

**B. PHARM.
EIGHTH SEMESTER
PHARMACEUTICAL REGULATORY SCIENCE
BP804ET [SPECIAL REPEAT]
[USE OMR SHEET FOR OBJECTIVE PART]**

**SET
A**

Duration : 3 hrs.

Full Marks : 75

[PART-A: Objective]

Time : 30 min.

Marks : 20

1×20=20

Choose the correct answer from the following:

- List of approved drugs and their associated IPR is available in _____
 - Pink book
 - Orange book
 - Red book
 - Indian pharmacopoeia
- Identify the relevant regulatory body in USFDA for approval of drugs
 - BLA
 - IND
 - CBER
 - CDER
- Rule 96 means-
 - Manner of packaging
 - Manner of importing
 - Manner of labelling
 - Manner of distributing
- How many types of DMF are there-
 - 5
 - 3
 - 2
 - 4
- Placebo means-
 - New medicine
 - Medically effectual treatment
 - Medically ineffectual treatment
 - Clinical study
- Drug regulatory body of Brazil is-
 - TGA
 - SFDA
 - MHLW
 - ANVISA
- Which regulatory body reviews and updates the Indian pharmacopoeia.
 - WHO
 - CDSCO
 - IPC
 - DTAB
- The Clinical trial legislative requirements are guided under-
 - Schedule Y
 - Schedule T
 - Schedule C
 - Schedule X
- On _____, the CTD became the mandatory format for NDA in the EU and Japan.
 - June 2003
 - January 2003
 - July 2003
 - July 2013

10. Comparative trial is also known as-
 - a. Three arm head-to-head study
 - b. Placebo
 - c. Two arm head-to-head study.
 - d. Phase I clinical study
11. IRB means
 - a. Institutional review board
 - b. Independent review board
 - c. Indian review board
 - d. International review board
12. Type I DMF deals with -
 - a. Packaging materials
 - b. Manufacturing site
 - c. Drug substance
 - d. Excipients
13. Who reviews,evaluates and approves the scientific and ethical aspects of a clinical trial
 - a. ICH
 - b. CPCSEA
 - c. IPC
 - d. IRB/IEC
14. The CFR is divided into -
 - a. 40 titles
 - b. 25 titles
 - c. 50 titles
 - d. 10 titles
15. Narcotic and psychotropic substances Act and rules was established in-
 - a. 1985
 - b. 1986
 - c. 1954
 - d. 1956
16. Informed consent form is also known as-
 - a. Informed consent document
 - b. Patient consent form
 - c. Both (a) and (b)
 - d. None of the above
17. The standard format of submitting the regulatory information to health authority is -
 - a. CTD
 - b. eCTD
 - c. DMF
 - d. ACTD
18. Which schedule specifies the general requirements for factory premises and materials-
 - a. Schedule Y
 - b. Schedule G
 - c. Schedule T
 - d. Schedule M
19. MHLW is a regulatory body of -
 - a. India
 - b. Europe
 - c. Japan
 - d. China
20. NPPA means-
 - a. National pharmaceutical pricing authority
 - b. National pharmacy product act
 - c. New pharmacy product advertisement
 - d. National public protection act

(PART-B : Descriptive)

Time : 2 hrs. 30 min.

Marks : 35

[Answer any seven (7) questions]

1. Write about the Drug and cosmetic Act. 5
2. What is an orange book, explain briefly. 1+4=5
3. Explain NDA and classify the drugs in NDA. 1+4=5
4. What a short note on eCTD. 5
5. Write about IRB/IEC mentioning its composition and procedures. 2+3=5
6. Write short notes on: 2.5+2.5
 - a. CDSCO =5
 - b. DTAB
7. Explain about "Concept of generics". 5
8. Write about 2.5+2.5
 - a. Narcotic and psychotropic Act. =5
 - b. Consumer Protection act.
9. Write about the rule no 96 from export of drugs in india. 5

-- -- --

PART-C: Long type questions

[Answer any two (2) questions]

1. Explain in details about the steps involved in developing clinical trial protocols. 10

2. Write short notes on: 5+5=10
 - a. Generic drugs
 - b. Purple book

3. Write and explain in details about the stages of drug discovery. 10

== *** ==