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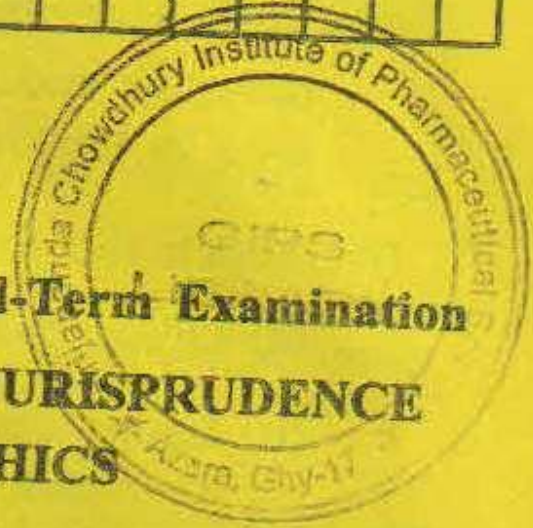
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Roll No. of candidate

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2016

B. Pharm 6th Semester End-Term Examination
PHARMACEUTICAL JURISPRUDENCE
AND ETHICS



Full Marks-100 Pass Marks-35 Time-Three hours

The figures in the margin indicate full marks for the questions.

1. Answer any six of the following : 2×6=12
- (i) Match the following :
- | | |
|--|--------------------------------|
| (a) Vending license is not issued for | (i) Schedule H drugs. |
| (b) Physician sample should not be supplied for | (ii) Schedule J drugs. |
| (c) A drug may not purport to prevent or cure | (iii) Schedule C and C1 drugs. |
| (d) A drug not to be sold without a prescription | (iv) Schedule X drugs. |

[Turn over

(ii) What are the circumstances due to which the name of a person is removed from the register of State Pharmacy Council ?

(iii) What are the controlled operations of narcotics and psychotropic substances powdered to the State Government ?

(iv) Match the following :

(a) Drugs Bill (i) 1930

(b) Drugs Enquiry Committee (ii) 1940

(c) Pharmacy Council of India (iii) 1943

(d) Health Survey and Development Committee (iv) 1949

(v) Fill in the blanks :

(a) Drug Inspectors are appointed under the Section of D&C Act, 1940.

(b) Manufacturing of opium is carried out at the places and

(vi) Name the government laboratories under D & C Act, responsible for the test and analysis of veterinary products and oral polio vaccine.

(vii) Define the terms- magic remedy, advertisement.

2. Answer any six of the following : 6×3=18

(i) How the ophthalmic preparations are packaged ? What are the quantities of dose permitted in a single packing of Schedule X drugs ?

(ii) What do you mean by spurious and misbranded drugs ?

(iii) What do you mean by 'illicit traffic' under NDPS act and rules ?

(iv) What are the essential requirements of the premise involved in manufacturing outside bond ?

(v) Define : ceiling price, pre-tax return and sale turnover.

(vi) What are coca derivatives and opium derivatives ?

(vii) What do you know about General Agreement on Trade and Services (GATS) and GATT ?

3. Answer any eight of the following : 8×5=40

(i) Give the composition of Drug Technical Advisory Board (DTAB). What are its functions ?

- (ii) Discuss the provisions in the Factories Act, 1948.
- (iii) Give the Pharmacist's Oath as approved by PCI.
- (iv) What are the circumstances under which pregnancies can be terminated ?
- (v) Write briefly about Prevention of Cruelty to Animal act, 1960.
- (vi) What are the directions in the Minimum Wages Act, 1948 ?
- (vii) Explain the regulations under the Poison Act, 1919.
- (viii) Explain manufacturing in bond.
- (ix) What categories of advertisements are banned or prohibited under the Drugs and Magic Remedies Act ?
- (x) How the retail price of drug formulations is calculated ?

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(4)

120(Y)

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(5)

120(Y)

4. Answer any *three* questions. (Essay type questions) : 3×10=30
- (i) (a) What are the required qualification and duties of government analyst ?
 - (b) Discuss the offences and penalties under narcotic drugs and psychotropic substances act. 5+5=10
 - (ii) (a) Discuss the provisions pertaining to cosmetics under the D&C Act 1940 and rules 1945.
 - (b) Write a note on manufacturing of Homeopathic medicines. 6+4=10
 - (iii) (a) Explain manufacturing of Schedule C, C1 and Schedule X drugs.
 - (b) Draw an indicative label of a Schedule X drug. 7+3=10
 - (iv) (a) What is intellectual property right ? What are the inventions not patentable ?
 - (b) Give a brief account of historical development of pharmaceutical legislations. 5+5=10