

QP Code : 717401

(2 Hours)

[Total Marks : 35

- N.B. : (1) All questions are **compulsory**.
(2) **Figures** to the **right** indicate **full** marks.

1. Answer the following :

- (i) Define community pharmacy.
- (ii) Enlist any four reasons for patient non-compliance.
- (iii) Classify adverse drug reactions.
- (iv) Justify digitalis toxicity is increased on co-current administration with loop diuretics.
- (v) Enlist name of antimicrobials safe in pregnancy.
- (vi) Enlist categories of drugs which require therapeutic drug monitoring.
- (vii) State any one aim of pharmacovigilance.

2. (a) Answer **any one** of the following:

- (i) Define patient counselling. Explain with examples role of hospital pharmacist in patient counselling.
- (ii) Define clinical pharmacy. Explain scope and objectives of clinical pharmacy.

(b) Answer **any one** of the following :

- (i) Write in short methods of assessment of Compliance.
- (ii) Discuss role of pharmacist ensuring rational use of medication.

3. (a) Answer **any one** of the following:

- (i) Explain detection and reporting methods of Adverse drug reaction.
- (ii) Explain in brief manifestation of Adverse drug reaction.

(b) Answer **any one** of the following:

- (i) What are the strategies used for therapeutic drug monitoring.
- (ii) Discuss in brief criteria for valid therapeutic drug monitoring.

4. (a) Answer **any one** of the following:

- (i) Classify drug interactions. Explain drug interactions due to alterations in metabolism with suitable examples.
- (ii) Write a short note on drug-food interactions.

(b) Answer **any one** of the following:

- (i) Define paediatrics. Explain the factors affecting drug therapy in paediatrics.
- (ii) Discuss dose adjustment in geriatric patient with kidney failure.

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5. (a) Answer **any one** of the following:

- (i) What is double blind method used in clinical trial. Explain in detail phase III of clinical trial.
- (ii) Explain principles of ICH GCP

(b) Answer **any one** of the following:

- (i) Define the following term .
 - (a) New drug (as per schedule Y)
 - (b) Case report form. (CRF)
 - (c) Serious adverse event (SAE)
- (ii) Write a note on Bioequivalence studies.
