

**B. PHARM.
FIFTH SEMESTER
PHARMACEUTICAL JURISPRUDENCE
BP505T [SPECIAL REPEAT]
[USE OMR SHEET FOR OBJECTIVE PART]**

**SET
A**

Duration : 3 hrs.

Full Marks : 75

[PART-A: Objective]

Time : 30 min.

Marks : 20

Choose the correct answer from the following:

1×20=20

- Schedule P as per D&C Act deals with the following
 - Standard for cosmetics
 - Biological and special product
 - Life period of drug
 - Clinical Trial
- Requirements of factory premises for manufacture of medical devices is dealt under...
 - Schedule M1
 - Schedule M2
 - Schedule M3
 - Schedule N
- Blood Bank comes under the schedule
 - Schedule B
 - Schedule D
 - Schedule F
 - Schedule G
- If the drug contains in filthy, putrid or decomposed substance then is known as
 - Spurious Drug
 - Adulterated drug
 - Misbranded drug
 - None of the above
- "Schedule F3" is related with
 - Standard for surgical dressing
 - Standard for sterilized umbilical tapes
 - Standard for ophthalmic preparation
 - Standard for production of sera
- Who among the following is the Chairman of Drug Technical Advisory Board?
 - The Drugs Controller of India
 - The President of Pharmacy Council of India
 - The President of Medical Council of India
 - The Director General of Health Services
- Which amongst the following is the duty of a Drug Inspector ?
 - Seizure of stocks
 - Take samples of any drug or cosmetics
 - Inspection
 - All of the above
- Central drug laboratory is located at..
 - Mumbai
 - Calcutta
 - Delhi
 - Lucknow
- Schedule P is related to..
 - Packing of drugs
 - Toilet preparation
 - Ophthalmic preparation
 - Biologicals

10. The premises for manufacturing of drug must fulfil the condition according..
- Schedule N.
 - Schedule M
 - Schedule O
 - Schedule X
11. The Narcotic Drugs and Psychotropic Substances Act was passed in the year..
- 1940
 - 1955
 - 1985
 - 2000
12. What is the definition of a "Magic Remedy"?
- Talisman, mantra, kavacha or any other charm or any substance alleged to possess miraculous powers powers to diagnose, cure, mitigate, treat or prevent a disease in humans or animals
 - Any substance natural or synthetic or any salt or preparation of such substance or material, included in the list of psychotropic substances specified in the
 - Any substance, whether processed, partially processed or unprocessed, which is intended for human consumption.
 - Ethyl alcohol of any strength and purity having the chemical composition C₂H₅OH
13. Which of the following is/are non-patentable?
- A new medical device for diagnosis
 - Admixture of two known substances to give a new effect
 - Algorithms
 - A genetically modified microorganism
14. Plant of the species of Papaver, from which opium or any phenanthrene alkaloid can be extracted, is called..
- opium poppy
 - coca
 - heroin
 - Cannabis sativum
15. Who has the power to fix the ceiling price of scheduled formulations?
- State Government
 - Central Government
 - Lok Sabha
 - Rajya Sabha
16. NDPS Act provide licensing system to regulate.....of narcotic and psychotropic substances
- Manufacturing
 - Cultivation
 - Sale and export
 - All of the above
17. Manufacture of Opium can be made only by the Central Government at its two factories situated at..
- Ghazipur and Neemuch
 - Kolkata and Delhi
 - Mumbai and Lucknow
 - None
18. The non bonded laboratory shall be inspected by the proper Excise officer..
- Once in two months
 - Once in a month
 - Once in a year
 - Once in six months

9. Discuss the advertisements which are prohibited under objectionable advertisement. 5

PART-C: Long type questions

[Answer any two (2) questions]

1. Elaborately discuss loan license and Repacking license 5+5=10
2. Elaborately discuss the Good Manufacturing Practises (GMP) required for factory premises. 10
3. Write the constitution of IAEC and describe the CPSCEA guidelines as per the prevention of cruelty to Animals act, 1960 10

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