QP Code: 722006 Reg. No......

Seventh Semester B. Pharm Degree Regular/Supplementary Examinations February 2023 Industrial Pharmacy

(2017 Scheme)

Time: 3 Hours Max. Marks: 75

- Answer all questions to the point neatly and legibly
 Do not leave any blank pages between answers
 Indicate the question number correctly for the answer in the margin space
- Answer all parts of a single question together Leave sufficient space between answers
- Draw diagrams wherever necessary

Essays (2x10=20)

- 1. Explain technology transfer protocol and quality risk management in technology development and also process involved in technology transfer from R and D to production.
- 2. Explain the general consideration of pilot plant scale up techniques for the development of liquid orals forms with relevant documentation.

Short Notes (7x5=35)

- 3. Explain any two hazards and their industrial safety measures.
- 4. Explain SUPAC guidelines.
- 5. Explain organization and responsibilities of CDSCO.
- 6. What are confidentiality agreements and MoUs.
- 7. Explain the granularity of TT process involved in technology development and transfer.
- 8. Explain Data presentation for FDA submissions.
- 9. Explain Certificate of Pharmaceutical Product (COPP).

Answer Briefly (10x2=20)

- 10. List out the major utility and service systems used in pharma industry.
- 11. Importance of Investigators Brochure (IB) and NDA.
- 12. Define TIFAC and TBSE.
- 13. What are regulatory requirements and approval procedure for new drug.
- 14. List out the different modules of CTD.
- 15. What are general consideration of Investigational New Drug (IND) application.
- 16. Format of COPP.
- 17. Define technology transfer protocol.
- 18. Define accident records.
- 19. What are the steps involved in scale up process.
