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Paper Id: 256104

Sub Code: MPH104T

Roll No.

M PHARM (SEM I) THEORY EXAMINATION 2018-19 REGULATORY AFFAIRS

Time: 3 Hours Total Marks: 75

Note: 1. Attempt all Sections. If require any missing data; then choose suitably.

SECTION A

1. Attempt all questions in brief.

 $10 \times 2 = 20$

- a. What do you mean by 'DMF'?
- b. State Hatch-Waxman Act.
- c. State the regulatory requirements for product approval of MHRA countries.
- d. Define 'CTD' and 'ECTD format'.
- e. Mention the role of IMPD in non clinical drug development.
- f. What do you mean by global submission of ANDA?
- g. Mention the functions of the Institutional Review Board.
- h. Mention the steps for developing clinical trial protocols.
- i. Explain the term 'CFR'.
- j. What do you mean by post approval regulatory affairs?

SECTION B

2. Attempt any two parts of the following:

 $2 \times 10 = 20$

- a. Describe the regulatory requirement for product approval of API and biologics.
- b. State and explain the regulation for combination products and medical devices.
- c. Explain the ANDA regulatory approval process.

SECTION C

3. Attempt any *five* parts of the following:

 $7 \times 5 = 35$

- a. Write short notes on 'master formula record' and 'code of federal regulation'.
- b. Describe the ways and means of US registration for foreign drugs.
- c. Write short notes on 'CTD' and 'ECTD format'.
- d. Write a brief note on 'non clinical drug development'.
- e. Describe the formulation and working procedure of Independent Ethics Committee.
- f. Write a brief note on 'informed consent process and procedures'.
- g. Write short note on 'industry and FDA liaison'.