near, labeled diagranis wherever necessary.

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- How are ICH guidelines organized by strategy the strategy explain by the First Ranker GOR lines of 1. ICH.
- 2. With a floor plan the layout of a production facility for solid oral dosage forms as mandated by Schedule M.
- 3. What is a Drug Master File? List its contents.
- 4. Explain the process of IPQC. Why is it essential? Illustrate the IPQ process for manufacturing process of a capsule formulation.

SHORT ESSAY (Answer any Nine)

9 X 5 = 45 Marks

3 X 10 = 30 Marks

- 5. What details should the protocol for a GLP study encompass?
- 6. Compare the scope of QA and QC
- 7. Explain the environmental control requirements for various production areas.
- 8. Give the IPQC requirements for topical dosage forms as per IP.
- 9. Explain the QC tests prescribed for ophthalmic products.
- 10. How do you audit your vendor for ensuring your purchase specifications?
- 11. Explain the weight variation test for solid oral dosage forms as per IP.
- 12. Explain the Common Technical Documentation triangle and it's modules.
- 13. Explain how you will compute expiry date of a drug product as per ICH requirements.
- 14. How will you handle disposal of waste products incurred during a pharmaceutical production process?

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