the grades of all VIII semesters and their courses. The CGPA shall reflect the failed status in case of F grade(s),till the course(s) is/are passed. When the course(s)is/are passed by obtaining a pass grade on subsequent examination(s) the CGPA shall only reflect the new grade and not the fail grades earned earlier. The CGPA is calculated as:

 $CGPA = \frac{C1S1 + C2S2 + C3S3 + C4S4 + C5S5 + C6S6 + C7S7 + C8S8}{C1 + C2 + C3 + C4 + C5 + C6 + C7 + C8}$

where $C_1, C_2, C_3,...$ is the total number of credits for semester I,II,III,.... and $S_1, S_2, S_3,...$ is the SGPA of semester I,II,III,....

20. Declaration of class

The class shall be awarded on the basis of CGPA as follows

First Class with Distinction	= CGPA of. 7.50 and above
First Class	= CGPA of. 6.00 to7.49
Second Class	= CGPA of. 5.00 to 5.99

21. Project work

A] Selection of the Project Topic

All the students shall undertake a projectunder the supervision of a teacher and submit a report. The project can be based on Lab oriented(small part of original research work) Study /Survey oriented or Computational studies or oriented. / Review topic/ Extension of Practice school work etc., based on Current Trends in Pharmaceutical science. The project shall be carried out in group not exceeding 5 in number. The project report shall be submitted in triplicate (typed &hard bound copy not less than 25 pages).

The internal and external examiner appointed for evaluation of the project shall be approved teachers of SPPU /Industrial Experts appointed by Principal of the respective institute. Students shall be evaluated in groups for four hours (i.e., about half an hour for a group of five students). The projects shall be evaluated as per the criteria given below

Evaluation of Dissertation Book:

15Marks
20Marks
20Marks
20Marks

```
Total 75Marks
```

<i>Evaluation of Presentation:</i> Presentation of work	25Marks
Communications kills	20Marks
Question and answers kills	30Marks

Total

Explanation: All the students should be evaluated thoroughly based on their performance in the Laboratory /Literature work and presentation done as individual student under given criteria.

75Marks

B] Practice School /Project Coordinator:

One of the Staff members shall be assigned as the Project coordinator for a given Academic Year.

Duties of the Coordinator:

- a. Overall co-ordination
- b. Facilitator in Guide-Student allotment.
- c. Preparation of schedules and Time tables.
- d. All relevant documentation and filing
- e. Submission of marks to and communication with College and University exam sections.

22. Industrial training (Desirable)

Every candidate shall be required to work for at least 150 hours spread over four weeks in a Pharmaceutical Industry/Hospital. It includes Production unit, Quality Control department, Quality Assurance department, Analytical laboratory, Chemical manufacturing unit, Pharmaceutical R&D, Hospital (Clinical Pharmacy), Clinical Research Organization, Community Pharmacy, etc. After the Semester – VI and before the commencement of Semester – VII, and shall submit satisfactory report of such work and certificate duly signed by the authority of training organization to the head of the institute.

AND/OR

Every candidate shall be required to undergo any one of the Skill development modules mentioned below (Duration – Min. 04 weeks)

- a) Hands on Training (Central instrumentation lab/Machine room etc)
- **b)** UGC/AICTE recognized online courses (SWAYAM/NPTEL etc)

After the successful completion of the module the candidate shall submit satisfactory report and certificate duly signed by the authority of training organization/Head of the institute

23. Practice School

In the VII semester, every candidate shall undergo practice school for a period of 150 hours evenly distributed throughout the semester. The student shall opt any one of the domains for practice school declared by the program committee from time to time.

At the end of the practice school, every student shall submit a printed report (in triplicate) on the practice school he/she attended (not more than 25 pages). Along with the exams of semester VII, the report submitted by the student, knowledge and skills acquired by the student through practice school shall be evaluated by the subject experts at college level and grade point shall be awarded.

24. Award of Ranks

Ranks and Medals shall be awarded on the basis of final CGPA. However, candidates who fail in one or more courses during the B.Pharm program shall not be eligible for award of ranks. Moreover, the candidates should have completed the B. Pharm program in minimum prescribed number of years, (four years) for the award of Ranks.

25. Award of degree

Candidates who fulfill the requirements mentioned above shall be eligible for award of degree during the ensuing convocation.

26. Duration for completion of the program of study

The duration for the completion of the program shall be fixed as double the actual duration of the program and the students have to pass within the staid period, otherwise they have to get fresh Registration.

27. Re-admission after break of study

Candidate who seeks re-admission to the program after break of study has to get the approval from the university by paying a condonation fee.

No condonation is allowed for the candidate who has more than 2 years of break up period and he/she has to rejoin the program by paying the requiredfees.

FINAL YEAR B. PHARM SEMESTER – VII

BP701T	INSTRUMENTAL METHODS OF ANALYSIS	
	(Theory)	Hours

Scope:

This subject deals with the application of instrumental methods in qualitative and quantitative analysis of drugs. This subject is designed to impart a fundamental knowledge on the principles and instrumentation of spectroscopic and chromatographic technique. This also emphasizes on theoretical and practical knowledge on modern analytical instruments that are used for drug testing.

Objectives:

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

- 1. Upon completion of the course the student shall be ableto
- 2. Illustrate the interaction of matter with electromagnetic radiations and justify its applications in drug analysis
- 3. Classifythechromatographicseparationmethodsandchooseappropriatetechniquefor analysis of drugs.
- 4. Design methods for performing quantitative & qualitative analysis of drugs using various analytical instruments.

Course Content:

UNIT - I

UV Visible spectroscopy

Introduction to spectroscopy, Electronic transitions, chromophores, auxochromes, spectral shifts, solvent effect on absorption spectra, Beer and Lambert's law, Derivation and deviations.

Instrumentation - Sources of radiation, wavelength selectors, sample cells, detectors- Photo tube, Photomultiplier tube, Photo voltaic cell, Silicon Photodiode. Hours

10

Applications - Spectrophotometric titrations, Single component and multi component Analysis

Fluorimetry

Theory, Concepts of singlet, doublet and triplet electronic states, internal and external conversions, factors affecting fluorescence, quenching, instrumentation and applications

UNIT	-11	
FTIR	spectroscopy	
	Introduction, fundamental modes of vibrations in poly atomic molecules, sample handling, factors affecting vibrations	10
	Instrumentation - Sources of radiation, wavelength selectors, detectors - Golay cell, Bolometer, Thermocouple, Thermister, Pyroelectric detector, FTIR instrument, sample handling attachments –DRS and ATR and applications	Hours
Flame	Photometry	
	Principle, interferences, instrumentation and applications	
	Atomic absorption spectroscopy	
	Principle, interferences, instrumentation and Applications	
	Nepheloturbidimetry	
	Introduction	
UNIT	-111	
	Introduction to chromatography -	
	Adsorption and partition column chromatography:	
	Methodology, advantages, disadvantages and applications.	
	Paper chromatography:	
	Introduction, methodology, development techniques, advantages, disadvantages and applications	10 Hours
	Thin layer chromatography:	
	Introduction, Principle, Methodology, Rf values, advantages, disadvantages and applications.	
	HPTLC:	
	Introduction, Instrumentation and applications	
UNIT	-IV	
	Theory of Chromatography	
	Plate theory, Rate theory, System suitability parameters	
	Gas chromatography	08
	Introduction, theory, instrumentation, temperatureprogramming, advantages, disadvantages and applications	Hours
	High performance liquid chromatography (HPLC)	
	Introduction, theory, instrumentation, advantages and applications.	

UNIT –V	UNIT –V	
---------	---------	--

UNIT	-V	
	Ion exchange chromatography-	
	Introduction, classification, ion exchange resins, properties, mechanism of ion exchange process, factors affecting ion exchange, methodology and applications	07 Hours
	Gel chromatography-	
	Introduction, theory, instrumentation and applications Affinity chromatography- Introduction	
Recon	nmended Books (Latest Editions):	
1.	Instrumental Methods of Chemical Analysis by B.K Sharma	
2.	Organic spectroscopy by Y.RSharma	
3.	Text book of Pharmaceutical Analysis by Kenneth A.Connors	
4.	Vogel's Text book of Quantitative Chemical Analysis by A.I.Vogel	
5.	Practical Pharmaceutical Chemistry by A.H. Beckett and J.B.Stenlake	
6.	Organic Chemistry by I. L.Finar	
7.	Organic spectroscopy by WilliamKemp	
8.	Quantitative Analysis of Drugs by D. C.Garrett	
9.	Quantitative Analysis of Drugs in Pharmaceutical Formulations by P. D.Sethi	
10.	Spectrophotometric identification of Organic Compounds bySilverstein.	

Ι

BP70	2T INDUSTRIAL PHARMACY -II (Theory)	45 Hours
Scope	:	
	ourse is designed to impart fundamental knowledge on pharmaceutical product opment and translation from laboratory to market.	
Obje	tives: Upon completion of the course, the student shall be able to:	
1.	Know the process of pilot plant and scale up of pharmaceutical dosage forms	
2.	Understand the process of technology transfer from lab scale to commercial ba	atch
3.	Know different Laws and Acts that regulate pharmaceutical industry	
4.	Understand the approval process and regulatory requirements for drug product	S
Cour	se Content:	
re or	eneral considerations - including significance of personnel requirements, space quirements, raw materials, Pilot plant scale up considerations for solids, liquid als, semi solids and relevant documentation, SUPAC guidelines, Introduction platform technology.	10 Hours
UNIT		
Tech	ology development and transfer:	
tra (P fin ec tra pr	HO guidelines for Technology Transfer (TT): Terminology, Technology insfer protocol, Quality risk management, Transfer from R & D to production rocess, packaging and cleaning), Granularity of TT Process (API, excipients, hished products, packaging materials) Documentation, Premises and uipments, qualification and validation, quality control, analytical method insfer, Approved regulatory bodies and agencies, Commercialization- acticalaspectsandproblems(casestudies), TTagencies in India - APCTD, RDC, TIFAC, BCIL, TBSE / SIDBI; TT related documentation -	10 Hours

	Regulatory affairs: Introduction, Historical overview of Regulatory Affairs, Regulatory authorities, Role of Regulatory affairs department, Responsibility of Regulatory Affairs Professionals	
Re	gulatory requirements for drug approval:	
	Drug Development Teams, Non-Clinical Drug Development, Pharmacology, Drug MetabolismandToxicology,GeneralconsiderationsofInvestigationalNewDrug(IN D)	10 Hours
	Application,Investigator'sBrochure(IB)andNewDrugApplication(NDA),Clinical research / BE studies, Clinical Research Protocols, Biostatistics in Pharmaceutical	
	Product Development, Data Presentation for FDA Submissions, Management of Clinical Studies.	
	UNIT-IV	
	Indian Regulatory Requirements: Central Drug Standard Control Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities, Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval procedures for New Drugs.	07 Hours
U	Central Drug Standard Control Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities, Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval procedures for New	-
UI Qu	Central Drug Standard Control Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities, Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval procedures for New Drugs. NIT-V rality management systems: Quality management & Certifications: Concept of Quality, Total Quality Management, Quality by Design (QbD), Six Sigma concept, Out of Specifications (OOS), Change control, Introduction to ISO 9000 series of quality systems standards, ISO 14000, NABL, GLP	Hours 08
UI Qu	Central Drug Standard Control Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities, Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval procedures for New Drugs. NIT-V Tality management systems: Quality management & Certifications: Concept of Quality, Total Quality Management, Quality by Design (QbD), Six Sigma concept, Out of Specifications (OOS), Change control, Introduction to ISO 9000 series of quality systems standards, ISO 14000, NABL, GLP	Hours 08
UI Qu Ra 1.	Central Drug Standard Control Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities, Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval procedures for New Drugs. NIT-V Tality management systems: Quality management & Certifications: Concept of Quality, Total Quality Management, Quality by Design (QbD), Six Sigma concept, Out of Specifications (OOS), Change control, Introduction to ISO 9000 series of quality systems standards, ISO 14000, NABL, GLP recommended Books: (Latest Editions) Regulatory Affairs from Wikipedia, the free encyclopedia modified on 7 th April	Hours 08
UI Qu	Central Drug Standard Control Organization (CDSCO) and State Licensing Authority: Organization, Responsibilities, Certificate of Pharmaceutical Product (COPP), Regulatory requirements and approval procedures for New Drugs. NIT-V nality management systems: Quality management & Certifications: Concept of Quality, Total Quality Management, Quality by Design (QbD), Six Sigma concept, Out of Specifications (OOS), Change control, Introduction to ISO 9000 series of quality systems standards, ISO 14000, NABL, GLP ecommended Books: (Latest Editions) Regulatory Affairs from Wikipedia, the free encyclopedia modified on 7 th April available at http://en.wikipedia.org/wiki/Regulatory_Affairs. International Regulatory Affairs Updates, 2005.available	Hours 08

Scope:

In the changing scenario of pharmacy practice in India, for successful practice of Hospital Pharmacy, the students are required to learn various skills like drug distribution, drug information, and therapeutic drug monitoring for improved patient care. In community pharmacy, students will be learning various skills such and dispensing of drugs, responding to minor ailments by providing suitable safe medication, patient counseling for improved patient care in the community setup.

Objectives:

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

- 1. Know various drug distribution methods in a hospital
- 2. Appreciate the pharmacy stores management and inventory control
- 3. Monitor drug therapy of patient through medication chart review and clinical review.
- 4. Obtain medication history interview and counsel the patients
- 5. Identify drug related problems
- 6. Detect and assess adverse drug reactions
- 7. Interpret selected laboratory results (as monitoring parameters in therapeutics) of specific disease states
- 8. Know pharmaceutical care services
- 9. Do patient counseling in community pharmacy;
- 10. Appreciate the concept of rational drug therapy.

Course Content:

UNIT-I

Hospital and it's organization

Definition, Classification of hospital- Primary, Secondary and Tertiary hospitals, Classification based on clinical and non- clinical basis, Organization Structure of a Hospital, and Medical staffs involved in the hospital and their functions.

Hospital pharmacy and its organization

Definition, functions of hospital pharmacy, Organization structure, Location, Layout and staff requirements, and Responsibilities and functions of hospital pharmacists.

Adverse drug reaction

Classifications - Excessive pharmacological effects, secondary pharmacological effects, idiosyncrasy, allergic drug reactions, genetically determined toxicity, toxicity following sudden withdrawal of drugs, Drug interaction- beneficial interactions, adverse interactions, and pharmacokinetic drug interactions, Methods for detecting drug interactions ,spontaneous case reports and record linkage

studies, and Adverse drug reaction reporting and management.	
Community Pharmacy	
Organization and structure of retail and wholesale drug store, types and design,	
Legal	
requirementsforestablishmentandmaintenanceofadrugstore, Dispensing of propriet	
ary products, maintenance of records of retail and wholesale drugstore.	
UNIT-II	10
Drug distribution system in a hospital	Hour
Dispensing of drugs to inpatients, types of drug distribution systems, charging	
policy and labelling, dispensing of drugs to ambulatory patients ,and Dispensing	
of controlled drugs. Hospital formulary	
Definition, contents of hospital formulary, Differentiation of hospital formulary	
and Drug list, preparation and revision, and addition and deletion of drug from	
hospital formulary. Therapeutic drug monitoring	
Need for Therapeutic Drug Monitoring, Factors to be considered during the	
Therapeutic Drug Monitoring, and Indian scenario for Therapeutic Drug	
Monitoring. Medication adherence	
Causes of medication non-adherence, pharmacist role in the medication	
adherence, and monitoring of patient medication adherence.	
Patient medication history interview	
Need for the patient medication history interview, medication interview forms.	
Community pharmacy management	
Financial, materials, staff, and infrastructure requirements.	
UNIT-III	
Pharmacy and therapeutic committee	
Organization, functions, Policies of the pharmacy and therapeutic	
committee in including drugs into formulary, inpatient and outpatient	
prescription, automatic stop order, and emergency drug list preparation.	
Drug information services	
-	
Drug and Poison information centre, Sources ofdrug information,	
-	10
Drug and Poison information centre, Sources ofdrug information,	
Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information.	Hou
Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling	Hou s
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and 	
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and Special cases that require the pharmacist 	
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and Special cases that require the pharmacist Education and training program in the hospital 	
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and Special cases that require the pharmacist Education and training program in the hospital Role of pharmacist in the education and training program, Internal and external training program, Services to the nursing homes/clinics, Code of 	
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and Special cases that require the pharmacist Education and training program in the hospital Role of pharmacist in the education and training program, Internal and 	
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and Special cases that require the pharmacist Education and training program in the hospital Role of pharmacist in the education and training program, Internal and external training program, Services to the nursing homes/clinics, Code of ethics for community pharmacy, and Role of pharmacist in the 	Hou s
 Drug and Poison information centre, Sources ofdrug information, Computerized services, and storage and retrieval of information. Patient counseling Definition of patient counseling; steps involved in patient counseling, and Special cases that require the pharmacist Education and training program in the hospital Role of pharmacist in the education and training program, Internal and external training program, Services to the nursing homes/clinics, Code of ethics for community pharmacy, and Role of pharmacist in the interdepartmental communication and community health education. 	

UNI	IT-IV	
	get preparation and implementation Budget preparation and lementation Clinical Pharmacy	
resp char	oduction to Clinical Pharmacy, Concept of clinical pharmacy, functions and onsibilities of clinical pharmacist ,Drug therapy monitoring-medication t review, clinical review, pharmacist intervention, Ward round participation, lication history and Pharmaceutical care.	08 Hour
Dos	ing pattern and drug therapy based on Pharmacokinetic & disease pattern.	S
Ove	er the counter (OTC) sales	
	Introduction and sale of over the counter, and Rational use of common over the counter medications.	
UNI	IT-V	
Dru	g store management and inventory control	
	Organization of drug store, types of materials stocked and storage conditions, Purchase and inventory control: principles, purchase procedure, purchase order, procurement and stocking, Economic order quantity, Reorder quantity level, and Methods used for the analysis of the drug expenditure.	07 Hour
Inve	estigational use of drugs	S
	Description, principals involved, classification, control, identification, role of hospital pharmacist, advisory committee.	
Inte	rpretation of Clinical Laboratory Tests	
Bloc	od chemistry, hematology, and urinalysis	
Rec	ommended Books (Latest Edition):	
1.	Merchant S.H. and Dr. J. S. Quadry. A textbook of hospital pharmacy, 4th ed. Ahmadabad: B.S. Shah Prakakshan;2001.	
2.	Parthasarathi G, Karin Nyfort-Hansen, Milap C Nahata. A textbook of Clinical Pharmacy Practice- essential concepts and skills, 1 st ed. Chennai: Orient Longman Private Limited;2004.	
3.	William E. Hassan. Hospital pharmacy, 5 th ed. Philadelphia: Lea &Febiger1986.	
4.	Tipnis Bajaj. Hospital Pharmacy, 1 st ed. Maharashtra: Career Publications;2008.	
5.	Scott LT. Basic skills in interpreting laboratory data, 4thed. American Society of Health System Pharmacists Inc;2009.	
6.	Parmar N.S. Health Education and Community Pharmacy, 18th ed. India:	

CBS Publishers & Distributers;2008.

Journals:

- 1. Therapeutic drug monitoring. ISSN:0163-4356
- 2. Journal of pharmacy practice. ISSN:0974-8326
- 3. American journal of health system pharmacy. ISSN: 1535-2900(online)
- 4. Pharmacy times (Monthly magazine)

Scope:

This subject is designed to impart basic knowledge on the area of novel drug delivery systems.

Objectives:

Upon completion of the course student shall be able

- 1. To understand various approaches for development of novel drug delivery systems.
- 2. To understand the criteria for selection of drugs and polymers for the development of novel drug delivery systems, their formulation and evaluation.

Course Content:

UNIT-I	
Controlled drug delivery systems:	
Introduction, terminology/definitions and rationale, advantages, disadvantages, selection of drug candidates. Approaches to design controlled release formulations based on diffusion, dissolution and ion exchange principles. Physicochemical and biological properties of drugs relevant to controlled release formulations	10 Hours
Polymers:	
Introduction, classification, properties, advantages and application of polymers in formulation of controlled release drug delivery systems.	
UNIT-II	
Microencapsulation:	
Definition, advantages and disadvantages, microspheres /microcapsules, microparticles, methods of microencapsulation, applications	
Mucosal Drug Delivery system:	10 Hawne
Introduction, Principles of bioadhesion / mucoadhesion, concepts, advantages and disadvantages, transmucosal permeability and formulation considerations of buccal delivery systems	10 Hours
Implantable Drug Delivery Systems:	
Introduction, advantages and disadvantages, concept of implants and osmotic pump.	

UNIT-III			
Transdermal Drug Delivery Systems:			
Introduction, Permeation through skin, factors affecting permeation, permeation enhancers, basic components of TDDS, formulation approaches.			
Gastroretentive drug delivery systems:			
Introduction, advantages, disadvantages, approaches for GRDDS $-$ Floating, high density systems, inflatable and gastro adhesive systems and their applications	10 Hours		
Nasopulmonary drug delivery system:			
Introduction to Nasal and Pulmonary routes of drug delivery ,Formulation of Inhalers(dry powder and metered dose), nasal sprays,nebulizers.			
UNIT-IV			
Targeted drug Delivery:			
Concepts and approaches advantages and disadvantages, introduction to liposomes, niosomes, nanoparticles, monoclonal antibodies and their applications.	08 Hours		
UNIT-V			
Ocular Drug Delivery Systems:			
Introduction, intra ocular barriers and methods to overcome –Preliminary study, ocular formulations and ocuserts	07.11		
Intrauterine Drug Delivery Systems:	07 Hours		
Introduction, advantages and disadvantages, development of intra uterine devices (IUDs) and applications			
Recommended Books: (Latest Editions)			
1. Y W. Chien, Novel Drug Delivery Systems, 2nd edition, revised and expanded, Marcel Dekker, Inc., New York, 1992.			
 Robinson, J. R., Lee V. H. L, Controlled Drug Delivery Systems, Marcel Dekker, Inc., New York, 1992. 			
3. Encyclopedia of Controlled Delivery. Edith Mathiowitz, Published by Wiley Interscience Publication, John Wiley and Sons, Inc, New York.Chichester/Weinheim			
 N.K. Jain, Controlled and Novel Drug Delivery, CBS Publishers &Distributors, New Delhi, First edition 1997 (reprint in 2001). 			
5. S.P. Vyas and R.K. Khar, Controlled Drug Delivery -concepts and advances, VallabhPrakashan, New Delhi, First edition2002.			
Jo	ournals		
---	---	---	-------------------
1. Indian Journal of Pharmaceutical Sciences(IPA)			
2. Indian Drugs(IDMA)			
3.	Journa	l of Controlled Release (Elsevier Sciences)	
4.	Drug I	Development and Industrial Pharmacy (Marcel & Decker)	
Inter	rnation	al Journal of Pharmaceutics (Elsevier Sciences)	
BP7	705P	INSTRUMENTAL METHODS OF ANALYSIS (Practical)	04 Hours/ Week
1.	Weig	hts and measures and pharmacopoeia inanalysis	
2.	Determination of absorption maxima and effect of solvent on absorption maxima of organiccompounds		
3.	Assay of Drug product as per IP (Assay of Paracetamol tablet by UV- Spectrophotometry)		
4.	Assay of Drug product by Calibration curvemethod		
5.	Assay of any drug/drug product bycolorimetry.		
6.	Simultaneous estimation of multicomponent formulation by UV spectroscopy(SE/Q analysis)		
7.	Estim	ation of drug by fluorimetry	
8.	Study	of quenching of fluorescence	
9.	Deter	mination of sodium and potassium by flame photometry	
10.	Separ	ation of amino acids by paper chromatography	
11.	Separ	ation of sugars by thin layer chromatography	
12.	Separ	ation of plant pigments by columnchromatography	
13.	Demo	onstration of HPLC instrument	
14.	Demo	onstration of FTIRinstrument	
15.	15. Interpretation of spectra of organic compounds by IR spectroscopy asper pharmacopoeia		

Recommended Books (Latest Editions)

- 1. Instrumental Methods of Chemical Analysis by B.KSharma
- 2. Organic spectroscopy by Y.RSharma
- 3. Text book of Pharmaceutical Analysis by Kenneth A. Connors
- 4. Vogel's Text book of Quantitative Chemical Analysis by A.I.Vogel
- 5. Practical Pharmaceutical Chemistry by A.H. Beckett and J.B.Stenlake
- 6. Organic Chemistry by I. L.Finar
- 7. Organic spectroscopy by WilliamKemp
- 8. Quantitative Analysis of Drugs by D. C. Garrett
- 9. Quantitative Analysis of Drugs in Pharmaceutical Formulations by P. D. Sethi
- 10. HPLC by P.D.Sethi
- 11. HPTLC by P.D. Sethi
- 12. Spectrophotometric identification of Organic Compounds bySilverstein

In the VII semester, every candidate shall undergo practice school for a period of 150 hours evenly distributed throughout the semester. The student shall opt any one of the domains for practice school declared by the program committee from time to time.

At the end of the practice school, every student shall submit a printed report (in triplicate) on the practice school he/she attended (not more than 25 pages). Along with the exams of semester VII, the report submitted by the student, knowledge and skills acquired by the student through practice school shall be evaluated by the subject experts at college level and grade point shall be awarded.

SEMESTER – VIII

BP801T BIOSTATISTICS AND RESEARCH METHODOLOGY (Theory)

Scope:

To understand the applications of Biostatics in Pharmacy. This subject deals with descriptive statistics, Graphics, Correlation, Regression, logistic regression Probability theory, Sampling technique, Parametric tests, Non Parametric tests, ANOVA, Introduction to Design of Experiments, Phases of Clinical trials and Observational and Experimental studies, SPSS, R and MINITAB statistical software's, analyzing the statistical data using Excel.

Objectives:

Upon completion of the course the student shall be able to

- 1. Know the operation of M.S. Excel, SPSS, R and MINITAB®, DoE (Design of Experiment)
- 2. Know the various statistical techniques to solve statistical problems
- 3. Appreciate statistical techniques in solving the problems.

Course content:

UNIT-I Introduction: Statistics, Biostatistics, Frequency distribution Measures of central tendency: Mean, Median, Mode- Pharmaceutical examples Measures of dispersion: Dispersion, Range, standard deviation, Pharmaceutical problems Correlation: Definition,KarlPearson'scoefficientofcorrelation,Multiplecorrelation- Pharmaceuticals examples	10 Hours
UNIT-II Regression: Curve fitting by the method of least squares, fitting the lines y= a + bx and x = a + by, Multiple regression, standard error of regression– Pharmaceutical Examples Probability: Definition of probability, Binomial distribution, Normal distribution, Poisson's distribution, properties– problems, Sample, Population, largesample, smallsample, Null hypothesis, alternative hypothesis, sampling, essence of sampling, types of sampling, Error-I type, Error-II type, Standard error of mean (SEM) - Pharmaceutical examples Parametric test: t-test(Sample, Pooled or Unpaired and Paired), ANOVA, (Oneway and Two way), Least Significance difference	10 Hours

UNIT-III	
Non Parametric tests:	
Wilcoxon Rank Sum Test, Mann-Whitney U test, Kruskal-Wallis test, Friedman	
Test	
Introduction to Research:	
Need for research, Need for design of Experiments, Experiential Design	
	10
Technique, plagiarism	Hours
Graphs:	
Histogram, Pie Chart, Cubic Graph, response surface plot, Counter Plot graph	
Designing the methodology:	
Sample size determination and Power of a study, Report writing and presentation	
ofdata, Protocol, Cohortsstudies, Observational studies, Experimental	
studies, Designing clinical trial, various phases.	
UNIT-IV	
Blocking and confounding system for Two-level factorials	
Regression modeling:	
Hypothesis testing in Simple and Multiple regression nmodels	08
Introduction to Practical components of Industrial and Clinical Trials	Hours
Problems: Statistical Analysis Using Excel, SPSS, MINITAB®, DESIGN OF	
EXPERIMENTS, R - Online Statistical Software's to Industrial and Clinical trial	
approach	
UNIT-V	
Design and Analysis of experiments:	
Factorial Design:	07
Definition, 2 ² , 2 ³ design. Advantage of factorial design	Hours
Response Surface methodology:	
Central composite design, Historical design, Optimization Techniques	
Recommended Books (Latest edition):	
1. Pharmaceutical statistics- Practical and clinical applications, Sanford Bolton,	
publisher Marcel Dekker Inc. NewYork.	
 Fundamental of Statistics – Himalaya Publishing House-S.C.Guptha 	
 Design and Analysis of Experiments –PHI Learning Private Limited, R. 	
Pannerselvam,	
andC.Montgomery	
	1

BP802T	SOCIAL AND PREVENTIVE PHARMACY (Theory)	45 Hours
 challenges. the pharmac Objectives: After the su 1. Acquire pharmac 2. Develop 	ccessful completion of this course, the student shall be able to: high consciousness/realization of current issues related to health and ceutical problems within the country andworldwide. a critical way of thinking based on current health care development. e alternative ways of solving problems related to health and pharmaceutic	e roles of
Definition, o Understand diseases and Sociology a Socio cultu health and c Hygiene an	ral factors related to health and disease, Impact of urbanization on lisease, Poverty and health	10 Hours
Ebola virus dengue, lyn	medicine nciples of prevention and control of diseases such as cholera, SARS, s, influenza, acute respiratory infections, malaria, chicken guinea, nphatic filariasis, pneumonia, hypertension, diabetes mellitus, cancer, on-drug substance abuse	10 Hours
following: surveillance mental hea deafness, U	ealth programs, its objectives, functioning and outcome of the HIV AND AIDS control programme, TB, Integrated disease program (IDSP), National leprosy control programme, National lth program, National programme for prevention and control of niversal immunization programme, National programme for control of Pulse polio programme.	10 Hours
welfare pro PreventionF	ealth intervention programme for mother and child, National family ogramme, National tobacco control programme, National Malaria Program, National programme for the health care for the elderly, Social ramme; role of WHO in Indian national program	08 Hours

UNIT-V Community services in rural, urban and school health: Functions of PHC, Improvement in rural sanitation, national urban health mission, Health promotion and education in school.	
Community services in rural, urban and school health: Functions of PHC, Improvement in rural sanitation, national urban health mission, Health promotion	

BP803ET	PHARMACEUTICAL MARKETING (Theory)	45 Hours	
Scope:			
technical peop managing and industry. The	eutical industry not only needs highly qualified researchers, che ole, but also requires skilled managers who can take the industry I taking the complex decisions which are imperative for the gro Knowledge and Know-how of marketing management groom the enging role in Sales and Product management.	forward by wth of the	
Objective:			
	ns to provide an understanding of marketing concepts and technique the pharmaceutical industry.	es and their	
Course Conte	ent:		
UNIT-I			
Marketing:			
marketing & s	eneral concepts and scope of marketing; Distinction between elling; Marketing environment; Industry and competitive analysis; sumer buying behavior; industrial buying behavior.		
Pharmaceutio	cal market:	10 Hours	
demographic consumer; ma prescribing ha	and qualitative aspects; size and composition of the market; descriptions and socio-psychological characteristics of the rket segmentation & targeting. Consumer profile; Motivation and abits of the physician; patients 'choice of physician and retail nalyzing the Market; Role of market research.		
UNIT-II			
Product decis	ion:		
Classification,	product line and product mix decisions, product life	10 Hours	
	t portfolio analysis; product positioning; New product decisions; ling, packaging and labelling decisions, Product management in l industry.		
UNIT-III			
Promotion:	Promotion:		
personal sellir	10Methods, determinants of promotional mix, promotional budget; An overview of ersonal selling, advertising, direct mail, journals, sampling, retailing, medical akhibition, public relations, online promotional techniques for OTC Products.10		

UNIT-IV	
Pharmaceutical marketing channels:	
Designing channel, channel members, selecting the appropriate channel, conflict in channels, physical distribution management: Strategic importance, tasks inphysical distributionmanagement.	08 Hours
Professional sales representative (PSR):	nours
Duties of PSR, purpose of detailing, selection and training, supervising, norms for customer calls, motivating, evaluating, compensation and future prospects of the PSR.	
UNIT-V	
Pricing:	
Meaning, importance, objectives, determinants of price; pricing methods and strategies, issues in price management in pharmaceutical industry. An overview of DPCO	07
(Drug Price Control Order) and NPPA (National Pharmaceutical Pricing Authority).	
Emerging concepts in marketing:	
Vertical & Horizontal Marketing; Rural Marketing; Consumerism; Industrial Marketing; Global Marketing.	
Recommended Books: (Latest Editions)	
1. Philip Kotler and Kevin Lane Keller: Marketing Management, Prentice Hall ofIndia, NewDelhi	
2. Walker, Boyd and Larreche : Marketing Strategy- Planning and Implementation, Tata MC GrawHill, New Delhi.	
3. Dhruv Grewal and Michael Levy: Marketing, Tata MC GrawHill	
 Arun Kumar and N Menakshi: Marketing Management, Vikas Publishing, India 	
5. RajanSaxena: Marketing Management; Tata MC Graw-Hill (IndiaEdition)	
6. Ramaswamy,U.S&Nanakamari,S:MarketingManagemnt:GlobalPerspective,I ndian Context, Macmilan India, NewDelhi.	
7. Shanker, Ravi: Service Marketing, Excell Books, NewDelhi	
 Subba Rao Changanti, Pharmaceutical Marketing in India (GIFT – Excel series)Excel Publications. 	

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

Objectives:

Upon completion of the subject student shall be able to;

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

Course content:

UNIT-I New Drug Discovery and development **10 Hours** Stages of drug discovery, Drug development process, pre-clinical studies, non-clinical activities, clinical studies, Innovator and generics, Concept of generics, Generic drug product development. **UNIT-II Regulatory Approval Process** Approval processes and timelines involved in Investigational New Drug (IND), New Drug Application (NDA), Abbreviated New Drug Application (ANDA). Changes to an **10 Hours** approved NDA / ANDA. **Regulatory authorities and agencies** Overview of regulatory authorities of India, United States, European Union, Australia, Japan, Canada (Organization structure and types of applications) UNIT-III **Registration of Indian drug product in overseas market 10 Hours** Procedure for export of pharmaceutical products, Technical documentation, Drug Master Files (DMF), Common Technical Document (CTD), electronic Common Technical Document (eCTD), ASEAN Common Technical Document (ACTD) research.

UNIT-IV	
Clinical trials	
Developing clinical trial protocols, Institutional Review Board / Independent Ethics committee - formation and working procedures, Informed consent process and procedures, GCP obligations of Investigators, sponsors & Monitors, Managing and Monitoring clinical trials, Pharmacovigilance -safety monitoring in clinical trials	08 Hours
UNIT-V	
Regulatory Concepts	07 Hours
Basic terminology, guidance, guidelines, regulations, Laws and Acts, Orange book, Federal Register, Code of Federal Regulatory, Purple book	
Recommended books (Latest edition):	
• Drug Regulatory Affairs by SachinItkar, Dr. N.S. Vyawahare, NiraliPrakashan.	
• The Pharmaceutical Regulatory Process, Second Edition Edited by Ira R. Berry and Robert P. Martin, Drugs and the Pharmaceutical Sciences, Vol.185. InformaHealth carepublishers.	
 New Drug Approval Process: Accelerating Global Registrations By Richard A Guarino, MD, 5thedition, Drugsand the Pharmaceutical Sciences, Vol.190. 	
 Guidebook for drug regulatory submissions / Sandy Weinberg. By John Wiley &Sons. Inc. 	
• FDA Regulatory Affairs: a guide for prescription drugs, medical devices, andbiologics	
 /edited by Douglas J. Pisano, David Mantus. 	
Generic Drug Product Development, Solid Oral Dosage forms, Leon Shargeland IsaderKaufer, Marcel Dekker series, Vol.143	
 Clinical Trials and Human Research: A Practical Guide to RegulatoryCompliance By Fay A. Rozovsky and Rodney K.Adams 	
• Principles and Practices of Clinical Research, Second Edition Edited by JohnI. Gallin and Frederick P.Ognibene	
Drugs: From Discovery to Approval, Second Edition By RickNg	

BP805ET PHARMACOVIGILANCE (Theory) 45 Ho	urs
--	-----

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

Objectives:

- At completion of this paper it is expected that students will be able to (know, do, and appreciate):
- Understand importance of drug safetymonitoring.
- Explain History, development, National and international scenario of pharmacovigilance & comprehend dictionaries, coding and terminologies used in pharmacovigilance
- Understand detection and assessment of new adverse drug reactions, Adverse drug reaction reporting systems and communication in pharmacovigilance, Pharmacovigilance Program of India (PvPI) requirement for ADR reporting in India ICH guidelines for ICSR, PSUR, expedited reporting, pharmacovigilance planning. CIOMS requirements for ADR reporting
- Comprehend methods of safety data during pre-clinical, clinical andpost approval phases of drugs' lifecycle.
- Write case narratives of adverse events and their quality.

Course Content:

UNIT-I

Introduction to Pharmacovigilance

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

Introduction to adverse drug reactions10 HoursDefinitions and classification of ADRs, Detection and reporting, Methods in Causality
assessment, Severity and seriousness assessment, Predictability and preventability
assessment, Management of adverse drug reactions10 Hours

Basic terminologies used in pharmacovigilance

Terminologies of adverse medication related events, Regulatory terminologies

UNIT-II	
Drug and disease classification	
Anatomical, therapeutic and chemical classification of drugs, International classification of diseases, Daily defined doses, International Nonproprietary Names for drugs	
Drug dictionaries and coding in pharmacovigilance	
WHO adverse reaction terminologies, MedDRA and Standardized MedDRA queries, WHO drug dictionary, Eudravigilance medicinal product dictionary	10 Hours
Information resources in pharmacovigilance	
Basic drug information resources, Specialized resources for ADRs	
Establishing pharmacovigilance programme	
Establishing in a hospital, Establishment & operation of drug safety department in industry, Contract Research Organizations (CROs), Establishing a national programme.	
UNIT-III	
Vaccine safety surveillance	
Vaccine Pharmacovigilance, Vaccination failure, Adverse events following immunization	
Pharmacovigilance methods	
Passive surveillance – Spontaneous reports and case series, Stimulated reporting,	10
Active surveillance – Sentinel sites, drug event monitoring and registries, Comparative observational studies – Cross sectional study, case control study and cohort study, Targeted clinical investigations	Hours
Communication in pharmacovigilance	
Effective communication in Pharmacovigilance, Communication in Drug Safety Crisis management, Communicating with Regulatory Agencies, Business Partners, Healthcare facilities & Media	
UNIT-IV	
Safety data generation	
Pre-clinical phase, Clinical phase, Post approval phase (PMS)	08
ICH Guidelines for Pharmacovigilance	08 Hours
Organization and objectives of ICH, Expedited reporting, Individual case safety reports, Periodic safety update reports, Post approval expedited reporting, Pharmacovigilance planning, Good clinical practice in pharmacovigilance studies	

UN	IT-V	
Pha	rmacogenomics of adverse drug reaction	
Gen	etics related ADR with example focusing PK parameters.	
CIC	DMS	07 Hours
	MSWorking Groups, CIOMS Form CDSCO (India) and Pharmaco - lance D&C Act and Schedule Y	nours
Diff	erences in Indian and global pharmacovigilance requirements	
Rec	ommended Books (Latest edition):	
1.	Textbook of Pharmacovigilance: S K Gupta, Jaypee Brothers, MedicalPublish	ners.
2.	Practical Drug Safety from A to Z By Barton Cobert, Pierre Biron, Jones and Publishers.	Bartlett
3.	Mann's Pharmacovigilance: Elizabeth B. Andrews, Nicholas, WileyPublisher	s.
4.	Stephens' Detection of New Adverse Drug Reactions: John Talbot, PatrickWa WileyPublishers.	alle,
5.	An Introduction to Pharmacovigilance: Patrick Waller, Wiley Publishers.	
6.	Cobert's Manual of Drug Safety and Pharmacovigilance: Barton Cobert, Jones BartlettPublishers.	&
7.	Textbook of Pharmacoepidemiolog edited by Brian L. Strom, Stephen E Kim Hennessy, WileyPublishers.	mel, Sean
8.	A Textbook of Clinical Pharmacy Practice -Essential Concepts and Skills:G. Parthasarathi, Karin NyfortHansen, Milap C.Nahata	
9.	National Formulary ofIndia	
10.	Text Book of Medicine by YashpalMunjal	
11.	Text book of Pharmacovigilance: concept and practice by GP Mohanta and P	K Manna
12.	http://www.whoumc.org/DynPage.aspx?id=105825&mn1=7347&mn2=7259. 3=7297	&mn
13.	http://www.ich.org/	
14.	http://www.cioms.ch/	
15.	http://cdsco.nic.in/	
16.	http://www.who.int/vaccine_safety/en/	
	http://www.ipc.gov.in/PvPI/pv home.html	

	QUALITY CONTROL AND STANDARDIZATION OF
DI OUOL I	HERBALS(Theory)

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

Objectives:

Upon completion of the subject student shall be able to;

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

Course Content

UNIT-I Basic tests for drugs – Pharmaceutical substances, Medicinal plants materialsand dosage forms, WHO guidelines for quality control of herbal drugs, Evaluation of commercial crude drugs intended foruse	10 Hours
 UNIT-II Quality assurance in herbal drug industry of cGMP, GAP, GMP and GLP in traditional system of medicine 	10 Hours
• WHO guidelines on current Good Manufacturing Practices (cGMP) for Herbal Medicines, WHO guidelines on GACP for Medicinal Plants.	
 UNIT-III EU and ICH guidelines for quality control of herbal drugs. Research Guidelines for Evaluating the Safety and Efficacy of Herbal Medicines 	10 Hours
 UNIT-IV Stability testing of herbal medicines. Application of various chromatographic techniques in standardization of herbal products. Preparation of documents for new drug application and export registration GMP requirements and Drugs & Cosmetics Act provisions. 	08 Hours

UNIT-V	
Regulatory requirements for herbal medicines.	07
WHO guidelines on safety monitoring of herbal medicines in pharmacovigilance systems.	Hours
Comparison of various Herbal Pharmacopoeias.	
Recommended Books (Latest Editions)	
Role Pharmacognosy by Trease and Evans	
Pharmacognosy by Kokate, Purohit andGokhale	
• Rangari, V.D., Text book of Pharmacognosy and Phytochemistry Vol. I,Carrier Pub., 2006.	
• Aggrawal, S.S., Herbal Drug Technology. Universities Press,2002.	
• EMEA. Guidelines on Quality of Herbal Medicinal Products/Traditional Medicinal Products,	
• Mukherjee, P.W. Quality Control of Herbal Drugs: An Approach to Evaluation of Botanicals. Business Horizons Publishers, New Delhi, India,2002.	
• Shinde M.V., Dhalwal K., Potdar K., Mahadik K. Application of quality control principles to herbal drugs. International Journal of Phytomedicine 1(2009); p.4-8.	
• WHO. Quality Control Methods for Medicinal Plant Materials, World Health Organization, Geneva, 1998. WHO. Guidelines for the Appropriate Use of Herbal Medicines. WHO Regional Publications, Western Pacific Series No 3, WHO Regional office for the Western Pacific, Manila,1998.	
• WHO. The International Pharmacopeia, Vol. 2: Quality Specifications, 3rdedn. World Health Organization, Geneva, 1981.	
• WHO. Quality Control Methods for Medicinal Plant Materials. World Health Organization, Geneva,1999.	
• WHO. WHO Global Atlas of Traditional, Complementary and Alternative Medicine. 2 vol. set. Vol. 1 contains text and Vol. 2, maps. World Health Organization, Geneva,2005.	
• WHO. Guidelines on Good Agricultural and Collection Practices (GACP)for Medicinal Plants. World Health Organization, Geneva, 2004.	

BP807ET	COMPUTER AIDED DRUG DESIGN (Theory)	45 Hours
techniques us Objectives: Upon complet 1. Understa 2. Classify 3. Understa 4. Analyse	is designed to provide detailed knowledge of rational drug design process and sed in rational drug design process. etion of the course, the student shall be able to understand and the design and discovery of leadmolecules the role of drug design tools for drug discoveryprocess and and analyse concepts of QSAR anddocking and apply various strategies to develop new drug likemolecules. ous molecular modeling software to design new drugmolecule	various
Course Cont UNIT-I		
Stages of dru Lead discove medicine, Ra discovery bas Introduction Analog Base Case studies Ligand based based (Desig	to Drug Discovery and Development - g discovery and development, ry approaches - Rational approaches to lead discovery based on traditional ndom screening, Non-random screening, serendipitous drug discovery, lead sed on drug metabolism, lead discovery based on clinical observation. to Ligand based and Structure Based DD d Drug Design - Bioisosterism, Bioisosteric replacement - I (Design of inhibitors of tubulin polymerization eg. Colchicine), Structure n of HMG-CoA reductase inhibitors. eg. Statins) and Analog based DD 2 histamine antagonist eg. Cimetidine)	14 Hours
UNIT- II Introduction Introduction Energy Minin minima deter Molecular d Rigid docking	to Computational tools Molecular Modeling - to molecular mechanics and quantum mechanics. nization methods and Conformational Analysis, global conformational mination.	10 Hours
Introduction SAR versu physicochem 2D QSAR - Experimental parameters su constant. Har 3D-QSAR ap COMFA and Pharmacoph	us QSAR, History and development of QSAR, Types of icalparameters and theoretical approaches for the determination of physicochemical uch as Partition coefficient, Hammet's substituent constant and Tafts steric nsch's analysis, Free Wilson analysis pproaches -	14 Hours

UN	NIT- IV	
	formatics & Methods in drug design Introduction to Bioinformatics, chemo formatics Databases -	07 Hours
	emical database, Natural compound database, Drug like compound database , ug bank	
Re	commended Books (Latest Editions)	
1.	Robert GCK, ed., "Drug Action at the Molecular Level" University PrakPress Baltimore.	
2.	Martin YC. "Quantitative Drug Design" Dekker, New York.	
3.	Delgado JN, Remers WA eds "Wilson & Gisvolds's Text Book of OrganicMedicinal & Pharmaceutical Chemistry" Lippincott, NewYork.	
4.	Foye WO "Principles of Medicinal chemistry 'Lea&Febiger.	
5.	Korolkovas A, BurckhalterJH. "Essentials of Medicinal Chemistry" Wiley Interscience.	
6.	WolfME,ed"TheBasisofMedicinalChemistry,Burger'sMedicinalChemistry" John Wiley & Sons,NewYork.	
7.	PatrickGraham,L.,AnIntroductiontoMedicinalChemistry,OxfordUniversityP ress.	
8.	Smith HJ, Williams H, eds, "Introduction to the principles of Drug Design" WrightBoston.	
9.	Silverman R.B. "The organic Chemistry of Drug Design and Drug Action" Academic Press NewYork.	
10.	D. J. Triggle, John Bodenhan Taylor, Peter Kennewell, Comprehensive Medicinal Chemistry, Volume I-VIII : Germany: Elsevier Science.	

BP808ET	CELL AND MOLECULAR BIOLOGY (Theory)	45 Hours
---------	-------------------------------------	----------

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

Objectives:

Upon completion of the subject student shall be able to:

1. Summarize cell and molecular biology history, cellular functioning and Composition & describe the chemical foundations of cell biology.

2. Describe cellular membrane structure and function properties and functions of DNA, CellCycle.

3. Describe basic molecular genetics mechanisms.

4. Understand the cell signaling pathways with their regulations and role indisease process.

Course contents

UNIT-I Cell and Molecular Biology: Definitions theory and basics and Applications. Cell and Molecular Biology: History and Summation. Properties of cells and cell membrane, Prokaryotic versus Eukaryotic, Cellular Reproduction, Chemical Foundations – an Introduction and Reactions (Types)	10 Hours
UNIT-II DNA and the Flow of Molecular Information, DNA Functioning, DNA and RNA, Types of RNA, Transcription and Translation	10 Hours
UNIT-III Proteins: Defined and Amino Acids, Protein Structure, Regularities in Protein Pathways, Cellular Processes, Positive Control and significance of Protein Synthesis	10 Hours
UNIT-IV Science of Genetics, Transgenics and Genomic Analysis, Cell Cycle analysis, Mitosis and Meiosis, Cellular Activities and Checkpoints Clinical phase, Post approval phase (PMS)	08 Hours

UN	T-V	
	Signals: Introduction, Receptors for Cell Signals, Signaling Pathways: rview, Misregulation of Signaling Pathways, Protein-Kinases: Functioning	07 Hours
Rec	ommended Books (latest edition):	
1.	W.B. Hugo and A.D. Russel: Pharmaceutical Microbiology,Blackwell Scientific publications, OxfordLondon.	
2.	Prescott and Dunn., Industrial Microbiology, 4th edition, CBS Publishers & Distributors, Delhi.	
3.	Pelczar, Chan Kreig, Microbiology, Tata McGraw Hill edn.	
4.	Malcolm Harris, Balliere Tindall and Cox: PharmaceuticalMicrobiology. Rose: IndustrialMicrobiology.	
5.	Probisher, Hinsdill et al: Fundamentals of Microbiology, 9th ed.Japan	
6.	Cooper and Gunn's: Tutorial Pharmacy, CBS Publisher and Distribution. Peppler: MicrobialTechnology.	
7.	Edward: Fundamentals of Microbiology.	
8.	N.K.Jain: Pharmaceutical Microbiology, VallabhPrakashan,Delhi	
9.	Bergeys manual of systematic bacteriology, Williams and Wilkins- A WaverlyCompany	
10.	B.R. Glick and J.J. Pasternak: Molecular Biotechnology: Principlesand	
11.	Applications of Recombinant DNA: ASM Press Washington D.C. RA Goldshyet. al., :KubyImmunology.	

BP	809ET	COSMETIC SCIENCE (Theory)	45 Hours
Sco	pe:		
	scourseisdesignedtoi eir formulationstudies	mpartfundamentalknowledgeofcosmeticandcosmeceutica	l products
Ob	jectives:		
Upo	on completion of the	course, the student shall be able to:	
1.	Understand the con incosmetics.	ncepts of cosmetics; anatomy of skin v/s hair, general exe	cipients used
2.		ept of cosmeceuticals, history, difference between c cosmeceuticals agents	cosmetics &
3.	Know different La	ws and Acts that regulate pharmaceutical industry	
4.	Understand the ap	proval process and regulatory requirements for drug produced	ucts
Сот	urse contents		
UN	IT-I		
as p		tic and cosmeceutical products, Definition of cosmetics egulations, Evolution of cosmeceuticals from cosmetics, DTC drugs	
Cos	smetic excipients:		
	factants, rheology ssification and applic	modifiers, humectants, emollients, preservatives. cation	10 Hours
Ski	n: Basic structure an	d function of skin.	
Hai	ir: Basic structure of	hair. Hair growth cycle.	
Ora	al Cavity: Common	problem associated with teeth and gums.	
UN	IT-II		
		on and building blocks of skin care products:	
Fac adv	e wash, Moisturizi	ng cream, Cold Cream, Vanishing cream and their ntages. Application of these products in formulation of	
for sha	mulation and buil mpoo, Hair condition	orants- Actives & mechanism of action. Principles of ding blocks of Hair care products: Conditioning oner, anti-dandruff shampoo. Hair oils, Chemistry and lene diamine based hairdye.	10 Hours
Prir	nciples of formulation	n and building blocks of oral care products:	
Таа	othnaste for bleeding	gums, sensitive teeth. Teeth whitening, Mouthwash.	

UN	NIT-III	
Su	n protection, Classification of Sunscreens and SPF.	
Ro	le of herbs in cosmetics:	
	in Care: Aloe and turmeric Hair care: Henna and amla. Oral care: Neem and ove Analytical cosmetics:	10 Hours
	S specification and analytical methods for shampoo, skin cream and othpaste.	
UN	NIT-IV	
Pr	inciples of Cosmetic Evaluation: Principles of sebumeter, corneometer.	
M	easurement	08 Hours
	TEWL, Skin Color, Hair tensile strength, Hair combing properties, Soaps d syndet bars. Evolution and skin benfits.	
UN	NIT-V	
un	ly and dry skin, causes leading to dry skin, skin moisturisation. Basic derstanding of the terms Comedogenic, dermatitis. Cosmetic problems sociated with Hair and scalp: Dandruff, Hair fall causes	07 Hours
	smetic problems associated with skin: blemishes, wrinkles, acne, prickly heat d body odor.	
Ar	tiperspirants and Deodorants- Actives and mechanism of action	
Re	ferences	
1)	Harry's Cosmeticology, Wilkinson, Moore, Seventh Edition, GeorgeGodwin.	
2)	Cosmetics – Formulations, Manufacturing and Quality Control, P.P. Sharma, 4th Edition, Vandana Publications Pvt. Ltd., Delhi.	
3)	Text book of cosmelicology by Sanju Nanda &Roop K. Khar, TataPublishers.	

	10ET	EXPERIMENTAL PHARMACOLOGY (Theory)	45 Hours
Scop	e:		
	•	t is designed to impart the basic knowledge of preclinical studies in ex luding design, conduct and interpretations of results.	perimental
Obje	ectives		
Upor	n comp	letion of the course the student shall be able to,	
1.	Under	stand the applications of various commonly used laboratory animals.	
2.	Demor	nstrate the various screening methods used in preclinical research.	
3.	Comp	rehend and demonstrate the importance of biostatistics and research met	hodology.
4.	Design	n and execute a research hypothesis independently.	
Cou	rse con	tents	
UNI	T-I		
Labo	oratory	Animals:	
cond	uct of	CPCSEA and OECD guidelines for maintenance, breeding and experiments on laboratory animals, Common lab animals: and applications of different species and strains of animals. Popular	10 Hours
	•	nd mutant animals.	
Tech	niques	nd mutant animals. for collection of blood and common routes of drug administration y animals, Techniques of blood collection and euthanasia.	
Tech	niques borator	for collection of blood and common routes of drug administration	
Tech in lat	iniques borator T-II	for collection of blood and common routes of drug administration	
Tech in lat	T-II Iinical Introdu drug s negati	for collection of blood and common routes of drug administration y animals, Techniques of blood collection and euthanasia. screening models uction: Dose selection, calculation and conversions, preparation of solution/suspensions, grouping of animals and importance of sham	10 Hours
Tech in lat UNI Prec	T-II Inical Introdu drug s negati positiv udy. Stud Parkin activit sedativ	for collection of blood and common routes of drug administration y animals, Techniques of blood collection and euthanasia. screening models uction: Dose selection, calculation and conversions, preparation of solution/suspensions, grouping of animals and importance of sham ve and	10 Hours
Tech in lat UNI Prec a.	T-II Inical Introdu drug s negati positiv udy. Stud Parkin activit sedativ	for collection of blood and common routes of drug administration y animals, Techniques of blood collection and euthanasia. screening models uction: Dose selection, calculation and conversions, preparation of solution/suspensions, grouping of animals and importance of sham we and vecontrolgroups.Rationaleforselectionofanimalspeciesandsexforthest by of screening animal models for Diuretics, nootropics, anti- ison's, antiasthmatics, Preclinical screening models: for CNS y- analgesic, antipyretic, anti-inflammatory, general anaesthetics, we and hypnotics, antipsychotic, antidepressant, antiepileptic,	10 Hours
Tech in lat UNI Prec a. b.	T-II Iinical Introdu drug s negati positiv udy. Stud Parkin activit sedativ antipat	for collection of blood and common routes of drug administration y animals, Techniques of blood collection and euthanasia. screening models uction: Dose selection, calculation and conversions, preparation of solution/suspensions, grouping of animals and importance of sham we and vecontrolgroups.Rationaleforselectionofanimalspeciesandsexforthest by of screening animal models for Diuretics, nootropics, anti- ison's, antiasthmatics, Preclinical screening models: for CNS y- analgesic, antipyretic, anti-inflammatory, general anaesthetics, we and hypnotics, antipsychotic, antidepressant, antiepileptic,	10 Hours

anti aggregatory, coagulants, and anticoagulants Preclinical screening models for other important drugs like antiulcer,	Hours
for CVS activity- antihypertensives, diuretics, antiarrhythmic, antidyslepidemic, anti aggregatory, coagulants, and anticoagulants Preclinical screening models for other important drugs like antiulcer,	Hours
anti aggregatory, coagulants, and anticoagulants Preclinical screening models for other important drugs like antiulcer,	Hours
antidiabetic, anticancer and antiasthmatics	
UNIT-V	
Research methodology and Bio-statistics.	
Selection of research topic, review of literature, research hypothesis and study design Pre- clinical data analysis and interpretation using Students't' test and One-way ANOVA. Graphical representation ofdata07	Hours
Recommended Books (latest edition):	
1. Fundamentals of experimental Pharmacology-byM. N.Ghosh	
2. Hand book of Experimental Pharmacology-S.K. Kulkarni	
3. CPCSEA guidelines for laboratory animal facility.	
4. Drug discovery and Evaluation by Vogel H.G.	
5. Drug Screening Methods by Suresh Kumar Gupta and S. K.Gupta	
6. Introduction to biostatistics and research methods by PSS Sundar Rao and J Richard	

This subject deals with the application of instrumental methods in qualitative and quantitative analysis of drugs. This subject is designed to impart advanced knowledge on the principles and instrumentation of spectroscopic and chromatographic hyphenated techniques. This also emphasizes on theoretical and practical knowledge on modern analytical instruments that are used for drugtesting.

Objectives:

Upon completion of the course the student shall be able to

- 1. Express the principle of the advanced instruments used and justify its applications in drug analysis
- 2. Understand the principles of analytical techniques and its application in analysis of drugs
- 3. Explain the importance and methods for the calibration of various analytical instruments
- 4. Formulate and justify techniques for the analysis of drugs using various analytical instruments.

Course contents

UNIT-I	
Nuclear Magnetic Resonance spectroscopy	
Principles of ¹ H-NMR, chemical shift, factors affecting chemical shift, coupling constant, Spin - spin coupling, relaxation, instrumentation and applications	
¹³ C-NMR- Introduction to ¹³ C-NMR spectroscopy	14 Hours
Mass Spectrometry	
Principles, , Ionization techniques –Electron impact, chemical ionization, MALDI, FAB, Analyzers-Time of flight and Quadrupole, instrumentation, Fragmentation, applications Simple structural elucidation problems	
UNIT-II	
Thermal Methods of Analysis	
Principles, instrumentation and applications of Thermogravimetric Analysis (TGA), Differential Thermal Analysis (DTA), Differential Scanning Calorimetry (DSC)	07 Hours
UNIT-III	
Electrophoresis	10 Houng
Introduction, factors affecting electrophoretic mobility, Techniques of paper, gel,capillary electrophoresis, applications	10 Hours
X-Ray Diffraction Methods	

Origin of X-rays, basic aspects of crystals, Xray Crystallography, rotating crystal technique, single crystal diffraction, powder diffraction, and applications. Calibration of following Instruments Electronic balance, UV-Visible spectrophotometer, IR spectrophotometer, Fluorimeter, HPLC.	
UNIT-IV Radio immuno assay Principle, different methods, Importance, various components, Limitation and Applications of Radioimmunoassay Extraction techniques General principle and procedure involved in the solid phase extraction and liquid-liquid extraction.	06 Hours
UNIT-V Hyphenated techniques Introduction to hyphenated techniques and types of techniques Details of LC- MS, GC-MS, HPTLC-MS, MS/MS.	08 Hours

45 Hours

BP812ET	DIETARY SUPPLEMENTS AND NUTRACEUTICALS (Theory)	45 Hours
---------	---	----------

This subject covers foundational topic that are important for understanding the need and requirements of dietary supplements among different groups in the population.

Objective:

This module aims to provide an understanding of the concepts behind the theoretical applications of dietary supplements. By the end of the course, students should be able to:

- 1. Understand the need of supplements by the different group of people to maintain healthy life.
- 2. Understand the outcome of deficiencies in dietary supplements.
- 3. Recognize the components in dietary supplements and the application.
- 4. Acquaint with the regulatory and commercial aspects of dietary supplements including healthclaims.

Course content:

UNIT-I

Definitions of Functional foods, Nutraceuticals and Dietary supplements. Classification of Nutraceuticals, Health problems and diseases that can be

prevented or cured by Nutraceuticals i.e. weight control, diabetes, cancer, heart disease, stress, osteoarthritis, hypertension etc.	
Public health nutrition, maternal and child nutrition, nutrition and ageing, nutrition education in community.	
Source, Name of marker compounds and their chemical nature, Medicinal uses and health benefits of following used as nutraceuticals/functional foods: Spirulina, Soyabean, Ginseng, Garlic, Broccoli, Gingko, Flaxseeds	
UNIT-II	
Phytochemicals as nutraceuticals : Occurrence and characteristic features (chemical nature medicinal benefits) of following	
Carotenoids - α and β -Carotene, Lycopene, Xanthophylls, leutin	
Sulfides: Diallyl sulfides, Allyl trisulfide.	
Polyphenolics: Reservetrol	15 Hours
Flavonoids- Rutin, Naringin, Quercitin, Anthocyanidins, catechins, Flavones	
Prebiotics / Probiotics.: Fructo oligosaccharides, Lacto bacillum Phyto estrogens : Isoflavones, daidzein, Geebustin, lignans Tocopherols	
Proteins, vitamins, minerals, cereal, vegetables and beverages as functional foods: oats, Wheat bran, rice bran, sea foods, coffee, tea and the like.	
UNIT-III	
Introduction to free radicals: Free radicals, reactive oxygen species, production of free radicals in cells, damaging reactions of free radicals on lipids, proteins, Carbohydrates, nucleic acids.	07 Hours
Dietary fibres and complex carbohydrates as functional food ingredients.	
UNIT-IV	
Free radicals in Diabetes mellitus, Inflammation, Ischemic reperfusion injury, Cancer, Atherosclerosis, Free radicals in brain metabolism and pathology, kidney damage, muscle damage. Free radicals involvement in other disorders. Free radicals theory of ageing.	10 Hours
Antioxidants: Endogenous antioxidants – enzymatic and nonenzymatic antioxidant defense, Superoxide dismutase, catalase, Glutathione peroxidase, Glutathione Vitamin C, Vitamin E, α - Lipoic acid, melatonin Synthetic antioxidants: Butylated hydroxy Toluene, Butylated hydroxy Anisole.	
Functional foods for chronic disease prevention.	
L	

UNI	T-V	
Effect of processing, storage and interactions of various environmental factors on the potential of nutraceuticals.		06 Hours
Regulatory Aspects; FSSAI,FDA, FPO,MPO, AGMARK. HACCP and GMPs on Food Safety. Adulteration of foods.		oo nours
Pha	Pharmacopoeial Specifications for dietary supplements and nutraceuticals.	
Refe	erences:	
1.	Dietetics by SriLakshmi	
2.	Role of dietary fibres and neutraceuticals in preventing diseases by K.T Agusti and P.Faizal: BSPublication.	
3.	Advanced Nutritional Therapies by Cooper. K.A.,(1996).	
4.	The Food Pharmacy by Jean Carper, Simon & Schuster, UK Ltd.,(1988).	
5.	Prescription for Nutritional Healing by James F.Balchand Phyllis A.Balch2 nd Edn., Avery Publishing Group, NY(1997).	
6.	G. Gibson and C.williams Editors 2000 Functional foods Woodhead Publ. Co.London.	
7.	Goldberg, I. Functional Foods. 1994. Chapman and Hall, NewYork.	
8.	Labuza, T.P. 2000 Functional Foods and Dietary Supplements: Safety,	
	Good Manufacturing Practice (GMPs) and Shelf Life Testing in Essentials	
	of Functional Foods M.K. Sachmidl and T.P. Labuza eds. AspenPress.	
9.	Handbook of Nutraceuticals and Functional Foods, Third Edition (Modern	
	Nutrition)	
10.	Shils, ME, Olson, JA, Shike, M. 1994 <i>Modern Nutrition in Health and Disease</i> . Eighth edition. Lea andFebiger	

BP 813 **PW PROJECT WORK**

150 Hours

A] Selection of the Project Topic

All the students shall undertake a project under the supervision of a teacher and submit a report. The project can be based on Lab oriented (small part of original research work) Study / Survey oriented or Computational studies or oriented. / Review topic/ Extension of Practice school work etc., based on Current Trends in Pharmaceutical science. The project shall be carried out in group not exceeding 5 in number. The project report shall be submitted in triplicate (typed & hard bound copy not less than 25 pages).

The internal and external examiner appointed for evaluation of the project shall be approved teachers of SPPU /Industrial Experts appointed by Principal of the respective institute. Students shall be evaluated in groups for four hours (i.e., about half an hour for a group of five students). The projects shall be evaluated as per the criteria given below

B]	Evaluation of Dissertation Book	:	
	Objective(s) of the work done	15M	larks
	Methodology adopted	20M	larks
	Results and Discussions	20M	larks
	Conclusions and Outcomes	20M	larks
	Total	75 M	larks
C]	Evaluation of Presentation:		
	Presentation of work	25M	larks
	Communication skills	20M	larks
	Question and answer skills	30M	larks
	Total	75 M	larks

Explanation: All the students should be evaluated thoroughly based on their performance in the Laboratory /Literature work and presentation done as individual student under given criteria.