



BRAINWARE UNIVERSITY

Term End Examination 2023

Programme – B.Pharm-2019

Course Name – Pharmacovigilance

Course Code - BP805ET

(Semester VIII)

Full Marks : 75

Time : 3:0 Hours

[The figure in the margin indicates full marks. Candidates are required to give their answers in their own words as far as practicable.]

Group-A

(Multiple Choice Type Question)

1 x 20=20

1. Choose the correct alternative from the following :

- (i) Identify the correct one, an IRB is also known as _____
- | | |
|---------|--------|
| a) ICH | b) IEC |
| c) DTAB | d) FDA |
- (ii) Indicate the correct one, In vitro study perform in or on _____
- | | |
|-----------------|--------------|
| a) Human | b) Animal |
| c) Living cells | d) Test tube |
- (iii) State the year in which Sulfanilamide tragedy occurred
- | | |
|---------|---------|
| a) 1838 | b) 1937 |
| c) 1951 | d) 1938 |
- (iv) Choose the appropriate full form of AMC
- | | |
|--------------------------------|------------------------------------|
| a) Allopathic motion center | b) ADR Monitoring Center |
| c) Alternative Medical Council | d) Ayurvedic Monitoring conference |
- (v) Choose the commonly reported ADR of diuretic class of drugs
- | | |
|-------------|----------------|
| a) Alopecia | b) Skin cancer |
| c) Rhinitis | d) Hypokalemia |
- (vi) Choose that happens to the case report forms (CRFs)
- | | |
|---|---|
| a) The CRF data is compiled and submitted to Regulatory Affairs | b) The CRF data is compiled and submitted to the FDA in the IND |
| c) The CRF data is aggregated by an external party if the trial was double blinded to assess the drug's safety and efficacy | d) The CRF data is aggregated and analyzed to assess the drug's safety and efficacy |
- (vii) Choose the appropriate full form of ICH is
- | | |
|--|---|
| a) Intermittent conference on harmonization | b) Intermittent council on harmonization |
| c) International conference on harmonization | d) International council on harmonization |
- (viii) choose the correct one in Pharmacovigilance activities, especially in preparation for the early post-marketing period of a new drug is known as _____planning.

- a) ABC
c) E2E Pharmacovigilance
- b) A2C
d) None
- (ix) Choose the schedule _____ is defined as the requirements and guidelines for permission to import and manufacture new drugs for sale or for clinical trials.
- a) Y
c) T
- b) M
d) A
- (x) Identify that the authorized advisory committee of Indian Govt. pertaining to Drug and Cosmetic act, is known as
- a) DTAB
c) CDSCO
- b) IPC
d) None
- (xi) Select the one is not related to the ADR type A
- a) It is predictable
c) It is pharmacodynamic effect
- b) It occurs due to genetic
d) It is dose related
- (xii) Identify that the ATC stands for :
- a) American Technical Council
c) Anatomical Theoretical Classification
- b) Anatomical Therapeutic Chemical Classification
d) Anatomical Therapeutic committee
- (xiii) Select the one is not related to Type B reaction
- a) Anaphylaxis
c) Drug allergy
- b) Dose
d) Idiosyncrasy
- (xiv) Select the correct one, An _____ is any untoward medical occurrence in a patient administered a medicinal product and which does not necessarily have a casual relationship with this treatment.
- a) Adverse event
c) Idiosyncrasy
- b) Adverse event
d) None
- (xv) Select the part of quality assurance which ensures that products are consistently produced and controlled to the quality standards appropriate to their intended use
- a) Drug master file
c) Inter company transfer
- b) Gap analysis
d) Good manufacturing practices
- (xvi) Identify the full form of ICSR
- a) WHO-UMC
c) International council for safety reports
- b) Indian council for survey and reports
d) Individual case safety reports
- (xvii) Identify that the nonproprietary name is also called _____.
- a) Classical
c) Generic
- b) Patent
d) Proper
- (xviii) Identify the name of the study of all gene variants/polymorphisms that influence patient response to given drug is known as _____.
- a) Pharmacodynamic
c) Pharmacokinetics
- b) Pharmacogenomics
d) Pharmacogenetics
- (xix) Indicate the number of groups working in CIOMS.
- a) 13
c) 4
- b) 8
d) 12
- (xx) Identify the correct one, Animal studies, clinical trials, bioavailability studies are part of which application process
- a) BLA
c) NDA
- b) ANDA
d) IND

Group-B

(Short Answer Type Questions)

5 x 7=35

2. Describe the safety Monitoring of Medicines (5)
3. Explain MedDRA and its standard queries. (5)

- 4. Explain CRO's and the importance of CRO's in the national program (5)
- 5. Describe specialized resources for ADRs (5)
- 6. Explain in detail about comparative observational studies (5)
- 7. State the importance of communication with Regulatory agencies and Business partners. (5)

OR

- Write the importance of communication in pharmacovigilance. (5)
- 8. Explain in detail about communication in drug safety crisis management (5)

OR

- Explain in detail the case-control study and cohort study (5)

Group-C

(Long Answer Type Questions)

10 x 2=20

- 9. compare ADR and ADE (10)
- 10. Explain in detail about Contact Research Organization. (10)

OR

- Explain in detail about Drug Event Monitoring and registries (10)
