# 2016

# THE MASTER OF PHARMACY (M. PHARM.) COURSE REGULATION 2014

(Based on notification in the Gazette of India No. 362, Dated December 11, 2014)

# SCHEME AND SYLLABUS



#### PHARMACY COUNCIL OF INDIA

Combined Council's Building, Kotla Road, Aiwan-E-Ghalib Marg, New Delhi-110 002. Website: www.pci.nic.

#### Table of Contents

S.No.	Content	Page.No.
	Regulations	01
1.	Short Title and Commencement	01
2.	Minimum qualification for admission	01
3.	Duration of the program	01
4.	Medium of instruction and examinations	01
5.	Working days in each semester	01
6.	Attendance and progress	02
7.	Program/Course credit structure	02
8.	Academic work	03
9.	Course of study	03
10.	Program Committee	15
11.	Examinations/Assessments	16
12.	Promotion and award of grades	32
13.	Carry forward of marks	32
14.	Improvement of internal assessment	33
15.	Reexamination of end semester examinations	33
16.	Allowed to keep terms (ATKT)	33
17.	Grading of performances	33
18.	The Semester grade point average (SGPA)	34
19.	Cumulative Grade Point Average (CGPA)	34
20.	Declaration of class	35
21.	Project work	35
22.	Award of Ranks	36
23.	Award of degree	36
24.	Duration for completion of the program of study	36
25.	Revaluation I Retotaling of answer papers	36
26.	Re-admission after break of study	36
27.	Pharmaceutics (MPH)	37
28.	Industrial Pharmacy (MIP)	55
29.	Pharmaceutical Chemistry (MPC)	73
30.	Pharmaceutical Analysis (MPA)	98
31.	Pharmaceutical Quality Assurance (MQA)	119
32.	Pharmaceutical Regulatory Affairs (MRA)	142
33.	Pharmaceutical Biotechnology (MPB)	165
34.	Pharmacy Practice (MPP)	188
35.	Pharmacology (MPL)	209
36.	Pharmacognosy (MPG)	232
37.	Research Methodology & Biostatistics (MRM)	252



असाधारण

#### EXTRAORDINARY

भाग III-खण्ड 4

PART III-Section 4

#### प्राधिकार से प्रकाशित PUBLISHED BY AUTHORITY

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NEW DELIH, THURSDAY, DECEMBER 11, 2014/AGRAHAYANA 20, 1936

#### PHARMACY COUNCIL OF INDIA NOTIFICATION

New Delhi, the 10th December, 2014

#### The Master of Pharmacy (M.Pharm) Course Regulations, 2014

No. 14-136/ 2014-PCL—In exercise of the powers conferred by Sections 10 and 18 of the Pharmacy Act, 1948 (8 of 1948), the Pharmacy Council of India, with the approval of the Central Government hereby makes the following regulations; namely—

#### CHAPTER -I:REGULATIONS

#### 1. Short Title and Commencement

These regulations shall be called as "The Revised Regulations for the Master of Pharmacy (M. Pharm.)Degree Program - Credit Based Semester System (CBSS) 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 the authorities of the university.

#### 2. Minimum qualification for admission

A Pass in the following examinations

- a) B. Pharm Degree examination of an Indian university established by law in India from an institution approved by Pharmacy Council of India and has scored not less than 55 % of the maximum marks (aggregate of 4 years of B.Pharm.)
- b) Every student, selected for admission to post graduate pharmacy program in any PCI approved institution should have obtained registration with the State Pharmacy Council or should obtain the same within one month from the date of his/her admission, failing which the admission of the candidate shall be cancelled.

Note: It is mandatory to submit a migration certificate obtained from the respective university where the candidate had passed his/her qualifying degree (B.Pharm.)

#### 3. Duration of the program

The program of study for M.Pharm. shall extend over a period of four semesters (two academic years). The curricula and syllabi for the program shall be prescribed from time to time by Phamacy Council of India, New Delhi.

#### 4. Medium of instruction and examinations

Medium of instruction and examination shall be in English.

#### 5. Working days in each semester

Each semestershall consist of not less than 100 working days. The odd semesters shall be conducted from the month of June July to November/December and the even semesters shall be conducted from the month of 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, practical classes, seminars, assignments, 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/extracurricular activities is dependent upon the quantum of work expected to be put in for each of these activities per week/per activity.

#### 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 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 a multiplier of half (1/2) for practical (laboratory) hours. Thus, for example, a theory course having four lectures 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.

The contact hours of seminars, assignments and research work shall be treated as that of practical courses for the purpose of calculating credits. i.e., the contact hours shall be multiplied by 1/2. Similarly, the contact hours of journal club, research work presentations and discussions with the supervisor shall be considered as theory course and multiplied by 1.

#### 7.2. Minimum credit requirements

The minimum credit points required for the award of M. Pharm. degree is 95. However based on the credit points earned by the students under the head of co-curricular activities, a student shall earn a maximum of 100 credit points. These credits are divided into Theory courses, Practical, Seminars, Assignments, Research work, Discussions with the supervisor, Journal club and Co-Curricular activities over the duration of four semesters. The credits

are distributed semester-wise as shown in Table 14. Courses generally progress in sequence, 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.

#### 8. Academic work

A regular record of attendance both in Theory, Practical, Seminar, Assignment, Journal club, Discussion with the supervisor, Research work presentation and Dissertation shall be maintained by the department / teaching staff of respective courses.

#### 9. Course of study

The specializations in M.Pharm program is given in Table 1.

Table – 1: List of M.Pharm. Specializations and their Code

S. No.	Specialization	Code
1.	Pharmaceutics	MPH
2.	Industrial Pharmacy	MIP
3.	Pharmaceutical Chemistry	MPC
4.	Pharmaceutical Analysis	MPA
5.	Pharmaceutical Quality Assurance	MQA
6.	Pharmaceutical Regulatory Affairs	MRA
7.	Pharmaceutical Biotechnology	MPB
8.	Pharmacy Practice	MPP
9.	Pharmacology	MPL
10.	Pharmacognosy	MPG

The course of study for M.Pharm specializations shall include Semester wise Theory & Practical as given in Table – 2 to 11. The number of hours to be devoted to each theory and practical course in any semester shall not be less than that shown in Table – 2 to 11.

Table - 2: Course of study for M. Pharm. (Pharmaceutics)

Course Code	Course	Credit Hours	Credit Points	Hrs./w k	Marks
	Seme	ester I			
MPH101T	Modern Pharmaceutical Analytical Techniques	4	4	4	100
MPH102T	Drug Delivery System	4	4	4	100
MPH103T	Modern Pharmaceutics	4	4	4	100
MPH104T	Regulatory Affair	4	4	4	100
MPH105P	Pharmaceutics Practical I	12	6	12	150
-	Seminar/Assignment	7	4	7	100
	Total	35	26	35	650
	Seme	ster II			
MPH201T	Molecular Pharmaceutics (Nano Tech and Targeted DDS)	4	4	4	100
MPH202T	Advanced Biopharmaceutics & Pharmacokinetics	4	4	4	100
МРН203Т	MPH203T Computer Aided Drug Delivery System		4	4	100
MPH204T	MPH204T Cosmetic and Cosmeceuticals		4	4	100
MPH205P	Pharmaceutics Practical II	12	6	12	150
	Seminar/Assignment	7	4	7	100
	Total	35	26	35	650

Table - 5: Course of study for M. Pharm. (Pharmaceutical Analysis)

Course Code	Course	Credit Hours	Credit Points	Hrs./wk	Marks
	Seme	ster I			
MPA101T	Modern Pharmaceutical Analytical Techniques	4	4	4	100
MPA102T	Advanced Pharmaceutical Analysis	4	4	4	100
MPA103T	Pharmaceutical Validation	4	4	4	100
MPA104T	Food Analysis	4	4	4	100
MPA105P	Pharmaceutical Analysis Practical I	12	6	12	150
-	Seminar/Assignment	7	4	7	100
	Total	35	26	35	650
	Semes	ster II			
MPA201T	Advanced Instrumental Analysis	4	4	4	100
MPA202T	Modern Bio-Analytical Techniques	4	4	4	100
MPA203T	Quality Control and Quality Assurance	4	4	4	100
MPA204T	Herbal and Cosmetic Analysis	4	4	4	100
MPA205P	Pharmaceutical Analysis Practical II	12	6	12	150
-	Seminar/Assignment	7	4	7	100
	Total	35	26	35	650

Table - 12: Course of study for M. Pharm. III Semester (Common for All Specializations)

Course Code	Course	Credit Hours	Credit Points
MRM 301T	Research Methodology and Biostatistics*	4	4
-	Journal club	1	1
-	Discussion / Presentation (Proposal Presentation)	2	2
-	Research Work	28	14
	Total	35	21

<sup>\*</sup> Non University Exam

Table - 13: Course of study for M. Pharm. IV Semester (Common for All Specializations)

Course Code	Course	Credit Hours	Credit Points
-	Journal Club	1	1
-	Research Work	31	16
-	Discussion/Final Presentation	3	3
	Total	35	20

Table - 14: Semester wise credits distribution

Table - 14. Semester wise credits distribution						
Semester	Credit Points					
I	26					
II	26					
III	21					
IV	20					
Co-curricular Activities (Attending Conference, Scientific Presentations and Other Scholarly Activities)	Minimum=02 Maximum=07*					
Total Credit Points	Minimum=95 Maximum=100*					

<sup>\*</sup>Credit Points for Co-curricular Activities

Table - 15: Guidelines for Awarding Credit Points for Co-curricular Activities

Name of the Activity	Maximum Credit Points Eligible / Activity
Participation in National Level Seminar/Conference/Workshop/Symposium/ Training Programs (related to the specialization of the student)	01
Participation in international Level Seminar/Conference/Workshop/Symposium/ Training Programs (related to the specialization of the student)	02
Academic Award/Research Award from State Level/National Agencies	01
Academic Award/Research Award from International Agencies	02
Research / Review Publication in National Journals (Indexed in Scopus / Web of Science)	01
Research / Review Publication in International Journals (Indexed in Scopus / Web of Science)	02

Note: International Conference: Held Outside India

International Journal: The Editorial Board Outside India

\*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 M. Pharm. programme shall have a Programme Committee constituted by the Head of the institution in consultation with all the Heads of the departments.
- 2. The composition of the Programme Committee shall be as follows: A teacher at the cadre of Professor shall be the Chairperson; One Teacher from eachM.Pharm specialization and four student representatives (two from each academic year), nominated by the Head of the institution.
- 3. Duties of the Programme 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 Programme Committee shall meet at least twice in a semester preferably at the end of each sessionalexam and before the end semester exam.

#### 11. Examinations/Assessments

The schemes for internal assessment and end semester examinations are given in Table – 16.

#### 11.1. End semester examinations

The End Semester Examinations for each theory and practical coursethrough semesters I to IVshall beconducted by the respective university except for the subject 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.

Tables – 1616 : Schemes for internal assessments and end semester
(Pharmaceutics- MPH)

		(Priarir	aceulic	:s- MPH)		_		
Course		Internal Assessment				End Semester Exams		Tota 1
Code	Course	Continu ous Mode		sional ams Durati on	Tot al	Mar ks	Durati on	Mar ks
		SE	EMESTE	R I				
MPH 101T	Modern Pharmaceuti cal Analytical Techniques	10	15	1 Hr	25	75	3 Hrs	100
MPH 102T	Drug Delivery System	10	15	1 Hr	25	75	3 Hrs	100
MPH 103T	Modern Pharmaceuti cs	10	15	1 Hr	25	75	3 Hrs	100
MPH 104T	Regulatory Affair	10	15	1 Hr	25	75	3 Hrs	100
MPH 105P	Pharmaceuti cs Practical I	20	30	6 Hrs	50	100	6 Hrs	150
-	Seminar /Assignment	-	-	-	-	-	-	100
			otal					650
		SE	MESTE	R II				
MPH 201T	Molecular Pharmaceuti cs(Nano Tech and Targeted DDS)	10	15	1 Hr	25	75	3 Hrs	100
MPH 202T	Advanced Biopharmac eutics & Pharmacokin etics	10	15	1 Hr	25	75	3 Hrs	100
MPH 203T	Computer Aided Drug Delivery System	10	15	1 Hr	25	75	3 Hrs	100
MPH	Cosmetic	10	15	1 Hr	25	75	3 Hrs	100

204T	and Cosmeceutic als							
MPH 205P	Pharmaceuti cs Practical I	20	30	6 Hrs	50	100	6 Hrs	150
-	Seminar /Assignment	-	-	-	-	-	-	100
Total							650	

al Chemistry Practical II						Hrs	
Seminar - /Assignment	-	-	-	-	-	-	100
Total						650	

Tables - 19: Schemes for internal assessments and end semester examinations (Pharmaceutical Analysis-MPA)

Course		Internal Assessment				End Semester Exams		Total
Code	Course	Contin uous Mode		sional ams Durati on	Tot al	Mark s	Dura tion	Marks
			SEMEST					
	Modern							
MPA101T	Pharmaceuti cal Analysis	10	15	1 Hr	25	75	3 Hrs	100
MPA102T	Advanced Pharmaceuti cal Analysis	10	15	1 Hr	25	75	3 Hrs	100
MPA103T	Pharmaceuti cal Validation	10	15	1 Hr	25	75	3 Hrs	100
MPA104T	Food Analysis	10	15	1 Hr	25	75	3 Hrs	100
MPA105P	Pharmaceuti cal Analysis- I	20	30	6 Hrs	50	100	6 Hrs	150
-	Seminar /Assignment	-	-	-	-	-	-	100
			otal					650
		,	SEMEST	ER II				
MPA201T	Advanced Instrumental Analysis	10	15	1 Hr	25	75	3 Hrs	100
MPA202T	Modern Bio- Analytical Techniques	10	15	1 Hr	25	75	3 Hrs	100
мРА203Т	Quality Control and Quality	10	15	1 Hr	25	75	3 Hrs	100

	Assurance							
MPA204T	Herbal and Cosmetic analysis	10	15	1 Hr	25	75	3 Hrs	100
MPA205P	Pharmaceuti cal Analysis- II	20	30	6 Hrs	50	100	6 Hrs	150
-	Seminar /Assignment	-	-	-	-	-	-	100
Total					650			

Tables - 26: Schemes for internal assessments and end semester examinations (Semester III&IV)

				seesement			emester	
Course Code		Internal Assessment			Exams		Tota	
	Course	Conti nuou s Mode	Sessional Exams		Tot	Mark	Durati	l Mark s
			Mark s	Durati on	al	s	on	
			SEMEST	ΓER III				
MRM30 1T	Research Methodology and Biostatistics*	10	15	1 Hr	25	75	3 Hrs	100
-	Journal club	-	-	-	25	-	-	25
-	Discussion / Presentation (Proposal Presentation)	-	-	-	50	-	-	50
-	Research work*	-	-	-	-	350	1 Hr	350
Total						525		
SEMESTER IV								
-	Journal club	-	-	-	25	-	-	25
-	Discussion / Presentation (Proposal Presentation)	-	-	-	75	-	-	75
-	Research work and Colloquium	-	-	-	-	400	1 Hr	400
Total					500			

<sup>\*</sup>Non University Examination

#### 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 - 27: Scheme for awarding internal assessment: Continuous mode

Theory				
Criteria	Maximum Marks			
Attendance (Refer Table – 28)	8			
Student - Teacher interaction	2			
Total	10			
Practical				
Attendance (Refer Table – 28	10			
Based on Practical Records, Regular viva voce, etc.	10			
Total	20			

Table - 28: Guidelines for the allotment of marks for attendance

Percentage of Attendance	Theory	Practical
95 – 100	8	10
90 – 94	6	7.5
85 – 89	4	5
80 – 84	2	2.5
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 in the table. The average marks of two sessional exams shall be computed for internal assessment as per the requirements given in tables.

#### 12. Promotion and award of grades

A student shall be declared PASS and eligible for getting grade in a course of M.Pharm.programme if he/she secures at least 50% marks in that particular courseincluding internal assessment.

#### 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 once in 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. Reexamination of end semester examinations

Reexamination of end semester examination shall be conducted as per the schedule given in table 29. The exact dates of examinations shall be notified from time to time.

Table - 29: Tentative schedule of end semester examinations

Semester	For Regular Candidates	For Failed Candidates
I and III	November / December	May / June
II and IV	May / June	November / December

#### 16. Allowed to keep terms (ATKT):

No student shall be admitted to any examination unless he/she fulfills the norms given in 6. ATKT rules are applicable as follows:

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

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

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

#### 17. Grading of performances

#### 17.1. 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 – 30.

Table – 30: Letter grades and grade points equivalent to Percentage of marks and performances

Percentage of Marks Obtained	Letter Grade	Grade Point	Performance
90.00 - 100	0	10	Outstanding
80.00 - 89.99	Α	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

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 and C4 and the student's grade points in these courses are G1, G2, G3 and G4, respectively, and then students' SGPA is equal to:

$$SGPA = C_1G_1 + C_2G_2 + C_3G_3 + C_4G_4$$

$$C_1 + C_2 + C_3 + C_4$$

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:

$$SGPA = \begin{array}{c} C_1G_1 + C_2G_2 + C_3G_3 + C_4* ZERO \\ \\ C_1 + C_2 + C_3 + C_4 \end{array}$$

#### 19. Cumulative Grade Point Average (CGPA)

The CGPA is calculated with the SGPA of all the IV semesters to two decimal points and is indicated in final grade report card/final transcript showing the grades of all IV semesters and their courses. The CGPA shall reflect the failed statusin case of F grade(s), till the course(s) is/are passed. When the course(s) is/are passedby 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:

$$CGPA = \begin{array}{c} C_1S_1 + C_2S_2 + C_3S_3 + C_4S_4 \\ \\ C_1 + C_2 + C_3 + C_4 \end{array}$$

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 to 7.49 Second Class = CGPA of 5.00 to 5.99

#### 21. Project work

All the students shall undertake a project under the supervision of a teacher in Semester III to IV and submit a report. 4 copies of the project report shall be submitted (typed & bound copy not less than 75 pages).

The internal and external examiner appointed by the University shall evaluate the project at the time of the Practical examinations of other semester(s). The projects shall be evaluated as per the criteria given below.

#### Evaluation of Dissertation Book:

Objective(s) of the work done	50 Marks
Methodology adopted	150 Marks
Results and Discussions	250 Marks
Conclusions and Outcomes	50 Marks

500 Marks	
	500 Marks

#### **Evaluation of Presentation:**

Presentation of work	100 Marks
Communication skills	50 Marks
Question and answer skills	100 Marks

#### 22. 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 M.Pharm program shall not be eligible for award of ranks. Moreover, the candidates should have completed the M. Pharm program in minimum prescribed number of years, (two years) for the award of Ranks.

#### 23. Award of degree

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

#### 24. 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 said period, otherwise they have to get fresh Registration.

#### 25. Revaluation I Retotaling of answer papers

There is no provision for revaluation of the answer papers in any examination. However, the candidates can apply for retotaling by paying prescribed fee.

#### 26. 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.

#### PHARMACEUTICS(MPH)

# MODERN PHARMACEUTICAL ANALYTICAL TECHNIQUES (MPH 101T)

#### Scope

This subject deals with various advanced analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are NMR, Mass spectrometer, IR, HPLC, GC etc.

#### Objectives

After completion of course student is able to know,

- Chemicals and Excipients
- The analysis of various drugs in single and combination dosage forms
- Theoretical and practical skills of the instruments

THEORY 60 HOURS

- a. UV-Visible spectroscopy: Introduction, Theory, Laws, 11
   Instrumentation associated with UV-Visible spectroscopy, Hrs
   Choice of solvents and solvent effect and Applications of UV Visible spectroscopy.
  - b. IR spectroscopy: Theory, Modes of Molecular vibrations, Sample handling, Instrumentation of Dispersive and Fourier -Transform IR Spectrometer, Factors affecting vibrational frequencies and Applications of IR spectroscopy
  - c. Spectroflourimetry: Theory of Fluorescence, Factors affecting fluorescence, Quenchers, Instrumentation and Applications of fluorescence spectrophotometer.
  - d. Flame emission spectroscopy and Atomic absorption spectroscopy: Principle, Instrumentation, Interferences and Applications.
- 2 NMR spectroscopy: Quantum numbers and their role in NMR, Principle, Instrumentation, Solvent requirement in NMR, Relaxation process, NMR signals in various compounds, Chemical shift, Factors influencing chemical shift, Spin-Spin coupling, Coupling constant, Nuclear magnetic double resonance, Brief outline of principles of FT-NMR and 13C NMR. Applications of NMR spectroscopy.

11 Hrs

- 3 Mass Spectroscopy: Principle, Theory, Instrumentation of Mass 11 Spectroscopy, Different types of ionization like electron impact, Hrs chemical, field, FAB and MALDI, APCI, ESI, APPI Analyzers of Quadrupole and Time of Flight, Mass fragmentation and its rules, Meta stable ions, Isotopic peaks and Applications of Mass spectroscopy
- 4 Chromatography: Principle, apparatus, instrumentation, 11 chromatographic parameters, factors affecting resolution and Hrs applications of the following:
  - a) Paper chromatography b) Thin Layer chromatography
  - c) Ion exchange chromatography d) Column chromatography
  - e) Gas chromatography f) High Performance Liquid chromatography
  - g) Affinity chromatography
- 5 a. Electrophoresis: Principle, Instrumentation, Working 11 conditions, factors affecting separation and applications of the Hrs following:
  - a) Paper electrophoresis b) Gel electrophoresis c) Capillary electrophoresis d) Zone electrophoresis e) Moving boundary electrophoresis f) Iso electric focusing
  - b. X ray Crystallography: Production of X rays, Different X ray diffraction methods, Bragg's law, Rotating crystal technique, X ray powder technique, Types of crystals and applications of X-ray diffraction.
- 6 Immunological assays : RIA (Radio immuno assay), ELISA, 5 Hrs Bioluminescence assays.

- 1. Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John Wiley & Sons, 2004.
- 2. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5th edition, Eastern press, Bangalore, 1998.
- 3. Instrumental methods of analysis Willards, 7th edition, CBS publishers.
- 4. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4th edition, CBS Publishers, New Delhi, 1997.
- 5. Organic Spectroscopy William Kemp, 3rd edition, ELBS, 1991.
- 6. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3rd Edition, CBS Publishers, New Delhi, 1997.
- 7. Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11. Marcel Dekker Series

# DRUG DELIVERY SYSTEMS (MPH 102T)

#### SCOPE

This course is designed to impart knowledge on the area of advances in novel drug delivery systems.

#### **OBJECTIVES**

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

The various approaches for development of novel drug delivery systems.

The criteria for selection of drugs and polymers for the development of delivering system

The formulation and evaluation of Novel drug delivery systems..

THEORY 60 Hrs

- 1. Release(SR) and Controlled Release Sustained (CR) formulations: Introduction & basic concepts, advantages disadvantages, factors influencing, Physicochemical & biological approaches for SR/CR formulation, Mechanism of Drug Delivery from SR/CR formulation. Polymers: introduction, definition, classification, properties and application Dosage Forms for Personalized Medicine: Introduction. Definition. Pharmacogenetics, Categories of Patients for Personalized Medicines: Customized drug delivery systems, Bioelectronic Medicines, 3D printing of pharmaceuticals, Telepharmacy.
- 2 Rate Controlled Drug Delivery Systems: Principles & 10 Fundamentals, Types, Activation; Modulated Drug Delivery Hrs Systems; Mechanically activated, pH activated, Enzyme activated, and Osmotic activated Drug Delivery Systems Feedback regulated Drug Delivery Systems; Principles & Fundamentals.
- 3 Gastro-Retentive Drug Delivery Systems: Principle, concepts 10 advantages and disadvantages, Modulation of GI transit time Hrs approaches to extend GI transit. Buccal Drug Delivery Systems: Principle of muco adhesion, advantages and disadvantages, Mechanism of drug permeation, Methods of formulation and its evaluations.
- Occular Drug Delivery Systems: Barriers of drug permeation, Methods to overcome barriers.

- 5 Transdermal Drug Delivery Systems: Structure of skin and 10 barriers, Penetration enhancers, Transdermal Drug Delivery Hrs Systems, Formulation and evaluation.
- 6 Protein and Peptide Delivery: Barriers for protein delivery. 08 Formulation and Evaluation of delivery systems of proteins and Hrs other macromolecules.
- Vaccine delivery systems: Vaccines, uptake of antigens, single shot vaccines, mucosal and transdermal delivery of vaccines.
   Hrs

#### REFERENCES

1. Y W. Chien, Novel Drug Delivery Systems, 2nd edition, revised and expanded,

Marcel Dekker, Inc., New York, 1992.

- 2. Robinson, J. R., Lee V. H. L, Controlled Drug Delivery Systems, Marcel Dekker, Inc., New York, 1992.
- 3. Encyclopedia of controlled delivery, Editor- Edith Mathiowitz, Published by WileyInterscience Publication, John Wiley and Sons, Inc, New York! Chichester/Weinheim
- 4. 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, Vallabh Prakashan, New Delhi, First edition 2002

#### **JOURNALS**

- 1. Indian Journal of Pharmaceutical Sciences (IPA)
- 2. Indian drugs (IDMA)
- 3. Journal of controlled release (Elsevier Sciences) desirable
- 4. Drug Development and Industrial Pharmacy (Marcel & Decker) desirable

### MODERN PHARMACEUTICS (MPH 103T)

#### Scope

Course designed to impart advanced knowledge and skills required to learn various aspects and concepts at pharmaceutical industries

#### Objectives

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

- The elements of preformulation studies.
- The Active Pharmaceutical Ingredients and Generic drug Product development
- Industrial Management and GMP Considerations.
- Optimization Techniques & Pilot Plant Scale Up Techniques
- Stability Testing, sterilization process & packaging of dosage forms.

#### THEORY 60 HRS

- a. Preformation Concepts Drug Excipient interactions 10 different methods, kinetics of stability, Stability testing. Theories of Hrs dispersion and pharmaceutical Dispersion (Emulsion and Suspension, SMEDDS) preparation and stability Large and small volume parental physiological and formulation consideration, Manufacturing and evaluation.
  - b. Optimization techniques in Pharmaceutical Formulation: Concept and parameters of optimization, Optimization techniques in pharmaceutical formulation and processing. Statistical design, Response surface method, Contour designs, Factorial designs and application in formulation

10

- Validation: Introduction to Pharmaceutical Validation, Scope & 10 merits of Validation, Validation and calibration of Master plan, Hrs ICH & WHO guidelines for calibration and validation of equipments, Validation of specific dosage form, Types of validation. Government regulation, Manufacturing Process Model, URS, DQ, IQ, OQ & P.Q. of facilities.
- 3 cGMP & Industrial Management: Objectives and policies of 10 current good manufacturing practices, layout of buildings, services, equipments and their maintenance Production management: Production organization, , materials management, handling and transportation, inventory management and control, production and planning control, Sales forecasting, budget and cost control, industrial and personal relationship. Concept of Total Quality Management.

- 4 Compression and compaction: Physics of tablet compression, 10 compression, consolidation, effect of friction, distribution of Hrs forces, compaction profiles. Solubility.
- 5 Study of consolidation parameters; Diffusion parameters, 10 Dissolution parameters and Pharmacokinetic parameters, Heckel Hrs plots, Similarity factors f2 and f1, Higuchi and Peppas plot, Linearity Concept of significance, Standard deviation, Chi square test, students T-test, ANOVA test.

- 1. Theory and Practice of Industrial Pharmacy By Lachmann and Libermann
- 2. Pharmaceutical dosage forms: Tablets Vol. 1-3 by Leon Lachmann.
- 3. Pharmaceutical Dosage forms: Disperse systems, Vol, 1-2; By Leon Lachmann.
- 4. Pharmaceutical Dosage forms: Parenteral medications Vol. 1-2; By Leon Lachmann.
- 5. Modern Pharmaceutics; By Gillbert and S. Banker.
- 6. Remington's Pharmaceutical Sciences.
- Advances in Pharmaceutical Sciences Vol. 1-5; By H.S. Bean & A.H. Beckett.
- 8. Physical Pharmacy; By Alfred martin
- 9. Bentley's Textbook of Pharmaceutics by Rawlins.
- 10. Good manufacturing practices for Pharmaceuticals: A plan for total quality control, Second edition; By Sidney H. Willig.
- 11. Quality Assurance Guide; By Organization of Pharmaceutical producers of India.
- 12.Drug formulation manual; By D.P.S. Kohli and D.H.Shah. Eastern publishers, New Delhi.
- 13. How to practice GMPs; By P.P.Sharma. Vandhana Publications, Agra.
- 14. Pharmaceutical Process Validation; By Fra. R. Berry and Robert A. Nash.
- 15. Pharmaceutical Preformulations; By J.J. Wells.
- 16. Applied production and operations management; By Evans, Anderson, Sweeney and Williams.
- 17. Encyclopaedia of Pharmaceutical technology, Vol I III.

# REGULATORY AFFAIRS (MPH 104T)

#### Scope

Course designed to impart advanced knowledge and skills required to learn the concept of generic drug and their development, various regulatory filings in different countries, different phases of clinical trials and submitting regulatory documents: filing process of IND, NDA and ANDA

- To know the approval process of
- To know the chemistry, manufacturing controls and their regulatory importance
- To learn the documentation requirements for
- To learn the importance and

#### Objectives:

Upon completion of the course, it is expected that the students will be able to understand

- The Concepts of innovator and generic drugs, drug development process
- The Regulatory guidance's and guidelines for filing and approval process
- Preparation of Dossiers and their submission to regulatory agencies in different countries
- Post approval regulatory requirements for actives and drug products
- Submission of global documents in CTD/ eCTD formats
- Clinical trials requirements for approvals for conducting clinical trials
- Pharmacovigilence and process of monitoring in clinical trials.

THEORY 60 Hrs

- 1. a. Documentation in Pharmaceutical industry: Master 12 formula record, DMF (Drug Master File), distribution records. Hrs Generic drugs product development Introduction , Hatch-Waxman act and amendments, CFR (CODE OF FEDERAL REGULATION) ,drug product performance, in-vitro, ANDA regulatory approval process, NDA approval process, BE and drug product assessment, in -vivo, scale up process approval changes, post marketing surveillance, outsourcing BA and BE to CRO.
  - b. Regulatory requirement for product approval: API, biologics, novel, therapies obtaining NDA, ANDA for generic drugs ways and means of US registration for foreign drugs

- 2 CMC, post approval regulatory affairs. Regulation for combination 12 products and medical devices.CTD and ECTD format, industry Hrs and FDA liaison. ICH Guidelines of ICH-Q, S E, M. Regulatory requirements of EU, MHRA, TGA and ROW countries.
- 3 Non clinical drug development: Global submission of IND, 12 NDA, ANDA. Investigation of medicinal products dossier, dossier Hrs (IMPD) and investigator brochure (IB).
- 4 Clinical trials: Developing clinical trial protocols. Institutional 12 review board/ independent ethics committee Formulation and Hrs working procedures informed Consent process and procedures. HIPAA- new, requirement to clinical study process, pharmacovigilance safety monitoring in clinical trials.

- 1. Generic Drug Product Development, Solid Oral Dosage forms, Leon Shargel and IsaderKaufer, Marcel Dekker series, Vol.143
- 2. The Pharmaceutical Regulatory Process, Second Edition Edited by Ira R. Berry and Robert P.Martin, Drugs and the Pharmaceutical Sciences, Vol. 185, Informa Health care Publishers.
- 3. New Drug Approval Process: Accelerating Global Registrations By Richard A Guarino, MD,5th edition, Drugs and the Pharmaceutical Sciences, Vol. 190.
- 4. Guidebook for drug regulatory submissions / Sandy Weinberg. By John Wiley & Sons.Inc.
- 5. FDA regulatory affairs: a guide for prescription drugs, medical devices, and biologics/edited By Douglas J. Pisano, David Mantus.
- 6. Clinical Trials and Human Research: A Practical Guide to Regulatory Compliance By Fay A.Rozovsky and Rodney K. Adams
- 7. www.ich.org/
- 8. www.fda.gov/
- 9. europa.eu/index\_en.htm
- 10. https://www.tga.gov.au/tga-basics

#### PHARMACEUTICS PRACTICALS - I (MPH 105P)

- Analysis of pharmacopoeial compounds and their formulations by UV Vis spectrophotometer
- Simultaneous estimation of multi component containing formulations by UV spectrophotometry
- 3. Experiments based on HPLC
- 4. Experiments based on Gas Chromatography
- 5. Estimation of riboflavin/quinine sulphate by fluorimetry
- 6. Estimation of sodium/potassium by flame photometry
- 7. To perform In-vitro dissolution profile of CR/SR marketed formulation
- 8. Formulation and evaluation of sustained release matrix tablets
- 9. Formulation and evaluation osmotically controlled DDS
- 10. Preparation and evaluation of Floating DDS- hydro dynamically balanced DDS
- 11. Formulation and evaluation of Muco adhesive tablets.
- 12. Formulation and evaluation of trans dermal patches.
- 13. To carry out preformulation studies of tablets.
- 14. To study the effect of compressional force on tablets disintegration time.
- 15. To study Micromeritic properties of powders and granulation.
- 16. To study the effect of particle size on dissolution of a tablet.
- 17. To study the effect of binders on dissolution of a tablet.
- 18. To plot Heckal plot, Higuchi and peppas plot and determine similarity factors.

# MOLECULAR PHARMACEUTICS (NANO TECHNOLOGY & TARGETED DDS) (NTDS) (MPH 201T)

#### Scope

This course is designed to impart knowledge on the area of advances in novel drug delivery systems.

#### **Objectives**

Upon completion of the course student shall be able to understand

- The various approaches for development of novel drug delivery systems.
- The criteria for selection of drugs and polymers for the development of NTDS
- The formulation and evaluation of novel drug delivery systems.

THEORY 60 Hrs

- 1. Targeted Drug Delivery Systems: Concepts, Events and 12 biological process involved in drug targeting. Tumor targeting and Hrs Brain specific delivery.
- 2 Targeting Methods: introduction preparation and evaluation. 12 Nano Particles & Liposomes: Types, preparation and evaluation. Hrs
- 3 Micro Capsules / Micro Spheres: Types, preparation and 12 evaluation, Monoclonal Antibodies; preparation and application, Hrs preparation and application of Niosomes, Aquasomes, Phytosomes, Electrosomes.
- 4 Pulmonary Drug Delivery Systems : Aerosols, propellents, 12 ContainersTypes, preparation and evaluation, Intra Nasal Route Hrs Delivery systems; Types, preparation and evaluation.
- Nucleic acid based therapeutic delivery system: Gene therapy, 12 introduction (ex-vivo & in-vivo gene therapy). Potential target diseases for gene therapy (inherited disorder and cancer). Gene expression systems (viral and nonviral gene transfer). Liposomal gene delivery systems.

Biodistribution and Pharmacokinetics. knowledge of therapeutic antisense molecules and aptamers as drugs of future.

- 1. Y W. Chien, Novel Drug Delivery Systems, 2nd edition, revised and expanded, Marcel Dekker, Inc., New York, 1992.
- 2. S.P.Vyas and R.K.Khar, Controlled Drug Delivery concepts and advances, Vallabh Prakashan, New Delhi, First edition 2002.
- 3. N.K. Jain, Controlled and Novel Drug Delivery, CBS Publishers & Distributors, NewDelhi, First edition 1997 (reprint in 2001).

# ADVANCED BIOPHARMACEUTICS & PHARMACOKINETICS (MPH 202T)

#### Scope

This course is designed to impart knowledge and skills necessary for dose calculations, dose adjustments and to apply biopharmaceutics theories in practical problem solving. Basic theoretical discussions of the principles of biopharmaceutics and pharmacokinetics are provided to help the students' to clarify the concepts.

#### Objectives

Upon completion of this course it is expected that students will be able understand.

- The basic concepts in biopharmaceutics and pharmacokinetics.
- The use raw data and derive the pharmacokinetic models and parameters the best describe the process of drug absorption, distribution, metabolism and elimination.
- The critical evaluation of biopharmaceutic studies involving drug product equivalency.
- The design and evaluation of dosage regimens of the drugs using pharmacokinetic and biopharmaceutic parameters.
- The potential clinical pharmacokinetic problems and application of basics of pharmacokinetic

THEORY 60 Hrs

12 1. Drug Absorption from the Gastrointestinal Tract: Gastrointestinal tract, Mechanism of drug absorption, Factors affecting drug absorption, pH-partition theory of drug absorption. Formuulation and physicochemical factors: Dissolution rate, Dissolution Noves-Whitney equation and drug process, dissolution, Factors affecting the dissolution rate. Gastrointestinal absorption: role of the dosage form: Solution (elixir, syrup and solution) as a dosage form, Suspension as a dosage form, Capsule as a dosage form, Tablet as a dosage form, Dissolution methods ,Formulation and processing factors, Correlation of in vivo data with in vitro dissolution data. Transport model: Permeability-Solubility-Charge State and the pH Partition Hypothesis, Properties of the Gastrointestinal Tract (GIT), pH Microclimate Intracellular Hq Environment. Tight-Junction Complex.

2 Biopharmaceutic considerations in drug product design Vitro Drug Product Performance: Introduction. biopharmaceutic factors affecting drug bioavailability, rate-limiting steps in drug absorption, physicochemical nature of the drug formulation factors affecting drug product performance, in vitro: dissolution and drug release testing, compendial methods of dissolution, alternative methods of dissolution testing, meeting dissolution requirements, problems of variable control in dissolution testingperformance of drug products. In vitro-in vivo correlation, dissolution profile comparisons, drua product stability, considerations in the design of a drug product.

12 Hrs

12

12

Hrs

12

Hrs

- 3 Pharmacokinetics: Basic considerations. pharmacokinetic models, compartment modeling; one compartment model- IV bolus, IV infusion, extra-vascular. Multi compartment model:two compartment - model in brief, non-linear pharmacokinetics: cause of non-linearity, Michaelis - Menten equation, estimation of k<sub>max</sub> and v<sub>max</sub>. Drug interactions: introduction, the effect of proteininteractions.the effect οf bindina tissue-binding interactions.cvtochrome p450-based drua interactions.drug interactions linked to transporters.
- 4 Drug Product Performance, In Vivo: Bioavailability and Bioequivalence: drug product performance. purpose bioavailability studies, relative and absolute availability, methods for assessing bioavailability, bioequivalence studies, design and evaluation of bioequivalence studies, study designs, crossover study designs, evaluation of the data, bioequivalence example, study submission and drug review process. biopharmaceutics classification system, methods. Permeability: In-vitro, in-situ and In-vivo methods.generic biologics (biosimilar products), clinical significance of bioequivalence studies, special concerns in bioavailability and bioequivalence studies, generic substitution.
- Application of Pharmacokinetics: Modified-Release Drug Products, Targeted Drug Delivery Systems and Biotechnological Products. Introduction to Pharmacokinetics and pharmacodynamic, drug interactions. Pharmacokinetics and pharmacodynamics of biotechnology drugs. Introduction, Proteins and peptides, Monoclonal antibodies, Oligonucleotides, Vaccines (immunotherapy), Gene therapies.

- 1. Biopharmaceutics and Clinical Pharmacokinetics by Milo Gibaldi, 4th edition, Philadelphia, Lea and Febiger, 1991
- 2. Biopharmaceutics and Pharmacokinetics, A. Treatise, D.M. Brahmankar and Sunil B. Jaiswal., VallabPrakashan, Pitampura, Delhi
- 3. Applied Biopharmaceutics and Pharmacokinetics by Shargel. Land YuABC, 2<sup>nd</sup>edition, Connecticut Appleton Century Crofts, 1985
- 4. Textbook of Biopharmaceutics and Pharmacokinetics, Dr. Shobha Rani R. Hiremath, Prism Book
- Pharmacokinetics by Milo Gibaldi and D. Perrier, 2nd edition, Marcel Dekker Inc., New York, 1982
- 6. Current Concepts in Pharmaceutical Sciences: Biopharmaceutics, Swarbrick. J., Leaand Febiger, Philadelphia, 1970
- Clinical Pharmacokinetics, Concepts and Applications 3rd edition by MalcolmRowland and Thom~ N. Tozer, Lea and Febiger, Philadelphia, 1995
- 8. Dissolution, Bioavailability and Bioequivalence, Abdou. H.M, Mack PublishingCompany, Pennsylvania 1989
- 9. Biopharmaceutics and Clinical Pharmacokinetics, An Introduction, 4th edition, revised and expande by Robert. E. Notari, Marcel Dekker Inc, New York and Basel, 1987.
- Biopharmaceutics and Relevant Pharmacokinetics by John. G Wagner and M.Pemarowski, 1st edition, Drug Intelligence Publications, Hamilton, Illinois, 1971.
- 11. Encyclopedia of Pharmaceutical Technology, Vol 13, James Swarbrick, James. G.Boylan, Marcel Dekker Inc, New York, 1996.
- 12. Basic Pharmacokinetics, 1 st edition, Sunil S Jambhekarand Philip J Breen, pharmaceutical press, RPS Publishing, 2009.
- 13. Absorption and Drug Development- Solubility, Permeability, and Charge State, Alex Avdeef, John Wiley & Sons, Inc, 2003.

# COMPUTER AIDED DRUG DEVELOPMENT (MPH 203T)

#### Scope

This course is designed to impart knowledge and skills necessary for computer Applications in pharmaceutical research and development who want to understand the application of computers across the entire drug research and development process. Basic theoretical discussions of the principles of more integrated and coherent use of computerized information (informatics) in the drug development process are provided to help the students to clarify the concepts.

#### Objectives

Upon completion of this course it is expected that students will be able to understand,

- History of Computers in Pharmaceutical Research and Development
- Computational Modeling of Drug Disposition
- Computers in Preclinical Development
- Optimization Techniques in Pharmaceutical Formulation
- Computers in Market Analysis
- Computers in Clinical Development
- Artificial Intelligence (AI) and Robotics
- Computational fluid dynamics(CFD)

THEORY 60 Hrs

- 1. a. Computers in Pharmaceutical Research and 12 Development: A General Overview: History of Computers in Hrs Pharmaceutical Research and Development. Statistical modeling in Pharmaceutical research and development: Descriptive versus Statistical Mechanistic Modeling, Parameters. Estimation. Confidence Regions, Nonlinearity at the Optimum, Sensitivity Analysis, Optimal Design, Population Modeling b. Quality-by-Design In Pharmaceutical Development: Introduction, ICH Q8 guideline, Regulatory and industry views on ObD. Scientifically based ObD - examples of application.
- Computational Modeling Of Drug Disposition: Introduction 12 ,Modeling Techniques: Drug Absorption, Solubility, Intestinal Hrs Permeation, Drug Distribution ,Drug Excretion, Active Transport; P-gp, BCRP, Nucleoside Transporters, hPEPT1, ASBT, OCT, OATP, BBB-Choline Transporter.

3 Computer-aided formulation development:: Concept of 12 Hrs optimization, Optimization parameters, Factorial design, Optimization technology & Screening design. Computers in Pharmaceutical Formulation: Development of pharmaceutical emulsions, microemulsion drug carriers Legal Protection of Innovative Uses of Computers in R&D, The Ethics of Computing in Pharmaceutical Research, Computers in Market analysis

12

- 4 a. Computer-aided biopharmaceutical characterization:
  Gastrointestinal absorption simulation. Introduction, Theoretical background, Model construction, Parameter sensitivity analysis, Virtual trial, Fed vs. fasted state, In vitro dissolution and in vitro-in vivo correlation, Biowaiver considerations
  - b. Computer Simulations in Pharmacokinetics and Pharmacodynamics: Introduction, Computer Simulation: Whole Organism, Isolated Tissues, Organs, Cell, Proteins and Genes.
  - c. Computers in Clinical Development: Clinical Data Collection and Management, Regulation of Computer Systems
- 5 Artificial Intelligence (AI), Robotics and Computational fluid dynamics: General overview, Pharmaceutical Automation, Hrs Pharmaceutical applications, Advantages and Disadvantages. Current Challenges and Future Directions.

- 1. Computer Applications in Pharmaceutical Research and Development, Sean Ekins, 2006, John Wiley & Sons.
- 2. Computer-Aided Applications in Pharmaceutical Technology, 1<sup>st</sup> Edition, Jelena Djuris, Woodhead Publishing
- 3. Encyclopedia of Pharmaceutical Technology, Vol 13, James Swarbrick, James. G.Boylan, Marcel Dekker Inc, New York, 1996.

# COSMETICS AND COSMECEUTICALS (MPH 204T)

## Scope

This course is designed to impart knowledge and skills necessary forthefundamental need for cosmetic and cosmeceutical products.

### **Objectives**

Upon completion of the course, the students shall be able to understand

- Key ingredients used in cosmetics and cosmeceuticals.
- Key building blocks for various formulations.
- Current technologies in the market
- Various key ingredients and basic science to develop cosmetics and cosmeceuticals
- Scientific knowledge to develop cosmetics and cosmeceuticals with desired Safety, stability, and efficacy.

THEORY 60 Hrs

- Cosmetics Regulatory: Definition of cosmetic products as per 12 Indian regulation. Indian regulatory requirements for labeling of Hrs cosmetics Regulatory provisions relating to import of cosmetics., Misbranded and spurious cosmetics. Regulatory provisions relating to manufacture of cosmetics Conditions for obtaining license, prohibition of manufacture and sale of certain cosmetics, loan license, offences and penalties.
- Cosmetics Biological aspects: Structure of skin relating to 12 problems like dry skin, acne, pigmentation, prickly heat, wrinkles and body odor. Structure of hair and hair growth cycle. Common problems associated with oral cavity. Cleansing and care needs for face, eye lids, lips, hands, feet, nail, scalp, neck, body and under-arm.
- Formulation Building blocks: Building blocks for different 12 product formulations of cosmetics/cosmeceuticals. Surfactants Hrs Classification and application. Emollients, rheological additives: classification and application. Antimicrobial used as preservatives, their merits and demerits. Factors affecting microbial preservative efficacy. Building blocks for formulation of a moisturizing cream, vanishing cream, cold cream, shampoo and toothpaste. Soaps and syndetbars.

Perfumes; Classification of perfumes. Perfume ingredients listed as allergens in EU regulation.

- Controversial ingredients: Parabens, formaldehyde liberators, dioxane.
- 4 Design of cosmeceutical products: Sun protection, sunscreens 12 classification and regulatory aspects. Addressing dry skin, acne, Hrs sun-protection, pigmentation, prickly heat, wrinkles, body odor., dandruff, dental cavities, bleeding gums, mouth odor and sensitive teeth through cosmeceutical formulations.
- 5 Herbal Cosmetics: Herbal ingredients used in Hair care, skin 12 care and oral care. Review of guidelines for herbal cosmetics by Hrs private bodies like cosmos with respect to preservatives, emollients, foaming agents, emulsifiers and rheology modifiers. Challenges in formulating herbal cosmetics.

- 1. Harry's Cosmeticology. 8th edition.
- 2. Poucher'sperfumecosmeticsandSoaps,10<sup>th</sup> edition.
- Cosmetics Formulation, Manufacture and quality control, PP.Sharma,4<sup>th</sup> edition
- 4. Handbook of cosmetic science and Technology A.O.Barel, M.Paye and H.I. Maibach. 3 <sup>rd</sup> edition
- 5. Cosmetic and Toiletries recent suppliers catalogue.
- 6. CTFA directory.

# PHARMACEUTICS PRACTICALS - II (MPH 205P)

- 1. To study the effect of temperature change, non solvent addition, incompatible polymer addition in microcapsules preparation
- 2. Preparation and evaluation of Alginate beads
- 3. Formulation and evaluation of gelatin /albumin microspheres
- 4. Formulation and evaluation of liposomes/niosomes
- 5. Formulation and evaluation of spherules
- 6. Improvement of dissolution characteristics of slightly soluble drug by Solid dispersion technique.
- 7. Comparison of dissolution of two different marketed products /brands
- 8. Protein binding studies of a highly protein bound drug & poorly protein bound drug
- 9. Bioavailability studies of Paracetamol in animals.
- 10. Pharmacokinetic and IVIVC data analysis by Winnoline<sup>R</sup> software
- 11. In vitro cell studies for permeability and metabolism
- 12. DoE Using Design Expert® Software
- 13. Formulation data analysis Using Design Expert Software
- 14. Quality-by-Design in Pharmaceutical Development
- 15. Computer Simulations in Pharmacokinetics and Pharmacodynamics
- 16. Computational Modeling Of Drug Disposition
- 17. To develop Clinical Data Collection manual
- 18. To carry out Sensitivity Analysis, and Population Modeling.
- 19. Development and evaluation of Creams
- 20. Development and evaluation of Shampoo and Toothpaste base
- 21. To incorporate herbal and chemical actives to develop products
- 22. To address Dry skin, acne, blemish, Wrinkles, bleeding gums and dandruff

## PHARMACEUTICALANALYSIS(MPA)

# MODERN PHARMACEUTICAL ANALYTICAL TECHNIQUES (MPA 101T)

### Scope

This subject deals with various advanced analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are NMR, Mass spectrometer, IR, HPLC, GC etc.

### **Objectives**

After completion of course student is able to know about chemicals and excipients

- The analysis of various drugs in single and combination dosage forms
- Theoretical and practical skills of the instruments

THEORY 60 Hrs

- 1. a. UV-Visible spectroscopy: Introduction, Theory, Laws, 10 Instrumentation associated with UV-Visible spectroscopy, Choice Hrs of solvents and solvent effect and Applications of UV-Visible spectroscopy, Difference/ Derivative spectroscopy.
  - b. IR spectroscopy: Theory, Modes of Molecular vibrations, Sample handling, Instrumentation of Dispersive and Fourier Transform IR Spectrometer, Factors affecting vibrational frequencies and Applications of IR spectroscopy, Data Interpretation.
  - c. Spectroflourimetry: Theory of Fluorescence, Factors affecting fluorescence (Characterestics of drugs that can be analysed by flourimetry), Quenchers, Instrumentation and Applications of fluorescence spectrophotometer.
  - d. Flame emission spectroscopy and Atomic absorption spectroscopy: Principle, Instrumentation, Interferences and Applications.

10

Hrs

- 2 NMR spectroscopy: Quantum numbers and their role in NMR, Principle, Instrumentation, Solvent requirement in NMR, Relaxation process, NMR signals in various compounds, Chemical shift, Factors influencing chemical shift, Spin-Spin coupling, Coupling constant, Nuclear magnetic double resonance, Brief outline of principles of FT-NMR and 13C NMR. Applications of NMR spectroscopy.
- 3 Mass Spectroscopy: Principle, Theory, Instrumentation of Mass 10

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

- 4 Chromatography: Principle. apparatus, instrumentation. 10 chromatographic parameters, factors affecting resolution, isolation of drug from excipients, data interpretation and applications of the following:
  - a. Thin Layer chromatography
  - b. High Performance Thin Layer Chromatography
  - c. Ion exchange chromatography
  - d. Column chromatography
  - e. Gas chromatography
  - f. High Performance Liquid chromatography
  - g. Ultra High Performance Liquid chromatography
  - h. Affinity chromatography
  - i. Gel Chromatography
- 5 Principle. a. Electrophoresis: Instrumentation. Working 10 conditions, factors affecting separation and applications of the following:
  - a) Paper electrophoresis b) Gel electrophoresis c) Capillary electrophoresis d) Zone electrophoresis e) Moving boundary electrophoresis f) Iso electric focusing
  - b. X ray Crystallography: Production of X rays, Different X ray methods, Bragg's law, Rotating crystal technique, X ray powder technique, Types of crystals and applications of X-ray diffraction
- 6 Potentiometry: Principle, working, Ion selective Electrodes and 10 Application of potentiometry. Hrs

Techniques: Principle, thermal Thermal transitions Instrumentation (Heat flux and power-compensation and designs), Modulated DSC, Hyper DSC, experimental parameters (sample preparation, experimental conditions, calibration, heating and cooling rates, resolution, source of errors) and their influence, advantage and disadvantages, pharmaceutical applications. Differential Thermal Analysis (DTA): Principle, instrumentation

and advantage and disadvantages, pharmaceutical applications, derivative differential thermal analysis (DDTA). TGA: Principle, instrumentation, factors affecting results, advantage and disadvantages, pharmaceutical applications.

- 1. Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John Wiley & Sons, 2004.
- 2. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5<sup>th</sup> edition, Eastern press, Bangalore, 1998.
- 3. Instrumental methods of analysis Willards, 7th edition, CBS publishers.
- 4. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4th edition, CBS Publishers, New Delhi, 1997.
- 5. Organic Spectroscopy William Kemp, 3rd edition, ELBS, 1991.
- 6. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3rd Edition, CBS Publishers, New Delhi, 1997.
- 7. Pharmaceutical Analysis Modern Methods Part B J W Munson, Vol 11, Marcel. Dekker Series
- 8. Spectroscopy of Organic Compounds, 2<sup>nd</sup> edn., P.S/Kalsi, Wiley estern Ltd., Delhi.
- 9. Textbook of Pharmaceutical Analysis, KA.Connors, 3<sup>rd</sup> Edition, John Wiley & Sons, 1982.

# ADVANCED PHARMACEUTICAL ANALYSIS (MPA 102T)

## Scope

This subject deals with the various aspects of Impurity, Impurities in new drug products, in residual solvents, Elemental impurities, Impurity profiling and characterization of degradents, Stability testing of phytopharmaceuticals and their protocol preparation. It also covers the biological testing of various vaccines and their principle and procedure.

## Objective

After completion of the course students shall able to know,

- Appropriate analytical skills required for the analytical method development.
- Principles of various reagents used in functional group analysis that renders necessary support in research methodology and demonstrates its application in the practical related problems.
- Analysis of impurities in drugs, residual solvents and stability studies of drugs and biological products

THEORY 60 Hrs

Impurity and stability studies: 10
 Definition, classification of impurities in drug Substance or Active Pharmaceutical Ingredients and quantification of impurities as per

Impurities in new drug products:

ICH auidelines

Rationale for the reporting and control of degradation products, reporting degradation products content of batches, listing of degradation products in specifications, qualification of degradation products

Impurities in residual solvents:

General principles, classification of residual solvents, Analytical procedures, limits of residual solvents, reporting levels of residual solvents

2 Elemental impurities: 10
Element classification, control of elemental impurities, Potential Hrs
Sources of elemental Impurities, Identification of Potential
Elemental Impurities, analytical procedures, instrumentation & C,
H, N and S analysis

## Stability testing protocols:

Selection of batches, container orientation, test parameters, sampling frequency, specification, storage conditions, recording of results, concept of stability, commitment etc. Important mechanistic and stability related information provided by results of study of factors like temperature, pH, buffering species ionic strength and dielectric constant etc. on the reaction rates. With practical considerations.

- Impurity profiling and degradent characterization: Method 10 development, Stability studies and concepts of validation Hrs accelerated stability testing & shelf life calculation, WHO and ICH stability testing guidelines, Stability zones, steps in development, practical considerations. Basics of impurity profiling and degradent characterization with special emphasis. Photostability testing guidelines, ICH stability guidelines for biological products
- 4 Stability testing of phytopharmaceuticals: 10 Regulatory requirements, protocols, HPTLC/HPLC finger printing, interactions and complexity.
- Biological tests and assays of the following:

  a. Adsorbed Tetanus vaccine b. Adsorbed Diphtheria vaccine Hrs

  c. Human anti haemophilic vaccine d. Rabies vaccine e.

  Tetanus Anti toxin f. Tetanus Anti serum g. Oxytocin h.

  Heparin sodium IP i. Antivenom. PCR, PCR studies for gene regulation, instrumentation (Principle and Procedures)
- 6 Immunoassays (IA) 10
  Basic principles, Production of antibodies, Separation of bound Hrs
  and unbound drug, Radioimmunoassay, Optical IA, Enzyme IA,
  Fluoro IA, Luminiscence IA, Quantification and applications of IA.

- Vogel's textbook of quantitative chemical analysis Jeffery J Bassett, J. Mendham, R. C. Denney, 5<sup>th</sup> edition, ELBS, 1991.
- 2. Practical Pharmaceutical Chemistry Beckett and Stenlake, Vol II, 4th Edition, CBS publishers, New Delhi, 1997.
- 3. Textbook of Pharmaceutical Analysis K A Connors, 3rd Edition, John Wiley & Sons, 1982.

- 4. Pharmaceutical Analysis Higuchi, Brochmman and Hassen, 2nd Edition, Wiley Inter science Publication, 1961.
- 5. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3rd Edition, CBS Publishers New Delhi, 1997.
- Pharmaceutical Analysis- Modern methods J W Munson Part B, Volume 11, Marcel Dekker Series.
- 7. The Quantitative analysis of Drugs D C Carratt, 3rd edition, CBS Publishers, NewDelhi, 1964.
- 8. Indian Pharmacopoeia Vol I, II & III 2007, 2010, 2014.
- 9. Methods of sampling and microbiological examination of water, first revision, BIS
- 10. Practical HPLC method development Snyder, Kirkland, Glajch, 2nd edition, John Wiley & Sons.
- 11. Analytical Profiles of drug substances Klaus Florey, Volume 1 20, Elsevier, 2005
- 12. Analytical Profiles of drug substances and Excipients Harry G Brittan, Volume 21 30, Elsevier, 2005.
- 13. The analysis of drugs in biological fluids Joseph Chamberlain, 2nd edition, CRC press, London.
- 14. ICH Guidelines for impurity profiles and stability studies.

# PHARMACEUTICAL VALIDATION (MPA 103T)

### Scope

The main purpose of the subject is to understand about validation and how it can be applied to industry and thus to improve the quality of the products. The subject covers the complete information about validation, types, methodology and application.

## Objectives

Upon completion of the subject student shall be able to

- Explain the aspect of validation
- Carryout validation of manufacturing processes
- Apply the knowledge of validation to instruments and equipments
- Validate the manufacturing facilities

- 1. Introduction: Definition of Qualification and Validation, 12 Advantage of Validation, Streamlining of Qualification & Validation Hrs process and Validation Master Plan.
  - Qualification: User Requirement Specification, Design Qualification, Factory Acceptance Test (FAT)/ Site Acceptance Test (SAT), Installation Qualification, Operational Qualification, Performance Qualification, Re- Qualification (Maintaining status-Calibration Preventive Maintenance, Change management), Qualification of Manufacturing Equipments, Qualification of Analytical Instruments and Laboratory equipments.
- Qualification of analytical instruments: Electronic balance, pH meter, UV-Visible spectrophotometer, FTIR, GC, HPLC, HPTLC Ualification of Glassware: Volumetric flask, pipette, Measuring cylinder, beakers and burette.
- 3 Validation of Utility systems: Pharmaceutical Water System & 12 pure steam, HVAC system, Compressed air and nitrogen. Hrs Cleaning Validation: Cleaning Validation Cleaning Method development, Validation and validation of analytical method used in cleaning. Cleaning of Equipment, Cleaning of Facilities. Cleaning in place (CIP).
- 4 Analytical method validation: General principles, Validation of analytical method as per ICH guidelines and USP.

- Computerized system validation: Electronic records and digital significance-21 CFR part 11 and GAMP 5.
- 5 General Principles of Intellectual Property: Concepts of 12 Intellectual Property (IP), Intellectual Property Protection (IPP), Hrs Economic Property Rights (IPR); importance. mechanism for protection of Intellectual Property -patents, Copyright, Trademark; Factors affecting choice of IP protection; Penalties for violation: Role of IP in pharmaceutical industry: Global ramification and financial implications. Filing a patent applications; patent application forms and guidelines. Types patent applications-provisional and non-provisional, PCT and convention patent applications; International patenting requirement procedures and costs; Rights and responsibilities of a patentee; Practical aspects regarding maintaining of a Patent file; Patent infringement meaning and scope. Significance of transfer technology (TOT), IP and ethics-positive and negative aspects of IPP; Societal responsibility, avoiding unethical practices.

- 1. B. T. Loftus & R. A. Nash, "Pharmaceutical Process Validation", Drugs and Pharm Sci. Series, Vol. 129, 3rd Ed., Marcel Dekker Inc., N.Y.
- 2. The Theory & Practice of Industrial Pharmacy, 3rd edition, Leon Lachman, Herbert A. Lieberman, Joseph. L. Karig, Varghese Publishing House, Bombay.
- 3. Validation Master plan by Terveeks or Deeks, Davis Harwood International publishing.
- 4. Validation of Aseptic Pharmaceutical Processes, 2nd Edition, by Carleton & Agalloco, (Marcel Dekker).
- 5. Michael Levin, Pharmaceutical Process Scale-Up∥, Drugs and Pharm. Sci. Series, Vol. 157,2nd Ed., Marcel Dekker Inc., N.Y.
- 6. Validation Standard Operating Procedures: A Step by Step Guide for Achieving Compliance in the Pharmaceutical, Medical Device, and Biotech Industries, Syed Imtiaz Haider
- 7. Pharmaceutical Equipment Validation: The Ultimate Qualification Handbook, Phillip A. Cloud, Interpharm Press
- 8. Validation of Pharmaceutical Processes: Sterile Products, Frederick J. Carlton (Ed.) and James Agalloco (Ed.), Marcel Dekker, 2nd Ed.
- 9. Analytical Method validation and Instrument Performance Verification by Churg Chan, Heiman Lam, Y.C. Lee, Yue. Zhang, Wiley Inter Science.

## FOOD ANALYSIS (MPA 104T)

## Scope

This course is designed to impart knowledge on analysis of food constituents and finished food products. The course includes application of instrumental analysis in the determination of pesticides in variety of food products.

## Objectives

At completion of this course student shall be able to understand various analytical techniques in the determination of

- Food constituents
- Food additives
- Finished food products
- Pesticides in food
- And also student shall have the knowledge on food regulations and legislations

- 1. Carbohydrates: classification and properties of food 12 carbohydrates, General methods of analysis of food Hrs carbohydrates, Changes in food carbohydrates during processing, Digestion, absorption and metabolism of carbohydrates, Dietary fibre. Crude fibre and application of food carbohydrates Proteins: Chemistry and classification of amino acids and proteins, Physico-Chemical properties of protein and their structure, general methods of analysis of proteins and amino acids, Digestion, absorption and metabolism of proteins.
- 2 Lipids: Classification, general methods of analysis, refining of fats 12 and oils; hydrogenation of vegetable oils, Determination of Hrs adulteration in fats and oils, Various methods used for measurement of spoilage of fats and fatty foods.
  Vitamins: classification of vitamins, methods of analysis of vitamins, Principles of microbial assay of vitamins of B-series.
- 3 Food additives: Introduction, analysis of Preservatives, 12 antioxidants, artificial sweeteners, flavors, flavor enhancers, Hrs stabilizers, thickening and jelling agents.

  Pigments and synthetic dyes: Natural pigments, their occurrence and characteristic properties, permitted synthetic

- dyes, Non-permitted synthetic dyes used by industries, Method of detection of natural, permitted and non-permitted dyes.
- 4 General Analytical methods for milk, milk constituents and milk 12 products like ice cream, milk powder, butter, margarine, cheese Hrs including adulterants and contaminants of milk.
  Analysis of fermentation products like wine, spirits, beer and
- Pesticide analysis: Effects of pest and insects on various food, 12 use of pesticides in agriculture, pesticide cycle, Hrs organophosphorus and organochlorine pesticides analysis, determination of pesticide residues in grain, fruits, vegetables, milk and milk products.

  Legislation regulations of food products with special emphasis on BIS, Agmark, FDA and US-FDA.

#### REFERENCES

vinegar.

- 1. The chemical analysis of foods David Pearson, Seventh edition, Churchill Livingstone, Edinburgh London, 1976
- 2. Introduction to the Chemical analysis of foods S. Nielsen, Jones & Bartlett publishers, Boston London, 1994.
- 3. Official methods of analysis of AOAC International, sixth edition, Volume I & II. 1997.
- 4. Analysis of Food constituents Multon, Wiley VCH.
- 5. Dr. William Horwitz, Official methods of analysis of AOAC International, 18th edition, 2005.

# PHARMACEUTICAL ANALYSIS PRACTICALS - II (MPA 105P)

- Analysis of Pharmacopoeial compounds and their formulations by UV Vis spectrophotometer
- 2. Simultaneous estimation of multi component containing formulations by UV spectrophotometry
- 3. Experiments based on HPLC
- 4. Experiments based on Gas Chromatography
- 5. Estimation of riboflavin/quinine sulphate by fluorimetry
- 6. Estimation of sodium/potassium by flame photometry
- 7. Assay of official compounds by different titrations
- 8. Assay of official compounds by instrumental techniques.
- 9. Quantitative determination of hydroxyl group.
- 10. Quantitative determination of amino group
- 11. Colorimetric determination of drugs by using different reagents
- 12. Imapurity profiling of drugs
- 13. Calibration of glasswares
- 14. Calibration of pH meter
- 15. Calibration of UV-Visible spectrophotometer
- 16. Calibration of FTIR spectrophotometer
- 17. Calibration of GC instrument
- 18. Calibration of HPLC instrument
- 19. Cleaning validation of any one equipment
- 20. Determination of total reducing sugar
- 21. Determination of proteins
- 22. Determination of saponification value, Iodine value, Peroxide value, Acid value in food products
- 23. Determination of fat content and rancidity in food products
- 24. Analysis of natural and synthetic colors in food
- 25. Determination of preservatives in food
- 26. Determination of pesticide residue in food products
- 27. Analysis of vitamin content in food products
- 28. Determination of density and specific gravity of foods
- 29. Determination of food additives

# ADVANCED INSTRUMENTAL ANALYSIS (MPA 201T)

## Scope

This subject deals with various hyphenated analytical instrumental techniques for identification, characterization and quantification of drugs. Instruments dealt are LC-MS, GC-MS, and hyphenated techniques.

## Objectives

After completion of course student is able to know,

- interpretation of the NMR, Mass and IR spectra of various organic compounds
- theoretical and practical skills of the hyphenated instruments
- identification of organic compounds

- 1. HPLC: Principle, instrumentation, pharmaceutical applications, 12 peak shapes, capacity factor, selectivity, plate number, plate Hrs height, resolution, band broadening, pumps, injector, detectors, columns, column problems, gradient HPLC, HPLC solvents, trouble shooting, sample preparation, method development. New developments in HPLC-role and principles of ultra, nano liquid chromatography in pharmaceutical analysis. Immobilized polysaccharide CSP's: Advancement in enantiomeric separations, revised phase Chiral method development and HILIC approaches. HPLC in Chiral analysis of pharmaceuticals. Preparative HPLC, practical aspects of preparative HPLC.
- Biochromatography: Size exclusion chromatography, ion 12 exchange chromatography, ion pair chromatography, affinity Hrs chromatography general principles, stationary phases and mobile phases.
  - Gas chromatography: Principles, instrumentation, derivatization, head space sampling, columns for GC, detectors, quantification. High performance Thin Layer chromatography: Principles, instrumentation, pharmaceutical applications.
- 3 Super critical fluid chromatography: Principles, 12 instrumentation, pharmaceutical applications. Hrs Capillary electrophoresis: Overview of CE in pharmaceutical analysis, basic configuration, CE characteristics, principles of CE, methods and modes of CE. General considerations and method

- development in CE, Crown ethers as buffer additives in capillary electrophoresis. CE-MS hyphenation.
- 4 Mass spectrometry: Principle, theory, instrumentation of mass spectrometry, different types of ionization like electron impact, chemical, field, FAB and MALD, APCI, ESI, APPI mass fragmentation and its rules, meta stable ions, isotopic peaks and applications of mass spectrometry. LC-MS hyphenation and DART MS analysis. Mass analysers (Quadrpole, Time of flight, FT-ICR, ion trap and Orbitrap) instruments. MS/MS systems (Tandem: QqQ, TOF-TOF;Q-IT, Q-TOF, LTQ-FT, LTQ-Orbitrap.
- NMR spectroscopy: Quantum numbers and their role in NMR, 12 Principle, Instrumentation, Solvent requirement in NMR, Hrs Relaxation process, NMR signals in various compounds, Chemical shift, Factors influencing chemical shift, Spin-Spin coupling, Coupling constant, Nuclear magnetic double resonance, Brief outline of principles of FT-NMR with reference to 13CNMR: Spin spin and spin lattice relaxation phenomenon. 13C NMR, 1-D and 2-D NMR, NOESY and COSY techniques, Interpretation and Applications of NMR spectroscopy. LC-NMR hyphenations.

- 1. Spectrometric Identification of Organic compounds Robert M Silverstein, Sixth edition, John Wiley & Sons, 2004.
- 2. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5<sup>th</sup> edition, Eastern press, Bangalore, 1998.
- 3. Instrumental methods of analysis Willards, 7<sup>th</sup> edition, CBS publishers.
- 4. Organic Spectroscopy William Kemp, 3<sup>rd</sup> edition, ELBS, 1991.
- 5. Quantitative analysis of Pharmaceutical formulations by HPTLC P D Sethi, CBS Publishers, New Delhi.
- 6. Quantitative Analysis of Drugs in Pharmaceutical formulation P D Sethi, 3<sup>rd</sup> Edition, CBS Publishers, New Delhi, 1997.
- 7. Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11. Marcel Dekker Series.
- 8. Organic Spectroscopy by Donald L. Paviya, 5th Edition.

## MODERN BIO-ANALYTICAL TECHNIQUES (MPA 202T)

## Scope

This subject is designed to provide detailed knowledge about the importance of analysis of drugs in biological matrices.

### Objectives

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

- Extraction of drugs from biological samples
- Separation of drugs from biological samples using different techniques
- Guidelines for BA/BF studies.

- Extraction of drugs and metabolites from biological matrices: 12
   General need, principle and procedure involved in the Bioanalytical methods such as Protein precipitation, Liquid Liquid extraction and Solid phase extraction and other novel sample preparation approach.
   Bioanalytical method validation: USFDA and EMEA guidelines.
- 2 Biopharmaceutical Consideration:
  Introduction, Biopharmaceutical Factors Affecting Drug Hrs
  Bioavailability, In Vitro: Dissolution and Drug Release Testing,
  Alternative Methods of Dissolution Testing Transport models,
  Biopharmaceutics Classification System. Solubility: Experimental
  methods. Permeability: In-vitro, in-situ and In-vivo methods.
- Basic consideration, Drug interaction (PK-PD interactions), The effect of protein-binding interactions, The effect of tissue-binding interactions, Cytochrome P450-based drug interactions, Drug interactions linked to transporters. Microsomal assays Toxicokinetics-Toxicokinetic evaluation in preclinical studies, Importance and applications of toxicokinetic studies. LC-MS in bioactivity screening and proteomics.
- 4 Cell culture techniques
  Basic equipments used in cell culture lab. Cell culture media, various types of cell culture, general procedure for cell cultures; isolation of cells, subculture, cryopreservation, characterization of

cells and their applications. Principles and applications of cell viability assays (MTT assays), Principles and applications of flow cytometry.

### 5 Metabolite identification:

12 Hrs

In-vitro / in-vivo approaches, protocols and sample preparation. Microsomal approaches (Rat liver microsomes (RLM) and Human liver microsomes (HLM) in Met -ID. Regulatory perspectives. In-vitro assay of drug metabolites & drug metabolizing enzymes.

Drug Product Performance, In Vivo: Bioavailability and Bioequivalence:

Drug Product Performance, Purpose of Bioavailability Studies, Relative and Absolute Availability. Methods for Assessing Bioavailability, Bioequivalence Studies, Design and Evaluation of Bioequivalence Studies, Study Designs, Crossover Study Designs, Generic Biologics (Biosimilar Drug Products), Clinical Significance of Bioequivalence Studies.

- Analysis of drugs in Biological fluids Joseph Chamberlain, 2<sup>nd</sup> Edition. CRC Press, Newyork. 1995.
- 2. Principles of Instrumental Analysis Doglas A Skoog, F. James Holler, Timothy A. Nieman, 5<sup>th</sup> edition, Eastern press, Bangalore, 1998.
- 3. Pharmaceutical Analysis Higuchi, Brochmman and Hassen, 2<sup>nd</sup> Edition, Wiley Interscience Publications, 1961.
- Pharmaceutical Analysis- Modern methods Part B J W Munson, Volume 11, Marcel Dekker Series
- 5. Practical HPLC method Development Snyder, Kirkland, Glaich, 2<sup>nd</sup> Edition, John Wiley & Sons, New Jercy. USA.
- 6. Chromatographic Analysis of Pharmaceuticals John A Adamovics, 2<sup>nd</sup> Edition, Marcel Dekker, Newyork, USA. 1997.
- 7. Chromatographic methods in clinical chemistry & Toxicology Roger L Bertholf, Ruth E Winecker, John Wiley & Sons, New Jercy, USA. 2007.
- 8. Good Laboratory Practice Regulations, 2<sup>nd</sup> Edition, Sandy Weinberg Vol. 69, Marcel Dekker Series, 1995.
- Good laboratory Practice Regulations Allen F. Hirsch, Volume 38, Marcel Dekker Series, 1989.
- 10. ICH, USFDA & CDSCO Guidelines.
- 11. Palmer

# QUALITY CONTROL AND QUALITY ASSURANCE (MPA 203T)

## Scope

This course deals with the various aspects of quality control and quality assurance aspects of pharmaceutical industries. It covers the important aspects like cGMP, QC tests, documentation, quality certifications, GLP and regulatory affairs.

### Objectives

At the completion of this subject it is expected that the student shall be able to know

- the cGMP aspects in a pharmaceutical industry
- to appreciate the importance of documentation
- to understand the scope of quality certifications applicable to Pharmaceutical industries
- to understand the responsibilities of QA & QC departments

- 1. Concept and Evolution of Quality Control and Quality 12
  Assurance Hrs
  Good Laboratory Practice, GMP, Overview of ICH Guidelines QSEM, with special emphasis on Q-series guidelines.
  Good Laboratory Practices: Scope of GLP, Definitions, Quality assurance unit, protocol for conduct of non clinical testing, control on animal house, report preparation and documentation.
- cGMP guidelines according to schedule M, USFDA (inclusive of CDER and CBER) Pharmaceutical Inspection Convention Hrs (PIC), WHO and EMEA covering: Organization and personnel responsibilities, training, hygiene and personal records, drug industry location, design, construction and plant lay out, maintenance, sanitation, environmental control, utilities and maintenance of sterile areas, control of contamination and Good Warehousing Practice. CPCSEA guidelines.
- 3. Analysis of raw materials, finished products, packaging 12 materials, in process quality control (IPQC), Developing Hrs specification (ICH Q6 and Q3)

Purchase specifications and maintenance of stores for raw materials. In process quality control and finished products quality control for following formulation in Pharma industry according to Indian, US and British pharmacopoeias: tablets, capsules, ointments, suppositories, creams, parenterals, ophthalmic and surgical products (How to refer pharmacopoeias), Quality control test for containers, closures and secondary packing materials.

12

Hrs

- 4. Documentation in pharmaceutical industry: Three tier documentation, Policy, Procedures and Work instructions, and records (Formats), Basic principles- How to maintain, retention and retrieval etc. Standard operating procedures (How to write), Master Formula Record, Batch Formula Record, Quality audit plan and reports. Specification and test procedures, Protocols and reports. Distribution records. Electronic data.
- 5. Manufacturing operations and controls: Sanitation of 12 manufacturing premises, mix-ups and cross contamination, Hrs processing of intermediates and bulk products, packaging operations, IPQC, release of finished product, process deviations, charge-in of components, time limitations on production, drug product inspection, expiry date calculation, calculation of yields, production record review, change control, sterile products, aseptic process control, packaging.

- 1. Quality Assurance Guide by organization of Pharmaceutical Procedures of India, 3<sup>rd</sup> revised edition, Volume I & II, Mumbai, 1996.
- 2. Good Laboratory Practice Regulations, 2<sup>nd</sup> Edition, Sandy Weinberg Vol. 69, Marcel Dekker Series, 1995.
- 3. Quality Assurance of Pharmaceuticals- A compedium of Guide lines and Related materials Vol I & II, 2<sup>nd</sup> edition, WHO Publications, 1999.
- 4. How to Practice GMP's P P Sharma, Vandana Publications, Agra, 1991.
- 5. The International Pharmacopoeia vol I, II, III, IV & V General Methods of Analysis and Quality specification for Pharmaceutical Substances, Excepients and Dosage forms, 3<sup>rd</sup> edition, WHO, Geneva, 2005.
- 6. Good laboratory Practice Regulations Allen F. Hirsch, Volume 38, Marcel Dekker Series, 1989.
- 7. ICH guidelines
- 8. ISO 9000 and total quality management

- 9. The drugs and cosmetics act 1940 Deshpande, Nilesh Gandhi, 4<sup>th</sup> edition, Susmit Publishers, 2006.
- 10. QA Manual D.H. Shah, 1<sup>st</sup> edition, Business Horizons, 2000.
- 11. Good Manufacturing Practices for Pharmaceuticals a plan for total quality control Sidney H. Willig, Vol. 52, 3<sup>rd</sup> edition, Marcel Dekker Series.
- 12. Steinborn L. GMP/ISO Quality Audit Manual for Healthcare Manufacturers and Their Suppliers, Sixth Edition, (Volume 1 With Checklists and Software Package). Taylor & Francis; 2003.
- 13. Sarker DK. Quality Systems and Controls for Pharmaceuticals. John Wiley & Sons; 2008.

# HERBAL AND COSMETIC ANALYSIS (MPA 204T)

### Scope

This course is designed to impart knowledge on analysis of herbal products. Regulatory requirements, herbal drug interaction with monographs. Performance evaluation of cosmetic products is included for the better understanding of the equipments used in cosmetic industries for the purpose.

### **Objectives**

At completion of this course student shall be able to understand

- Determination of herbal remedies and regulations
- Analysis of natural products and monographs
- Determination of Herbal drug-drug interaction
- Principles of performance evaluation of cosmetic products.

- Herbal remedies- Toxicity and Regulations: Herbals vs 12
  Conventional drugs, Efficacy of herbal medicine products, Hrs
  Validation of Herbal Therapies, Pharmacodynamic and
  Pharmacokinetic issues. Herbal drug standardization: WHO and
  AYUSH guidelines.
- Adulteration and Deterioration: Introduction, types of 12 adulteration/substitution of herbal drugs, Causes and Measure of Hrs adulteration, Sampling Procedures, Determination of Foreign Matter, DNA Finger printing techniques in identification of drugs of natural origin, heavy metals, pesticide residues, phototoxin and microbial contamination in herbal formulations. Regulatory requirements for setting herbal drug industry: Global marketing management, Indian and international patent law as applicable herbal drugs and natural products and its protocol.
- 3 Testing of natural products and drugs: Effect of herbal 12 medicine on clinical laboratory testing, Adulterant Screening using Hrs modern analytical instruments, Regulation and dispensing of herbal drugs, Stability testing of natural products, protocol.
  - Monographs of Herbal drugs: Study of monographs of herbal drugs and comparative study in IP, USP, Ayurvedic

Pharmacopoeia, American herbal Pharmacopoeia, British herbal Pharmacopoeia, Siddha and Unani Pharmacopoeia, WHO guidelines in quality assessment of herbal drugs.

4 Herbal drug-drug interaction: WHO and AYUSH guidelines for 12 safety monitoring of natural medicine, Spontaneous reporting Hrs schemes for bio drug adverse reactions, bio drug-drug and bio drug-food interactions with suitable examples. Challenges in monitoring the safety of herbal medicines.

12

5 Evaluation of cosmetic products: Determination of acid value, ester value, saponification value, iodine value, peroxide value, rancidity, moisture, ash, volatile matter, heavy metals, fineness of powder, density, viscosity of cosmetic raw materials and finished products. Study of quality of raw materials and general methods of analysis of raw material used in cosmetic manufacture as per BIS.

Indian Standard specification laid down for sampling and testing of various cosmetics in finished forms such as baby care products, skin care products, dental products, personal hygiene preparations, lips sticks. Hair products and skin creams by the Bureau Indian Standards.

- 1. Pharmacognosy by Trease and Evans
- 2. Pharmacognosy by Kokate, Purohit and Gokhale
- 3. Quality Control Methods for Medicinal Plant, WHO, Geneva
- 4. Pharmacognosy & Pharmacobiotechnology by Ashutosh Kar
- 5. Essential of Pharmacognosy by Dr.S.H.Ansari
- 6. Cosmetics Formulation, Manufacturing and Quality Control, P.P. Sharma, 4<sup>th</sup> edition, Vandana Publications Pvt. Ltd., Delhi
- 7. Indian Standard specification, for raw materials, BIS, New Delhi.
- 8. Indian Standard specification for 28 finished cosmetics BIS. New Delhi
- 9. Harry's Cosmeticology 8th edition
- 10. Suppliers catalogue on specialized cosmetic excipients
- 11. Wilkinson, Moore, seventh edition, George Godwin. Poucher's Perfumes, Cosmetics and Soaps
- 12. Hilda Butler, 10th Edition, Kluwer Academic Publishers. Handbook of Cosmetic Science and Technology, 3rd Edition,

# PHARMACEUTICAL ANALYSIS PRACTICALS - I (MPA 205P)

- 1. Comparison of absorption spectra by UV and Wood ward Fiesure rule
- 2. Interpretation of organic compounds by FT-IR
- 3. Interpretation of organic compounds by NMR
- 4. Interpretation of organic compounds by MS
- 5. Determination of purity by DSC in pharmaceuticals
- 6. Identification of organic compounds using FT-IR, NMR, CNMR and Mass spectra
- 7. Bio molecules separation utilizing various sample preparation techniques and Quantitative analysis of components by gel electrophoresis.
- 8. Bio molecules separation utilizing various sample preparation techniques and Quantitative analysis of components by HPLC techniques.
- 9. Isolation of analgesics from biological fluids (Blood serum and urine).
- 10. Protocol preparation and performance of analytical/Bioanalytical method validation.
- 11. Protocol preparation for the conduct of BA/BE studies according to guidelines.
- 12. In process and finished product quality control tests for tablets, capsules, parenterals and creams
- 13. Quality control tests for Primary and secondary packing materials
- 14. Assay of raw materials as per official monographs
- 15. Testing of related and foreign substances in drugs and raw materials
- 16. Preparation of Master Formula Record.
- 17. Preparation of Batch Manufacturing Record.
- 18. Quantitative analysis of rancidity in lipsticks and hair oil
- 19. Determination of aryl amine content and Developer in hair dye
- 20. Determination of foam height and SLS content of Shampoo.
- 21. Determination of total fatty matter in creams (Soap, skin and hair creams)
- 22. Determination of acid value and saponification value.
- 23. Determination of calcium thioglycolate in depilatories

# Semester III MRM 301T - Research Methodology & Biostatistics

#### UNIT – I

General Research Methodology: Research, objective, requirements, practical difficulties, review of literature, study design, types of studies, strategies to eliminate errors/bias, controls, randomization, crossover design, placebo, blinding techniques.

### UNIT - II

Biostatistics: Definition, application, sample size, importance of sample size, factors influencing sample size, dropouts, statistical tests of significance, type of significance tests, parametric tests(students "t" test, ANOVA, Correlation coefficient, regression), non-parametric tests (wilcoxan rank tests, analysis of variance, correlation, chi square test), null hypothesis, P values, degree of freedom, interpretation of P values.

#### UNIT - III

Medical Research: History, values in medical ethics, autonomy, beneficence, non-maleficence, double effect, conflicts between autonomy and beneficence/non-maleficence, euthanasia, informed consent, confidentiality, criticisms of orthodox medical ethics, importance of communication, control resolution, guidelines, ethics committees, cultural concerns, truth telling, online business practices, conflicts of interest, referral, vendor relationships, treatment of family members, sexual relationships, fatality.

#### UNIT - IV

CPCSEA guidelines for laboratory animal facility: Goals, veterinary care, quarantine, surveillance, diagnosis, treatment and control of disease, personal hygiene, location of animal facilities to laboratories, anesthesia, euthanasia, physical facilities, environment, animal husbandry, record keeping, SOPs, personnel and training, transport of lab animals.

#### UNIT - V

Declaration of Helsinki: History, introduction, basic principles for all medical research, and additional principles for medical research combined with medical care.

