



GUJARAT TECHNOLOGICAL UNIVERSITY

Bachelor of Pharmacy

Subject Code: BP805TT

SEMESTER: VIII

Subject Name: PHARMACOVIGILANCE

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

Objectives: *At completion of this paper it is expected that students will be able to (know, do, and appreciate):*

1. Why drug safety monitoring is important?
2. History and development of pharmacovigilance
3. National and international scenario of pharmacovigilance
4. Dictionaries, coding and terminologies used in pharmacovigilance
5. Detection of new adverse drug reactions and their assessment
6. International standards for classification of diseases and drugs
7. Adverse drug reaction reporting systems and communication in pharmacovigilance
8. Methods to generate safety data during pre clinical, clinical and post approval phases of drugs' life cycle
9. Drug safety evaluation in paediatrics, geriatrics, pregnancy and lactation
10. Pharmacovigilance Program of India (PvPI) requirement for ADR reporting in India
11. ICH guidelines for ICSR, PSUR, expedited reporting, pharmacovigilance planning
12. CIOMS requirements for ADR reporting
13. Writing case narratives of adverse events and their quality.

Teaching scheme and examination scheme:

Teaching Scheme				Evaluation Scheme			
Theory	Tutorial	Practical	Total	Theory		Practical	
				External	Internal	External	Internal
3	1	0	4	80	20	0	0

Sr No	Topics	% weightage
1.	<p>Introduction to Pharmacovigilance</p> <ul style="list-style-type: none"> <input type="checkbox"/> History and development of Pharmacovigilance <input type="checkbox"/> Importance of safety monitoring of Medicine <input type="checkbox"/> WHO international drug monitoring programme <input type="checkbox"/> Pharmacovigilance Program of India(PvPI) <p>Introduction to adverse drug reactions</p> <ul style="list-style-type: none"> <input type="checkbox"/> Definitions and classification of ADRs <input type="checkbox"/> Detection and reporting <input type="checkbox"/> Methods in Causality assessment <input type="checkbox"/> Severity and seriousness assessment <input type="checkbox"/> Predictability and preventability assessment <input type="checkbox"/> Management of adverse drug reactions <p>Basic terminologies used in pharmacovigilance</p> <ul style="list-style-type: none"> <input type="checkbox"/> Terminologies of adverse medication related events <input type="checkbox"/> Regulatory terminologies 	10
2.	<p>Drug and disease classification</p> <ul style="list-style-type: none"> <input type="checkbox"/> Anatomical, therapeutic and chemical classification of drugs <input type="checkbox"/> International classification of diseases 	10



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	<input type="checkbox"/> Daily defined doses <input type="checkbox"/> International Non proprietary Names for drugs Drug dictionaries and coding in pharmacovigilance <input type="checkbox"/> WHO adverse reaction terminologies <input type="checkbox"/> MedDRA and Standardised MedDRA queries <input type="checkbox"/> WHO drug dictionary <input type="checkbox"/> Eudravigilance medicinal product dictionary Information resources in pharmacovigilance <input type="checkbox"/> Basic drug information resources <input type="checkbox"/> Specialised resources for ADRs Establishing pharmacovigilance programme <input type="checkbox"/> Establishing in a hospital <input type="checkbox"/> Establishment & operation of drug safety department in industry <input type="checkbox"/> Contract Research Organisations (CROs) <input type="checkbox"/> Establishing a national programme	
3.	Vaccine safety surveillance <input type="checkbox"/> Vaccine Pharmacovigilance <input type="checkbox"/> Vaccination failure <input type="checkbox"/> Adverse events following immunization Pharmacovigilance methods <input type="checkbox"/> Passive surveillance – Spontaneous reports and case series <input type="checkbox"/> Stimulated reporting <input type="checkbox"/> Active surveillance – Sentinel sites, drug event monitoring and registries <input type="checkbox"/> Comparative observational studies – Cross sectional study, case control study and cohort study <input type="checkbox"/> Targeted clinical investigations Communication in pharmacovigilance <input type="checkbox"/> Effective communication in Pharmacovigilance <input type="checkbox"/> Communication in Drug Safety Crisis management <input type="checkbox"/> Communicating with Regulatory Agencies, Business Partners, Healthcare facilities & Media	10
4.	Safety data generation <input type="checkbox"/> Pre clinical phase <input type="checkbox"/> Clinical phase <input type="checkbox"/> Post approval phase (PMS) ICH Guidelines for Pharmacovigilance <input type="checkbox"/> Organization and objectives of ICH <input type="checkbox"/> Expedited reporting <input type="checkbox"/> Individual case safety reports <input type="checkbox"/> Periodic safety update reports <input type="checkbox"/> Post approval expedited reporting <input type="checkbox"/> Pharmacovigilance planning <input type="checkbox"/> Good clinical practice in pharmacovigilance studies	8
5.	Pharmacogenomics of adverse drug reactions <input type="checkbox"/> Genetics related ADR with example focusing PK parameters. Drug safety evaluation in special population <input type="checkbox"/> Paediatrics <input type="checkbox"/> Pregnancy and lactation <input type="checkbox"/> Geriatrics CIOMS <input type="checkbox"/> CIOMS Working Groups <input type="checkbox"/> CIOMS Form CDSCO (India) and Pharmacovigilance <input type="checkbox"/> D&C Act and Schedule Y	7



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<input type="checkbox"/> Differences in Indian and global pharmacovigilance requirements
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Recommended Books (Latest edition):

1. Textbook of Pharmacovigilance: S K Gupta, Jaypee Brothers, Medical Publishers.
2. Practical Drug Safety from A to Z By Barton Cobert, Pierre Biron, Jones and Bartlett Publishers.
3. Mann's Pharmacovigilance: Elizabeth B. Andrews, Nicholas, Wiley Publishers.
4. Stephens' Detection of New Adverse Drug Reactions: John Talbot, Patrick Walle, Wiley Publishers.
5. An Introduction to Pharmacovigilance: Patrick Waller, Wiley Publishers.
6. Cobert's Manual of Drug Safety and Pharmacovigilance: Barton Cobert, Jones & Bartlett Publishers.
7. Textbook of Pharmacoepidemiology edited by Brian L. Strom, Stephen E Kimmel, Sean Hennessy, Wiley Publishers.
8. A Textbook of Clinical Pharmacy Practice -Essential Concepts and Skills: G. Parthasarathi, Karin Nyfort Hansen, Milap C. Nahata
9. National Formulary of India
10. Text Book of Medicine by Yashpal Munjal
11. Text book of Pharmacovigilance: concept and practice by GP Mohanta and PK Manna
12. <http://www.who.int/dynpage.aspx?id=105825&mn1=7347&mn2=7259&mn3=7297>
13. <http://www.ich.org/>
14. <http://www.cioms.ch/>
15. <http://cdsco.nic.in/>
16. http://www.who.int/vaccine_safety/en/
17. http://www.ipc.gov.in/PvPI/pv_home.html