

[LI 825] APRIL 2016 Sub. Code: 3825

PHARM. 'D' AND PHARM. 'D' (POST BACCALAUREATE) DEGREE EXAMINATION (2009-2010 Regulation) FIFTH YEAR PAPER I – CLINICAL RESEARCH

Q.P. Code: 383825

Time: Three hours Maximum: 70 Marks

I. Elaborate on: $(4 \times 10 = 40)$

1. Explain the Pharmacological and Toxicological approaches to Drug discovery.

- 2. Describe the roles and responsibilities of Investigators and Clinical Research associates.
- 3. Discuss the challenges in the implementation of Good Clinical Practice guidelines.
- 4. Describe the guidelines of Central drug standard control organization.

II. Write notes on: $(6 \times 5 = 30)$

- 1. Abbreviated New Drug Application.
- 2. Various types of post marketing surveillance.
- 3. Ethical principles in Clinical research.
- 4. Explain the contents in Investigational new drug application.
- 5. Safety monitoring in Clinical research.
- 6. Responsibilities of Institutional review board.
