X(4th Sm.)-Chemistry-H/CC-8/CBCS

2022

CHEMISTRY — HONOURS

Paper : CC-8

(Organic Chemistry - 4)

Full Marks : 50

The figures in the margin indicate full marks. Candidates are required to give their answers in their own words as far as practicable.

Answer question no. 1 and any eight questions from the rest (question no. 2 to 13).

1. Answer any ten questions :

1×10

(a) Give the product(s) of the following reaction :

$$H_2C = CH - CH_2$$

 $H_2C = CH - CH_2$
 $CH_2 - CH = CH_2$
 $Heat$

- (b) The IR spectrum of benzene shows many peaks but its UV spectrum is very simple. Explain.
- (c) Phenol is directly converted to anisole on reaction with diazomethane but an aluminium alkoxide catalyst is required to convert ethanol to ethyl methyl ether with the same reagent. Explain.

(d) [B]
$$\checkmark$$
 Na/ether CH₃CH₂CN $\frac{1. \text{ CH}_3\text{Mg I/dry ether}}{2. \text{ H}^{\oplus}/\text{H}_2\text{O}}$ [A]

Give the structures of [A] and [B] (structures only).

- (e) The difference in precessional frequency of a proton from TMS is 186 Hz in a 60 MHz NMR machine. Find its δ value.
- (f) How do you protect propane-1,3-diol? Write down also the deprotecting agent.
- (g) Explain why the normal isotope of carbon, ¹²C is NMR inactive.
- (h) Write down the products (only write down the structures of the products) obtained by diazocoupling of benzenediazonium chloride with alkaline 2-naphthol and aniline separately.
- Explain why cis- cinnamic acid absorbs at a higher frequency than its trans- isomer in the IR spectrum.
- (j) Give one example of each of the following :
 - (i) Illogical electrophile
 - (ii) Illogical nucleophile

Please Turn Over

X(4th Sm.)-Chemistry-H/CC-8/CBCS

- (k) Write down the structures of the products when RCOOH and $R_2C = O$ are separately subjected to Schmidt reaction.
- (l) Write down the synthetic equivalents corresponding to the following synthons :

(i) : ČHO (ii) Ph

- 2. (a) How is *threo* (active) isomer of butane-2,3-diol be distinguished from its *erythro* (*meso*) isomer by IR spectroscopy?
 - (b) Explain why is tetramethylsilane (Me₄Si) (TMS) used as an internal standard in NMR spectral studies. Write down the unit used to measure coupling constant. 3+2

OH

(b) Show the retrosynthetic pathway and the synthesis of target molecule (TM) as follows : 3+2

(T.M)

4. (a) $+ CH_3 - CH = CH_2 \xrightarrow{H_3PO_4} [G] \xrightarrow{O_2/OH} [H]$ (High Pressure) $|H_2SO_4, 100^{\circ}C$ |H] $|H_2SO_4, 100^{\circ}C$

Give the structures of [G], [H], [I] and [J]. Show the mechanism involved in conversion of [H] to [I] and [J]

(b) Give the product(s) of the following reaction along with the mechanism involved. 3+2



- 5. (a) Define stereospecific and stereoselective reactions and justify the difference between the two terms with the example of addition of singlet and triplet carbene to Z-2-butene.
 - (b) Primary and secondary nitroalkanes can take part in Nef carbonyl synthesis, but tertiary nitroalkanes can not. Explain. 3+2
- 6. (a) Explain why anisole with a mixture of nitric and sulphuric acid gives *o*-nitroanisole in 31% yield whereas with $HNO_3 Ac_2O$ gives the same product in 71% yield. Provide a suitable mechanism to justify the above observation.
 - (b) Show how a single reagent can be used to distinguish between primary, secondary and tertiary aromatic amines (No mechanism is needed). 3+2
- 7. (a) Account for the following trends in $\lambda_{max}(nm)$: ethylene (175), 1.3-butadiene (217); and 1.3,5-hexatriene (250). Explain why 1,5-hexadiene ($\lambda_{max} = 185 \text{ nm}$) does not absorb light above 200 nm.
 - (b) Discuss the difficulties of synthesising $Me_3C NH_2$ by Gabriel phthalimide synthesis. Show how Me_3CNH_2 can be prepared from $Me_3C OH$. 3+2
- 8. (a) Using Woodward-Fieser rule, calculate λ_{max} of the UV absorption for the following compounds :



(b) Give the product(s) of the following reaction along with the mechanism involved. 3+2



 (a) An organic compound with molecular formula C₆H₁₂O gives positive iodoform test. Its IR and ¹H NMR spectral data are as follows.

IR : $v_{cm^{-1}} = 1710 \text{ cm}^{-1} \text{ (strong)}$

¹H NMR : $\delta_{2,1}$ (3H, s) and 1.1 (9H, s).

Deduce the structure of the molecule with proper justification.

(b) What is the range of 'finger print region' in IR spectroscopy in cm⁻¹? Justify the naming of this range. 3+2

Please Turn Over

X(4th Sm.)-Chemistry-H/CC-8/CBCS

(4)

10. (a) The following reaction gives a single product. Give the structure of the product and also e_{xplain} mechanistically why the other isomeric product is not formed.



(b) Use Felkin-Anh's model to determine the stereochemistry of the major product of the following reaction : 3+2



11. (a) Give the retrosynthetic pathway followed by the synthesis of the following target molecule (TM):



(b) Give the product(s) of the following reaction along with the mechanism involved : 3+2



12. (a) $Ph_2C - CMe_2 + Ph_2C - CEt_2 - H^{\oplus}$ [K] + [L] OH OH ОН ОН (II) (I)

Explain the products of the reaction mechanistically.

(b) Give possible modes of retrosynthetic analysis and efficient synthesis for



Which mode is better choice?

3+2





(b) Define the terms 'chemically equivalent' and 'magnetically equivalent' used in ¹H NMR spectroscopy. Give an example of a molecule with chemically equivalent but magnetically non-equivalent protons with proper justification. 3+2