

OUTCOME

On completion, the enrolled participants will be able to:

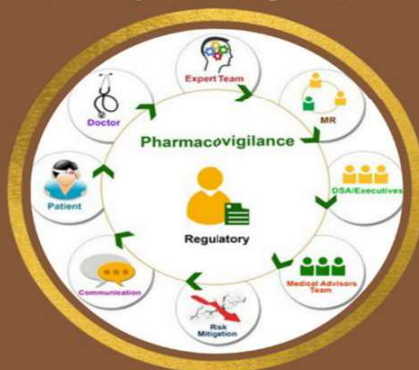
- Identify ADRs / signals in real world setup.
- Understand the need of ADR reporting.
- Assess causality of ADRs.

GUIDELINES

- MBBS 2nd year onwards (incl PGs), UGs & PGs of allied health sciences from nursing, dental, pharmacy & physiotherapy colleges can enrol.
- A minimum of 80% attendance will be required to be eligible for final assessment.
- Final assessment would include objective questions as well as case discussion.
- Certificate on completion.

DESCRIPTION

A series of lectures & workshops to enhance, enrich & equip students with skills required for future endeavors. Extensively designed course of offline lectures to comprehend nuances of why adverse drug monitoring is required.



BENEFITS

- Introduces to the process of detecting & monitoring ADRs through signal detection.
- Prepares for clinical application.
- In demand career opportunity in industrial field.

TEERTHANKER MAHAVEER
MEDICAL COLLEGE &
RESEARCH CENTRE



PHARMACOVIGILANCE MONITORING & TRAINING



Value Added Course
Department of Pharmacology,
TMMC&RC

COURSE SCHEDULE

- Start Date: 30/11/2022
- End Date: 31/3/2023
- Duration: 19 hours
- Registration: From 15/11/2022



CONTACT US

We are readily available to solve your queries regarding this course. You can reach out to us at Department of Pharmacology, 3rd floor Medical College Building, TMU.



RESOURCE FACULTY

1. Dr. PS Matreja (Professor & HOD)
2. Dr. Shipa Patrick (Professor)
3. Dr. Preeti Singh (Professor)
4. Dr. Meenu Thomas (Asst Professor)
5. Dr. Pooja Agarwal (Asst Professor)
6. Dr. Dhyuti Gupta (Senior Resident)



COURSE CO-ORDINATOR

Dr Prithpal Singh Matreja
Professor & HOD
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TMMC & RC
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Contact number: 9855001847

COURSE DETAILS

MODULE	COURSE
1	Introduction to need of Pharmacovigilance
2	What are adverse drug reactions (ADRs)
3	ADR – Focusing on drug hypersensitivity
4	Global pharmacovigilance status
5	What is the national pharmacovigilance status
6	Assessing causality, severity & preventability of ADRs
7	Hands on – Filling up ADR form
8	Hands on – Online reporting of ADR
9	Hands on – Case based workup
10	Feedback & Evaluation