

 $3 \times 10 = 30 \text{ Marks}$

- 1. Discuss Regulatory requirement of ANDA generic drug approval in US.
- 2. Explain the *In vitro* drug product performance and its limitations.
- 3. Describe the regulatory guidelines for approval of biologics.
- 4. Discuss development of clinical trial protocol and add a note on working of institutional review board for clinical trials.

SHORT ESSAY (Answer any Nine)

9 X 5 = 45 Marks

- 5. What are the amendments of Hatch Waxman Act?
- 6. Discuss the importance of informed consent in clinical trial.
- 7. Discuss ICH guidelines related to maintenance.
- 8. Write a note on Master formula record and distribution records.
- 9. Discuss the regulatory requirements of investigator brochure.
- 10. Explain non clinical development related to global submission of NDA.
- 11. Discuss about CMC in regulations.
- 12. Explain the Regulatory requirements of ROW countries.
- 13. Write a note on post approval regulatory affairs.
- 14. Discuss regulation of combination products.

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