

### AY 2020-21 Onwards

## Hindi Vidya Prachar Samiti's RAMNIRANJAN JHUNJHUNWALA COLLEGE (AUTONOMOUS)

(Also known as R. J. College of Arts, Science & Commerce as per UGC Notification)

Affiliated to UNIVERSITY OF MUMBAI II Recognized by UGC under 2f & 12B NAAC Accredited 'A GRADE' with CGPA 3.50

Knowledge is all Ambrosia

Postgraduate Diploma in

**Clinical Studies,** 

**Data Management** 

&

**Medical Writing** 

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Opposite Railway Station, Ghatkopar (W), Mumbai 400 086, Maharashtra, INDIA. F.Y.B.Sc Botany Syllabus Semester I & II



### Hindi Vidya Prachar Samiti's RamniranjanJhunjhunwala College Of Arts, Science & Commerce (Autonomous College)

Affiliated to UNIVERSITY OF MUMBAI

Syllabus for the Post Graduate Diploma

# Program: Post Graduate Diploma in Clinical Studies, Data Management and Medical Writing

Program Code: RJSPGDCSDMMW

(Revised Syllabus Academic year 2020-2021)

#### Post Graduate Diploma in

#### **Clinical Studies, Data Management and Medical Writing**

#### Program Code: RJPGDCSDMMW

#### Preamble:

Clinical research has led to medical advances benefiting mankind. Clinical research is performed on human beings to evaluate safety and efficacy of a new treatment or a new drug. Clinical trial is aimed to evaluate if the medical intervention is more effective and less harmful than the standard treatment. Regulatory authorities approve or reject the marketing of the drug depending on the data obtained from clinical trial. Professional in the specialized industry of clinical research need to have good knowledge of International Conference on Harmonization Good Clinical Practices guidelines. Different level of training is needed at different functional levels of clinical research due to the complexity of overall drug development process. Clinical data management focuses on clinical trial data collection and data management. Methodical and scientific knowledge with good basic writing skills is essential for medical writing. In additional to technical skills, good interpersonal skills and excellent communication is essential for clinical research professional.

Post Graduate Diploma in Clinical Studies, data management and medical writing offers essentials of clinical research in a profound and concise manner to understand the overall process of drug development. The course emphasizes the theoretical and ethical clinical research and practical aspects of conducting clinical trials. The course is ideal for Graduates with the background of Life Science / Biotechnology / Chemistry / Statistics to build qualification and expertise to enter into the exciting world of clinical research. Real-world clinical trial case studies reinforce the knowledge of clinical studies that will help individuals accelerate their career in clinical research.

#### **Program Objectives and Outcomes:**

This course is based on systematic teaching, learning and evaluation techniques. Quality based training, comprehensive coaching by experts from industry professionals and dynamic mentoring will fulfill following objectives.

- To understand the overall drug development process in compliance with ICH GCP, regulatory guidelines and ethics
- To identify and differentiate designs and phases of clinical trial and comprehend about the responsibilities of different stakeholders
- To consider the process of monitoring and managing a clinical trial conducted at the study site
- To understand the basics of clinical data management
- To contemplate the rationale of pharmacovigilance and medical writing

Title: Post Graduate Diploma in Clinical Studies, Data Management and Medical Writing

Eligibility: Bachelor's Degree in Life Science / Biotechnology / Chemistry / Statistics

Duration of the course: One Year (Part time) Blended teaching weekends

#### Fee Structure:

Tuition Fee	:	35,000.00
RFID	:	150.00
Examination Fees	:	500.00
Application form fees	:	Rs 100.00

Intake capacity: 40 students

**Faculty:** Industry and Academia Professional in area of Clinical research, Pharmacy, Clinical Data Management, Pharmacovigilance and Medical writing

#### Standard of Passing:

- a. Candidate who secures minimum 50% marks in each paper be declared to have passed the examination in that subject.
- b. A candidate who fails to secure 50% marks in a paper will be allowed to reappear in that paper.
- c. Candidate can carry forward at his/her option the marks in the paper in which he/she has passed, in such a case student is entitled for award of class.
- d. Candidate who secures a minimum of 50% marks in each paper and an aggregate of 60% and above marks on the whole shall be declared to have passed the examination in the First Class.
- e. Candidate who secures a minimum of 50% marks in each paper and an aggregate of 70% and above marks on the whole shall be declared to have passed the examination in First Class with Distinction.

#### Medium of Instruction: English

#### Scheme of Evaluation: There would be continuous evaluation

- Internal assessment: 40% weightage multiple choice questions, quiz, assignments, case studies
- External assessment: 60% weightage subjective questions

ER	PAPER	INTERNAL		EXTERNAL		ITS	CODE
PAPER	TITLE OF	Maximum	Minimum	Maximum	Minimum	CREDITS	COURSE
I	Clinical Pharmacology and Toxicology	40	16	60	24	8	RJSPGDCSDMMW101
П	Good Practices & Ethics	40	16	60	24	8	RJSPGDCSDMMW102
Ш	Clinical Trial Processes	40	16	60	24	8	RJSPGDCSDMMW103
	Total	120	48	180	72	24	

#### SEMESTER I

#### SEMESTER II

ER	PAPER	INTERNAL		EXTERNAL		ITS	CODE
PAPER	TITLE OF	Maximum	Minimum	Maximum	Minimum	CREDITS	COURSE
I	Regulations	40	16	60	24	8	RJSPGDCSDMMW201
II	Pharmacovigilance	40	16	60	24	8	RJSPGDCSDMMW202
111	Clinical Data Management & Medical Writing	40	16	60	24	8	RJSPGDCSDMMW203
	Total	120	48	180	72	24	

Total (600) = Internal (240) + External (360)

Total Credits = 48

Page **5** of **12** 

#### **SEMESTER I**

#### Paper I: Clinical Pharmacology and Toxicology

8 Credits

#### UNIT – 1:

**Pharmaceutical Industry & globalization** –Overview, Opportunities & Career options in Clinical Research

#### UNIT – 2:

**Pharmacy** - Physico-Chemical properties of drugs, different drug dosage forms, Formulation development and manufacture of drugs.

**Therapeutics** - Principles of Management & Drug Therapy

#### UNIT – 3:

**Pharmacokinetics** - Absorption, bioavailability, distribution, metabolism protein binding, excretion, placental and blood brain barrier

**Pharmacodynamics** - Mechanism of drug action, receptors, agonists, antagonists, side effects and adverse events

#### UNIT – 4:

**Toxicology** - Acute, Sub-acute and Chronic Toxicity, Mutagenicity, Teratogenecity, Oncogenicity and effects on **fertility; pre-clinical studies** 

#### Paper II: Good Practices & Ethics

8 Credits

#### UNIT – 1:

Good Manufacturing Practices, Good Laboratory Practices

#### UNIT – 2:

International Conference on Harmonization, Good Clinical Practices

#### UNIT – 3:

History of Ethics in Clinical Research and Ethics Committee

#### UNIT – 4:

Belmont Report, Nuremberg Codes, Declaration of Helsinki

#### Paper III: Clinical Trial Processes

#### 8 Credits

#### UNIT - 1:

Responsibilities of Stakeholders: Sponsors, Investigators, CROs, Monitors; Clinical Trial Designs

#### UNIT – 2:

Clinical Trial Phase I, Phase II, Phase IV

#### UNIT – 3: Essential Documents in Clinical Trials:

SOP, Protocol, Investigator Brochure, Master Files, Informed Consent Forms, Case Record Form

#### UNIT – 4:

**Managements of Clinical Trials** - Investigator's Meeting, Project management, Patient Recruitment & Retention, Trial Monitoring, Drug Resource and supplies; Trial Budget, Audit and Inspection

#### SEMESTER II

#### Paper I: Regulations

#### 8 Credits

UNIT – 1:

**BA/BE Studies** - Bioavailability and Bioequivalence - Methods and Procedures, regulatory requirements, planning & design, Protocol/ CRF outline, QA & QC, Drug accountability

#### UNIT – 2:

**Regulation in India**: Drugs and Cosmetics Act, Schedule 'Y', Quality in Regulatory Context, Patent laws; **ICMR** 

#### UNIT – 3:

USFDA: History, Structure & Function, Code of Federal Regulation

#### UNIT – 4:

**EMEA**: History, Structure & Function, Regulations;

JAPAN : History, Structure & Function

#### Paper II: Pharmacovigilance

8 Credits

UNIT – 1:

**Overview of Pharmacovigilance:** Importance; National & International Programs; Methods

UNIT – 2:

**Principles of Pharmacovigilance:** ADR; Assessment; Medication errors, Signal detection; Risk assessments

UNIT – 3:

Drug Dictionaries: Coding & Tools; Drug Safety: PSURs; Package inserts

UNIT – 4:

**Regulatory Guidelines:** ICH, EMEA, USFDA, Sch. 'Y'

Page **10** of **12** 

#### Paper III: Clinical Data Management & Medical Writing 8 Credits

#### UNIT – 1:

**Biostatistics: Descriptive Statistics** - Data Types; Collection; Sampling, Compilation; Tables & Graphs, Measures of Central Tendency, Measures of variation

#### UNIT – 2:

Clinical Data Management: Overview, scope, terminologies; Principles of CDM

#### UNIT – 3:

**Clinical Data Management**: Data Entry, Queries & Data Clarification, Electronic Data Capture, Software in CDM

#### UNIT – 4:

**Medical Writing:** Literature Search & Medical Articles, Contract writing, Publication, Abstracts, Bibliography, Clinical Study Reports

#### List of Reference Books:

- Research in education by J W Best and J V Khan Prentice Hall of India, New Delhi (1995).
- Pharmaceutical Statistics by Sanford Bolton, Marcel Dekker, New York, USA, Informa Healthcare; 4 edition (October 17, 2003).
- Elementary Statistical Quality Control, Volume 25, Burr, I. W. (1979), New York: Marcel Dekker, Inc.
- 4. Managing the clinical drug development process, C. Nardi, Marcel Dekker, New York, USA (1991).
- 5. Basic managerial skills for all by E H Mcgrath, Prentice Hall of India, N.Delhi (2002).
- 6. Clinical Research Environment in India by Umakanta Sahoo, Faiz Kermani, ICFAI University Press (2008).
- 7. Clinical Trials. Lelia Duley and Barbara Farrell (eds), BMJ Books, London, 2002.
- 8. Handbook for good clinical research practice WHO Library Catalogue.
- 9. Articles: ICH-GCP, Schedule Y, US FDA guidelines, WHO Guidelines.
- 10. Bioavailability and Bioequivalence in pharmaceutical technology by Tapan Kumar and Ganeshan M, CBS publishers and distributers(2006).
- Design of experiments. A realistic approach by V L Anderson and Robert Mclean, Marcel Dekker, New York, USA (1974).
- 12. Fundamentals of Clinical Research: Bridging Medicine, Statistics and Operations, Antonella Bacchieri and Giovanni Della Cioppa, Springer (2007)

### Postgraduate Diploma in Clinical Studies, Data Management & Medical Writing

Sr. No	Roll No	Name of the Student	Grade	Certificate No
1	1	CHAVAN AJIT VIJAY JYOTI	A	PGDCSDMMW-201920-01
2	2	JAWKE AKANKSHA BABAN ANKITA	A+	PGDCSDMMW-201920-02
3	3	SHINDE AKSHATA YASHWANT YOGITA	A	PGDCSDMMW-201920-03
4	4	BHOSALE ALISHA CHANDRASHEKHAR VIDYA	0	PGDCSDMMW-201920-04
5	5	JAKATE AMRUTA GANESH CHITRA	A	PGDCSDMMW-201920-05
6	6	GAIKWAD ANIKET SUNIL SUVARNA	A	PGDCSDMMW-201920-06
7	7	TANKARIA Anjaly SHAILESH PARUL	A+	PGDCSDMMW-201920-07
8	8	PAWAR ANKITA RAJARAM MANGAL	A+	PGDCSDMMW-201920-08
9	9	PANDEY ANVEKSHA ARUN KUMAR ARATI DEVI	A+	PGDCSDMMW-201920-09
10	10	SINGH ASHISHKUMAR DEVENDRABAHADUR KIRAN	A	PGDCSDMMW-201920-10
11	11	THOMBARE DHANASHREE MAHENDRA SINDHU	0	PGDCSDMMW-201920-11
12	12	PAKHARE DIVYA BALASAHEB PRADNYA	B+	PGDCSDMMW-201920-12
13	13	KHAN FAIZALAM RIYAZ ZARGUNJAAN	A+	PGDCSDMMW-201920-13
14	14	BAGATE HARSHAWARDHAN SUBHASH KEVAL	A	PGDCSDMMW-201920-14
15	15	YADAV JITESH ANAND APARNA	A+	PGDCSDMMW-201920-15
16	16	OVHAL KAJAL ARUN ASHA	A+	PGDCSDMMW-201920-16
17	17	MAHADIK KOMAL VIJAY SHOBHA	A+	PGDCSDMMW-201920-17
18	18	GUPTA KOMAL RAJKUMAR MAMTADEVI	A+	PGDCSDMMW-201920-18
19	19	KAMBLI MANSI SADANAND SANGITA	A+	PGDCSDMMW-201920-19
20	20	DHOLAKIA NOOPUR MRUGESH TAMANNA	0	PGDCSDMMW-201920-20
21	21	KANSE PANKAJ BALKRISHNA VIMAL	A	PGDCSDMMW-201920-21
22	22	DEOLEKAR POOJA NANDALAL NEETA	A+	PGDCSDMMW-201920-22
23	23	JOSHI POOJA SHRIRAM SEEMA	A+	PGDCSDMMW-201920-23
24	24	CHIVILKAR PRADNYA DAGADU SUMITRA	В	PGDCSDMMW-201920-24
25	25	MISHRA PRAGATI SUDHAKAR PRATIMA	F	PGDCSDMMW-201920-25
26	26	THAKKAR PRANAV RAVILAL DEVIBEN	0	PGDCSDMMW-201920-26
27	27	SHIVADE PRITAM ANANDA SUREKHA	A	PGDCSDMMW-201920-27
28	28	HALDANKAR RADHIKA MAHENDRA MANSI	A+	PGDCSDMMW-201920-28
29	29	GHADGE RESHMA PRADIP ASHA	A+	PGDCSDMMW-201920-29
30	30	GURAV RUCHIRA RAVINDRA SHARAYU	A+	PGDCSDMMW-201920-30
31	31	KAMBLE RUTUJA SANJAY MEGHA	A+	PGDCSDMMW-201920-31
32	32	MIRZA SAIMA KHALID FARZANA	A+	PGDCSDMMW-201920-32
33	33	KOTIAN SHWETA SADASHIV MOHINI	A+	PGDCSDMMW-201920-33
34	34	TODKAR SIDDHARTH PRALHAD SAMPADA	A+	PGDCSDMMW-201920-34
35	35	SITAP SIDDHI SANJAY VEENA	A+	PGDCSDMMW-201920-35
36	36	TIWARI SNEHA JAYNARAYAN MANJU	A+	PGDCSDMMW-201920-36
37	37	PATIL SOHA RAJDATTA NILAKSHI	0	PGDCSDMMW-201920-37
38	38	KONAR SUGANYA MUPPIDATHI PECHI	A	PGDCSDMMW-201920-38
39	39	PARKHE SUPRIYA BALASAHEB YOGESHRI	A+	PGDCSDMMW-201920-39
40	40	GHOSAL SUSHMITA JAGANNATH MANIDIPA	0	PGDCSDMMW-201920-40

41	41	AHUJA TANYA RANBIR KALPANA	A+	PGDCSDMMW-201920-41
42	42	ZADMALI TEJASVI DR ANAND DAISYRANI	A+	PGDCSDMMW-201920-42
43	43	VADHAR VIDHI MANISH SONAL	A+	PGDCSDMMW-201920-43
44	44	PRAJAPATI VIDHI SATISH LEENA	A+	PGDCSDMMW-201920-44
45	45	BATTULA VINIT RAVI BATTULA RUPA BATTULA	A+	PGDCSDMMW-201920-45
46	46	KALE VINIT VISHWAS VAISHALI	0	PGDCSDMMW-201920-46
47	47	QURESHI ZUBER ABDULAZIZ BILKIS	A	PGDCSDMMW-201920-47
48	48	SINGH SUMEET SANJAY BABITA	A+	PGDCSDMMW-201920-48
49	49	JAISWAR INDRAJEET BECHAN RAM MALTI	A+	PGDCSDMMW-201920-49
50	50	BORKHEDE DINESH HANUMANT RUKMINI	A+	PGDCSDMMW-201920-50

Hindi Vidya Prachar Samiti's RAMNIRANJAN JHUNJHUNWALA COLLEGE of Arts, Science & Commerce (Autonomous)

**Affiliated to UNIVERSITY OF MUMBAI** 

# **CERTIFICATE OF MERIT**



This is to certify that

#### CHAVAN AJIT VIJAY JYOTI

has secured

### "A"

in Postgraduate Diploma in Clinical Studies Data Management and Medical Writing

in the examination held in Aug 2020.

myak

Controller of Examination

Certificate ID: PGDCSDMMW-201920-01

Principal