## 11.

## ALKENES AND ALKYNES

## UNSATURATED HYDROCARBONS

Contain carbon-carbon double or triple bonds (more hydrogen can be added).

## 1. INTRODUCTION OF ALKENES

## Introduction - Structure and Bonding

- Alkenes are also called olefins, contain a carbon-carbon double bond.
- General formula: $\mathrm{C}_{\mathrm{n}} \mathrm{H}_{2 n}$ (for one double bond)
- Suffix $=-$ ene


## Calculating Degrees of Unsaturation

1. An acyclic alkene has the general structural formula $\mathrm{C}_{\mathrm{n}} \mathrm{H}_{2 n-2}$.
2. Cycloalkanes also have the general formula $\mathrm{C}_{\mathrm{n}} \mathrm{H}_{2 \mathrm{n}}$.
3. Each $\pi$-bond or ring removes two hydrogen atoms from a molecule and this introduces on degree of unsaturation.
4. The number of degrees of unsaturation for a given molecular formula can be calculated by comparing the actual number of H atoms in a compound and the maximum number of H atoms possible.
5. This procedure gives the total number of rings and or $\pi$ bonds in a molecule.

## 2. NOMENCLATURE OF ALKENES

## IUPAC Nomenclature of Alkenes

Step 1: Identify and name the longest continuous chain of $C$ atoms which contains the double bond(s) (\# C + -ene).
Step 2: Number the longest chain so that the C's joined by the double bond have the lowest numbers possible. If the double bond has the same position starting from either end, use the positions of the substituents to determine the beginning of the chain.
Step 3: Locate the double bond by the lower numbered C-atom joined by the double bond (e.g. 1-butene).
Step 4: Locate and name attached groups.
Step 5. Combine the names for the attached groups and the longest chain into the name.

If there is more than one double bond:

- A counting prefix (di-, tri-, tetra-, etc.) is placed immediately in front of the suffix -ene to indicate the number of double bond (diene, triene, tetraene, etc.).
- Usually, an " $a$ " is placed before the counting prefix to make pronunciation easier (e.g. butadiene).
- The starting position of each double bond is indicated by the lower number, separated by commas (e.g. 1, 3-butadiene).
- For cycloalkenes, the ring is named as cyclo- + \#C + -ene; one of the carbons of the double bond must be numbered "1".


## Nomenclature of Alkenes

Some alkene or alkenyl substituents have common names.
The simplest alkene, $\mathrm{CH}_{2}=\mathrm{CH}_{2 n}$ named in the IUPAC system as ethene, is often called ethylene.

Illustration 1: Name of the following as substituted derivatives of ethylene.
(JEE MAIN)
a. $\mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}-\mathrm{CH}_{3}$
b. $\quad \mathrm{Cl}_{2} \mathrm{C}=\mathrm{CHCl}$
c. $\left(\mathrm{CH}_{3}\right)_{2} \mathrm{C}=\mathrm{CH}_{2}$
d. $\left(\mathrm{CH}_{2}=\mathrm{CH}\right)_{2} \mathrm{C}=\mathrm{CH}_{2}$
e. $\left(\mathrm{CH}_{2}=\mathrm{CH}\right)-\mathrm{CH}=\mathrm{CH}-\left(\mathrm{CH}=\mathrm{CH}_{2}\right)$

Sol: When both the carbons contain same substituents use the symmetrical (sym) and when both substituents are on the same C atom use unsymmetrical (unsym)
a. Sym-Dimethyl ethylene
b. Trichloroethylene
c. unsym-Dimethyl ethylene
d. unsym-Divinyl ethylene
e. sym-Divinyl ethylene

Illustration 2: Write the IUPAC names of the following.
a.


b.

c.

(JEE ADVANCED)

Sol: (a)


4-Bromo-3-(1-methylpropyl)-7-methyl-1, 5-octadiene
Or
4-Bromo-3-(sec-butyl)-7-methyl-1, 5-octadiene
(b)

Choose the longest chain with two $(C=C)$. There $(C=C)$ cannot be taken into a single chain 3-Ethenyl-1, 5-heptadiene.
(c)


## 3. HYBRIDIZATION OF ALKENES

1. When a carbon is connected to three other atoms (that is, one of the bonds is a double bond), the molecules are modelled by combining the 2 s and two of the 2 p -orbitals to produce three $\mathbf{s p}^{2}$-orbitals.
2. Since only two of $2 p$ orbitals where hybridized, there is one leftover $p$ orbital in an $\mathrm{sp}^{2}$-hybridized carbon atom.

## Introduction - Structure and Bonding

Recall that the double bond consists of a $\pi$-bond and a $\sigma$-bond.
Each carbon is $\mathrm{sp}^{2}$ hybridized and trigonal planar, with bond angles of approximately $120^{\circ}$.

## 4. ISOMERIZATION

## Geometric Isomers in Alkenes

1. Free rotation is not possible around double bonds, therefore there are two different forms of 2-butene, which are geometric isomers of each other.
2. The prefix cis- is used when the two arms of the longest chain are on the same side of the double bond; the prefix trans- is used when they are on opposite sides of the double bond.
3. Geometric isomers can have drastically different chemical and physical properties.

cis-2-butene

trans-2-butene

Illustration 3: Which of the following do not show G.I.?
a.

b.

c.

d.


Sol: In order to show Geometrical isomerism, the alkene must contain different substituents.
(a) Two


Its show G.I.
(b)


Two different group
Option (b) does not show G.I
(c) Option (c) show G.I. at the double bond between $C_{1}-C_{2}$ but does not show G.I. at the double bond between $\mathrm{C}_{4}-\mathrm{C}_{5}$.

or

(d)


Shows G.I
Shows G.I

Illustration 4: From the following G.I. shown by:
(a)

(b)

(c)

(d)


Sol: In order to show Geometrical isomerism the alkene must contain different substituents .i.e. the substituents on both the carbon bearing double bonds must have different substituents.
(JEE ADVANCED)
(a)

(b)

Does not show G.I
(c)

(d)


## 5. PREPARATION OF ALKENES

### 5.1 Dehydration of alcohols



Ease of dehydration of alcohols: $3^{\circ}>2^{\circ}>1^{\circ}$, because the alkenes formed are more stable.
Various dehydrating agents can be used for the dehydration of alcohols:

1. Conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$ at $443-453 \mathrm{~K}\left(170-180^{\circ} \mathrm{C}\right)$ acts as Lowry Bronsted acid $\mathrm{H}^{\oplus}$ donor.
2. $\mathrm{H}_{3} \mathrm{PO}_{4}($ Phosphoric acid at 470 K$)$.
3. $\mathrm{Al}_{2} \mathrm{O}_{3}$ at $633 \mathrm{~K}\left(380^{\circ} \mathrm{C}\right)$ acts as a Lewis acid.
4. Heating with $\mathrm{P}_{2} \mathrm{O}_{5}$ or $\mathrm{P}_{4} \mathrm{O}_{10}$ (phosphorous pentoxide).
5. Heating with $\mathrm{POCl}_{3}$ (phosphorous oxychloride) + pyridine.
6. Heating with $\mathrm{KHSO}_{4}$.

Stability order of alkene: More substituted alkenes is more stable (due to hyperconjugation).



(Stabilisation by extended conjugation)

(Due to ring stability)

Mechanism of dehydration of $1^{\circ}$ alcohols: $\mathrm{E}_{2}$ mechanism


1. The first step in the dehydration of $1^{\circ}$ alcohols is protonation.
2. The second step is the attack of $\mathrm{C}_{\mathrm{B}}$ (conjugated base) at -proton $\left(-\mathrm{OH}_{2}{ }^{+}\right)$and the elimination of protonated hydroxyl group (a good leaving group) takes place simultaneously to form double bond.
Rate $=\mathrm{k}\left[\begin{array}{l}\ddot{\theta} \\ \mathrm{A}\end{array}\right]$ [protonated alcohol]
3. Kinetics of the reaction is of the second order and bimolecular.
4. Two eliminating groups (i.e., H and OH groups) must be in anti-position.
5. Reaction is stereospecific and stereo selective (or regioselective).
6. No rearrangement takes place.

Illustration 5: Arrange the following alcohols in the decreasing order of their ease of dehydration with $\mathrm{H}_{2} \mathrm{SO}_{4}$.
a.

b.

c.

d.

e. $\mathrm{Me} \sim(\sim \mathrm{OH}$
(JEE MAIN)

Sol: Order of stability of carbocation: $3^{\circ}>2^{\circ}>1^{\circ}$
Depending upon the formation of carbocation we can explain the order of reactivity.
We can see that options (b) and (c) are $3^{\circ}$ alcohols, since dehydration of alcohol proceeds through the formation of carbocation, we have to take into consideration the stability of carbocation. $3^{\circ}$ carbocation in (c) is more stable due to resonance and gives a more substituted alkene.
(b)

(c)



Remaining (a) and (d) both are $2^{\circ}$ alcohols, $2^{\circ}$ carbocation can be stabilized to $3^{\circ}$ via $1,2-\mathrm{H}^{\ominus}$ shift and gives a more substituted alkene
(d)


So, the order of dehydration is: $(\mathrm{c})>(\mathrm{b})>(\mathrm{d})>(\mathrm{a})>(\mathrm{e})$

Illustration 6: Give the products of the reaction with conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$ of:
(JEE ADVANCED)
(a)

(b)


Sol: (a)




Saytzeff product
(b)





Benzyl $C^{\oplus}$ (more stable than $\left.3^{\circ} \mathrm{C}\right)^{\oplus}$

(A)

3-Methyl-1-phenyl but-2-ene


### 5.2 Rearrangement and Ring Expansion

Rearrangement results in the substitution, elimination, and ring expansion or ring contraction. In case of cyclic alcohols, whenever a carbocation is formed outside, the next $C$ atom of a ring, ring expansion may take place, if a relief in the ring strain occurs.

## Examples of Ring Expansion:

1. 



2.




(More stable than but-1-ene)



Whenever a carbocation is formed on the ring, ring contraction takes place. But in this case, ring strain is not relieved (because higher homologue cyclic ring has to change to lower one), however, the products corresponding to ring contraction are formed in smaller amounts.

### 5.3 From Alkynes

1. By partial reduction of alkynes. The catalytic hydrogenation of alkynes to alkenes occurs faster than that of alkenes to alkanes. Therefore, by using a specific catalyst, it is possible to stop the reduction to give cis-or transalkenes depending upon the nature of the catalyst used.
(i) Cis-alkenes are formed by hydrogenation in presence of (i) $\mathrm{Pd} / \mathrm{BaSO}_{4}$ or $\mathrm{CaCO}_{3}$ poisoned with $\mathrm{PbCO}_{3}$ or Lindlar's catalyst(quinolone) (ii) $\mathrm{B}_{2} \mathrm{H}_{6}$ in THF/ ether $+\mathrm{CH}_{3} \mathrm{COOH}$.
(ii) Trans-alkenes are formed by hydrogenation in presence of $\mathrm{Na}+$ liq. $\mathrm{NH}_{3}$ (Birch reduction)
(a) With terminal alkyne.


(b) With internal alkynes:


Illustration 7: Explain
(JEE ADVANCED)


Sol: Here the reactant contains a chiral center hence it is optically active. On reduction with palladium it produces cis product hence the chirality is lost thus it is optically inactive. (Carbon contains two of the same groups)

On birch reduction Trans addition takes place thus product is trans and it is optically active. It gives two product d and I.


Illustration 8: Identify $A$ and $B$


Gas (B) gives test for unsaturation, i.e. it decolourises $\mathrm{Br}_{2}$ water and aq. $\mathrm{KMnO}_{4}$ solution.
Sol: From the given data it appears that the starting compound contains 6 carbon in its backbone. (Since product formed is butene (4 carbon) and 2 molecule of $\mathrm{CO}_{2}$ (2 carbon).

Thus the compound (A) should be 6 C-atom dibasic acid (four C atoms of $(B)$ or $(C)$ and two $(\mathrm{COOH})$ groups.
The first step is Kolbe's Electrolysis which proceeds through free radical mechanism.
The two possible structure of the dibasic acids are as follows:

OR




As given B gives test for unsaturation, therefore, (A) can be (II).
Possible reaction is as follows:


Disodium-2,3-dimethyl
Butane-1,4-dioate


cis or trans-But-2-ene

At the cathode, the reduction of $\mathrm{H}_{2} \mathrm{O}$ takes place (since reduction potential of $\mathrm{H}_{2} \mathrm{O}>$ reduction potential of $\mathrm{Na}^{+}$ion).
$\mathrm{H}_{2} \mathrm{O}+\mathrm{e}^{\Theta} \rightarrow \stackrel{\Theta}{\mathrm{O}} \mathrm{H}+1 / 2 \mathrm{H}_{2}(\mathrm{~g})$

### 5.4 From Alkyl Halides

From alkyl halides or haloalkanes: Alkyl halides on heating with a strong base such as sodium ethoxide or concentrated alcoholic solution of potassium hydroxide undergo dehydrohalogenation to give alkenes.


Alkyl halide



This process of the removal of a molecule of a halogen halide ( $\mathrm{HCl}, \mathrm{HBr}$ or HI ) from a haloalkane to form an alkene is called dehydrohalogenation. Dehydrohalogenation is an example of an elimination reaction. Since, in this reaction, a hydrogen is removed from $\beta$-carbon and halogen from the $\alpha$-carbon, therefore, it is called $\beta$-elimination reaction. The ease of dehydrohalogenating of alkyl halides having the same alkyl group but different halogens is: iodides > bromides > chlorides while for isomeric alkyl halides having the same halogen but different structures is: tertiary > secondary > primary. Thus, a tertiary alkyl iodide is most reactive.

Saytzeff rule: Depending upon the structure, alkyl halides may give one or more isomeric alkenes. For example, dehydrohalogenation of 1 -chlorobutane gives only one alkene, i.e., but-1-ene since only one type of $\beta$-hydrogen is available on the left side of the molecule.


1-Chlorobutane

1. When a bulky base is used, e.g. potassium tert-butoxide in tert-butyl alcohol or $\mathrm{Et}_{3} \ddot{\mathrm{~N}}, f$ for the dehydrohalogenation of RX, the less-substituted alkene (Hofmann product) is favoured.



It is due to the steric hindrance of the bulky base. The large tert-butoxide ion finds difficulty in removing one of the internal $2^{\circ} \mathrm{H}$ atoms because of greater steric hindrance (or crowding) at that site in the transition state (T.S.). It removes one of the more exposed $1^{\circ} \mathrm{H}$ atoms of the methyl group and gives less-substituted alkene, i.e. follows Hofmann's rule.

If the RX contains one or more double bonds, then the Saytzeff product is not formed, instead a product containing a conjugated double bond is formed rather than containing an isolated double bond, since conjugated double bonds are more stable than isolated double bonds.

When the more substituted alkene obtained by Saytzeff rule is sterically hindered, the less-substituted alkene is formed in a major amount.

Stereo chemical requirement: For $\mathrm{E}_{2}$ dehydrohalogenation, two eliminating groups must be in trans position and must yield anti-elimination reaction.
(i) Anti-elimination can occur in two ways with cis-isomer.

(ii) Anti-elimination can occur in one way with the trans- isomer.


Dehalogenation Geminal (1, 1-dihalogen) and vicinal (1, 2-dihalogen) derivatives of alkane with (Zn dust +EtOH or $\mathrm{CH}_{3} \mathrm{COOH}$ ) or $\mathrm{KI}+$ acetone or ( $\mathrm{Na}+$ ether) give alkene.



## $\mathrm{E}_{2}$ Dehalogenation is anti

Reaction is stereospecific
Note: The presence of tert-butyl group on equatorial position does not allow the flipping of the equatorial Br to axial Br.



A forms alkene $B$ does not.

Illustration 9: Identify all the possible alkenes that would be formed on the dehydrohalogenation of the following organic halides with alcoholic KOH. Also, identify the major alkene.
(a). 1-Chloropentane
(b). 2-Chloropentane
(c). 2-Chloro-2-methyl butane
(JEE ADVANCED)
Sol: Identify the carbocation that is going to form during the reaction and accordingly predict the product (minor and major).
Order of stability of carbocation - tert> sec>primary.
(a)

(b)




2-Methyl but-2-ene (Major)

2-Methyl but-1-ene (Minor)

Illustration 10: Predict the order of reactivity of the following compounds in dehydrohalogenation.
(JEE MAIN)
I.

II.

III.

IV.

V.


Sol: Order of formation of carbocation is $3^{\circ}>2^{\circ}>1^{\circ}$
Correct order of reactivity is (V) $>$ (III) $>$ (II) $>$ (IV) $>$ (I)


(III)

(II) and (III) both involves formation of tertiary carbocation but ease of formation of carbocation in (III) is more than in (II), because the reactivity order of RX is $\mathrm{RI}>\mathrm{RBr}>\mathrm{RCI}>\mathrm{RF}$ due the leaving group order of $\mathrm{I}^{\ominus}>\mathrm{Br}^{\ominus}>\mathrm{Cl}^{\ominus}>\mathrm{F}^{\ominus}$
(IV)

(V)


Illustration 11: Given the major and minor products of the following reactions.

b.

(JEE ADVANCED)
Sol: Depending upon the base used, predict the product. If the bulky base is used, the Hoffmann product (less substituted alkene, less stable) will predominate. If the non- bulky base is used, Saytzeff product (more substituted, more stable) will predominate.
(a) Here RX is RF, thus less-substituted alkene (Hofmann elimination) is formed, irrespective of whether the base is bulky or non-bulky. Both (A) and (B) are same.

(b) (A) Base used is non bulky thus the Saytzeff product is formed (More-substituted alkene) Me Me
(B) Base used is bulky thus the Hofmann product is formed (Less-substituted alkene)


### 5.5 Kolbe's Electrolytic Method

Electrolysis of sodium or potassium salts of saturated dicarboxylic acids gives alkenes. For example,


Pot. succinate
Ethylene

The reaction is believed to occur by the following steps:


At cathode. Since the electrode potential of $\mathrm{H}^{+}$ions is much higher than that of $\mathrm{K}^{+}$ions, therefore, $\mathrm{H}^{+}$ions are preferentially reduced to produce $\mathrm{H}_{2}$ while $\mathrm{K}^{+}$ions remain in the solution.
$2 \mathrm{H}^{+}+2 \mathrm{e}^{-} \longrightarrow[2 \mathrm{H}] \longrightarrow \mathrm{H}_{2}$

## 6. PHYSICAL PROPERTIES OF ALKENES

1. Only weak Vander Waals interactions, have low melting points and boiling points which increases with increased in surface area as the number of carbon increases.
2. Alkenes are soluble in organic solvents and insoluble in water.
3. Cis-2-Butene has a higher boiling point $\left\{4^{\circ} \mathrm{C}\right\}$ than trans-2-butene $\left\{1^{\circ} \mathrm{C}\right\}$ because in cis isomer, two C-C bond dipoles rain force each other yielding a small net molecular dipole.
4. Range of physical states: $\leq 4 \mathrm{C}$ 's - gases, $5-17 \mathrm{C}$ 's-liquid, $\geq 18 \mathrm{C}$ 's-solids.
5. Less dense than water as a result floats on water. Flammable and non-toxic.

### 6.1 Boiling Point

Boiling points of all these hydrocarbons increase with the increase in the carbon content; for each added $-\mathrm{CH}_{2}-$ ) group, boiling point rises from $20^{\circ} \mathrm{C}$ to $30^{\circ} \mathrm{C}$. Because of the presence of $\pi$-bonds, these molecules are slightly polar and, hence, have a higher boiling points than those of the corresponding alkanes.
In alkenes that show geometrical isomerism, generally boiling point of cis form is higher than the trans form due to the high polarity of the cis isomer compared to its trans isomer.


Net dipole
It shows dipole moment (Hence, slightly polar) b.p. ( 277 K )


Dipole moment $=0$
(Non-polar)
b.p. ( 274 K )
(All vectors cancel)

cis-Pent-2-ene
(A)

trans-Pent-2-ene
(B)

Dipole moment of $(A)>$ Dipole moment of $(B)$
(A) is more polar than (B)
b.p. of > b.p. of (B)

In alkenes that show geometrical isomers, trans forms have higher melting points than cis forms because the more symmetrical trans forms pack closely in the crystal lattice.

Similarly, out of $\mathrm{o}^{-}, \mathrm{m}^{-}$and p -xylenes, the p -isomers that are most symmetrical have the highest melting point.

### 6.2 Solubility

The solubility in non-polar solvents are in the following order: alkanes > alkenes > alkynes. Alkanes and alkynes are thereby, only very slightly soluble in water (with alkynes being slightly more soluble than alkenes). The densities of alkenes and alkynes are lower than that of water.

Illustration 12: i. Predict the more stable alkene of each pair.
ii. For which pairs could you use $\Delta \mathrm{H}_{\mathrm{h}}^{\circ}$ ?
iii. For which pairs could you use $\Delta \mathrm{H}_{\mathrm{C}}^{\circ}$ to determine their relative stabilities?
I. (a) 2-Methyl-pent-2-ene
(b) 2,3-Dimethyl-but-2-ene $\Delta \mathrm{H}_{\mathrm{C}}^{\circ}$
II. (c) Hexene
(d) cis-Hex-3-ene
III. (e) trans-Hex-2-ene
(f) 2-Methyl-pent-2-ene
IV. (g) cis-Hex-3-ene
(h) trans-Hex-3-ene
(JEE MAIN)

## Sol:

(I)

$\Delta H_{h}^{\circ}$ cannot be used because the same alkane is not produced on hydrogenation. So, $\Delta H_{C}^{\circ}$ is used to determine the relative stabilities of the alkene pairs because on complete combustion, the alkene requires same moles of $\mathrm{O}_{2}$ and produces same moles of $\mathrm{CO}_{2}$ and $\mathrm{H}_{2} \mathrm{O}$ (same formula: $\mathrm{C}_{6} \mathrm{H}_{12}$ ).
(II)

$\Delta H_{h}^{\circ}$ is used to determine the relative stabilities of the alkene pair because on hydrogenation two alkenes would yield the same alkane.
(III)

$\Delta H_{C}^{\circ}$ is used, since it will not produce the same alkane on hydrogenation but on combustion alkene requires same moles of $\mathrm{O}_{2}$ and produces same moles of $\mathrm{CO}_{2}$ and $\mathrm{H}_{2} \mathrm{O}$ (same formula: $\mathrm{C}_{6} \mathrm{H}_{12}$ ).


## 7. REACTIONS OF ALKENES

### 7.1 Hydrogenation of Alkenes

Hydrogenation of alkenes can be carried out by the following reagents:

1. Pt or Pd or Ni or $\mathrm{PtO}_{2}$
2. $\mathrm{Pd}+\mathrm{C}$ or $\mathrm{Pt}+\mathrm{C}(\mathrm{C} \Rightarrow$ charcoal $)$, Wilkinson's catalyst $\left[\mathrm{RhCl}_{\left.\left(\mathrm{Ph}_{3} \mathrm{P}\right)_{3}\right] \text { Chlorotris (triphenyl phosphine) rhodium (I) }}\right.$
3. $\mathrm{NaBH}_{4}+\mathrm{AlCl}_{3}$ in diglyme (solvent)
4. $\mathrm{NaBH}_{4}+\mathrm{PtCl}_{2}$

Hydrazine $\left(\mathrm{NH}_{2} \mathrm{NH}_{2}+\mathrm{O}_{2}\right)$

5. Birch Reduction: Birch reduction (alkali metals, such as Na and K in liquid $\mathrm{NH}_{3}+\mathrm{EtOH}$ or MeOH ), Mechanism of only terminal $(C=C)$ bond to $(C-C)$ bond:

$$
\mathrm{NH}_{2}-\mathrm{H}+\mathrm{HOO} \mathrm{Et} \longrightarrow\left(\mathrm{H}-\dot{\mathrm{N}} \mathrm{H}-\mathrm{Et}+\mathrm{H}_{2} \mathrm{O}\right.
$$

Source of H atom is from liquid $\mathrm{NH}_{3}$. If $\mathrm{ND}_{3}$ is used, D atom will be added.


Heat of hydrogenation: The energy released when one mole of alkenes is hydrogenated with $\mathrm{H}_{2}$ is called the heat of hydrogenation.

More negative the heat of hydrogenation of an alkene, less stable is the alkene. So, the stability order is as follows:
Trans-But-2-ene > cis-But-2-ene > But-1-ene




$$
\Delta \mathrm{H}^{\circ}=-120 \mathrm{~kJ} \mathrm{~mol}^{-1}
$$

cis-But-2-ene



$$
\Delta \mathrm{H}^{\circ}=-115 \mathrm{~kJ} \mathrm{~mol}^{-1}
$$

Heat of Combustion: When the hydrogenation of the alkene does not give the same alkane, the heat of combustion is used to measure their relative stabilities. More negative the value of $\Delta \mathrm{H}_{C}^{\circ}$, less stable is the compound or vice versa.
I.


But-1-ene
II.

III.

IV.


2-Methylprop-1-ene
Although each isomer mentioned above ( $\mathrm{I}-\mathrm{IV}$ ) consumes 6 moles of $\mathrm{O}_{2}$ and produces 4 moles of $\mathrm{CO}_{2}$ and 4 moles of $\mathrm{H}_{2} \mathrm{O}$, the comparison of $\Delta \mathrm{H}_{\mathrm{C}}^{\circ}$ shows that (IV) is most stable among the four isomers because it evolves the least heat. Thus, the order of stability is: (IV) $>$ (III) $>$ (II) $>$ (I).

Illustration 13: Give the products with their stereoisomers, if any.
(JEE ADVANCED)
(a) $\quad \mathrm{Me}-\equiv — \mathrm{Me}$

(b)

cis-2,3-Dibromo-but-2-ene
(c)


Sol: Identify the reducing agent used for the different reaction. Depending upon the type of reducing agent used and addition of hydrogen predict the formation of stereoisomer.
(a)

$$
\mathrm{Me} \text { — } \equiv — \mathrm{Me}
$$

(A)

| $\mathrm{H}_{2}+$ Poisoned Pd <br> Syn-add. | $\mathrm{Na}+$ liq. $\mathrm{NH}_{3}$ <br> + EtOH <br> Anti-add. | $\mathrm{Na}+$ liq. $\mathrm{ND}_{3}$ <br> + EtOH <br> Anti-add. <br> $\mathrm{ND}_{3}$ is a source <br> of D | $\mathrm{Na}+$ liq. $\mathrm{NH}_{3}$ <br> $+E t O D$ <br> Anti-add. <br> EtOD is not source |
| :---: | :---: | :---: | :---: |
|  |  |  | of $\mathrm{D} . \mathrm{NH}_{3}$ is source <br> of H |



Z or cis-But-2-ene
(I)


E or trans-But-2-ene


E or trans-2,3-
Dideuterio-but-2-ene


E or trans-But-2-ene
(IV)
(b) Note: Metal $+\mathrm{H}_{2}$ does not reduce $(\mathrm{R}-\mathrm{X})$ to $(\mathrm{R}-\mathrm{H})$ but reduced $(\mathrm{Ar}-\mathrm{X})$ to $(\mathrm{Ar}-\mathrm{H})$.

Reactant is cis + mechanism of addition is syn $\rightarrow$ product is meso.

(c) Reactant is trans + mechanism of addition is syn $\rightarrow$ product is dl or $( \pm)$ or racemic.


### 7.2 Electrophilic Addition Reactions

In electrophilic addition reactions of alkenes, a $\pi$ bond is broken and two new $\sigma$ bonds are formed. Formation of an electron deficient electrophile is the driving force for this reaction. This electrophile form a covalent bond with $\pi$ electrons of alkene and the positive charge is transferred to the remaining doubly bonded carbon.
In second step of E.A.R. positively charged intermediate combine with electron rich anion to form the second covalent bond. That is one weak $\pi$ bond is broken and two strong $\sigma$ bonds are formed.


Let us illustrate the mechanism of electrophilic addition reactions by taking the example of addition of $\mathrm{Br}_{2}$ to ethylene. The reaction occurs by a two-step ionic mechanism as discussed below:

Step-1: Bromine molecule is nonpolar but when it comes close to an ethylene molecule, the $\pi$-electrons of the double bond begin to repel the electron pair holding the two bromine atoms together in the bromine molecule. As a result, the bromine molecule gets polarized. The positive end of this bromine dipole behaves as an electrophile and is attracted by the $\pi$-electrons of the ethylene molecule to form a $\pi$-complex* which subsequently gives the carbocation and the bromide ion. This step is slow and hence is the rate determining step of the reaction.

This step can simply be represented as


Step-2: The carbocation formed in step-1 being a reactive chemical species immediately undergoes a nucleophilic attack by the bromide ion present in the solution forming the addition product. This step is fast and hence does not affect the rate of the reaction.


### 7.3 Addition of $\mathbf{X}_{2}$

Addition of halogens: Halogens such as chlorine and bromine readily add to alkenes to form 1, 2-dihaloalkanes. For example,



During the addition of bromine to alkenes, bromine loses its orange red colour, since the dibromide formed is colourless. This reaction is, therefore, used as a test for unsaturation in organic compounds. Reactivity order: $\mathrm{F}_{2}>\mathrm{Cl}_{2}>\mathrm{Br}_{2}>\mathrm{I}_{2}$ (because the $\mathrm{E}_{\text {activation }}$ order is: $\mathrm{F}_{2}<\mathrm{Cl}_{2}<\mathrm{Br}_{2}<\mathrm{I}_{2}$ ).

Reactant cis (with two same groups) + mechanism of addition of $\mathrm{Br}_{2}$ is anti (with two same groups) $\Rightarrow$ product is dl or $( \pm)$ of racemic. Reactant is trans + mechanism of addition of $\mathrm{Br}_{2}$ is anti $\Rightarrow$ product is meso. Similarly, maleic acid (cis) $+\mathrm{Br}_{2}$ (anti-addition) $\rightarrow$ dl product.


Fumaric acid (trans) $+\mathrm{Br}_{2}$ (anti-addition) $\rightarrow$ Meso product


### 7.4 Addition of HX

Addition of halogen halides. Mono-haloalkanes or alkyl halides are formed when alkenes react with halogen halides ( $\mathrm{HCl}, \mathrm{HBr}, \mathrm{HI}$ ),

e.g.


$$
\begin{array}{lc}
\text { Ethene } & \text { Bromoethane } \\
\text { (Ethylene) } & \text { (Ethyl bromide) }
\end{array}
$$

The order of reactivity of halogen halides in this reaction is: $\mathrm{HI}>\mathrm{HBr}>\mathrm{HCl}$. This order of reactivity can be explained on the basis of bond dissociation energies of the halogen halides; lower the bond dissociation energy, more reactive is the halogen halide ;
$\mathrm{HI}\left(300 \mathrm{~kJ} \mathrm{~mol}^{-1}\right)>\mathrm{HBr}\left(360 \mathrm{~kJ} \mathrm{~mol}^{-1}\right)>\mathrm{HCl}\left(430 \mathrm{~kJ} \mathrm{~mol}^{-1}\right)$
The actual product formed, however, depends upon whether the alkene is symmetrical or unsymmetrical as discussed below:
(i) Addition to symmetrical alkenes. When the alkene is symmetrical only one product is theoretically possible. For example,


Mechanism: Like the addition of halogen, addition of halogen halides to alkenes is also an electrophilic addition reaction and occurs by the following two steps:

Step 1.


Step 2.

(ii) Addition to unsymmetrical alkenes: When the alkene is unsymmetrical, two products are theoretically possible. For example, the addition of HBr to propene in the dark and in the absence of peroxides can, in principle, give two products. But experimentally, it has been found that under these conditions, the major product is 2-bromopropane and the minor product is 1-bromopropane.


Markovnikov's rule: The rule states that: "The addition of unsymmetrical reagents such as $\mathrm{HX}, \mathrm{H}_{2} \mathrm{O}, \mathrm{HOX}$, etc. to unsymmetrical alkenes occurs in such a way that the negative part of the addendum (i.e., adding molecule) goes to that carbon atom of the double bond which carries lesser number of hydrogen atoms."

For example,


Theoretical explanation of Markovnikov's rule: The addition of hydrogen halides to alkenes is an electrophilic addition reaction. Thus, during the addition of HBr to propene, the first step involves the addition of a proton. This addition, in principle, can occur in two ways. If the proton adds on the terminal carbon atom of the double bond, a $2^{\circ}$ carbocation (I) is formed and if addition occurs on the middle carbon atom. a $1^{\circ}$ carbocation (II) is produced.

Since, a $2^{\circ}$ carbocation (I) is more stable than $1^{\circ}$ carbocation (II), therefore, carbocation (I) is predominantly formed. This carbocation then rapidly undergoes nucleophilic attack by the $\mathrm{Br}^{-}$ion forming 2-bromopropane as the major product. Thus, Markovnikov's addition occurs through the more stable carbocation intermediate.


Peroxide effect: It should be noted that Markovnikov's rule is not always followed. In the presence of peroxides such as benzoyl peroxide ( $\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{CO}-\mathrm{O}-\mathrm{O}-\mathrm{COC}_{6} \mathrm{H}_{5}$ ), the addition of HBr (but not of HCl or HI ) to unsymmetrical alkenes takes place contrary to Markovnikov's rule. This is known as Peroxide effect or Kharasch effect. Thus,


Mechanism: The addition of HBr to alkenes in presence of peroxides occurs by a free radical mechanism. It consists of the following three steps.
(a) Initiation.
(i)

(ii)

(iii)

(b) Propagation: It consists of two steps.

During the first step, ${\mathrm{a} \mathrm{Br}^{-} \text {adds to the double bond in such a way so as to give the more stable free radical. In the }}^{\text {a }}$, second step, the free radical thus produced abstracts a H from HBr to complete the addition.
(i)

(ii)

(c) Termination:
(i) $2 \mathrm{Br} \longrightarrow \mathrm{Br}_{2}$
(ii)

(iii)


1,4-Dibromo-2,3-dimethylbutane
HCI, HF and HI do not obey Peroxide effect. Only HBr does WHY?

| $\mathbf{X}$ in HX | $\Delta \mathrm{H}$ (in kJ/mole) |  |
| :--- | :--- | :--- |
|  | (i) $\mathrm{X}+\mathrm{CH}_{2}=\mathrm{CHCH}_{3} \longrightarrow$ <br> $\mathrm{XCH}_{2}-\mathrm{CH}-\mathrm{CH}_{3}$ | (ii) $\mathrm{XCH}_{2}-\mathrm{CHCH}_{3}+\mathrm{HX} \longrightarrow \mathrm{XCH}_{2}-\mathrm{CH}_{2} \mathrm{CH}_{3}+\mathrm{X}$ |
| F | -209 | +159 |
| Cl | -101 | +27 |
| Br | -42 | -37 |
| I | +12 | -104 |

In HBr , both the steps being exothermic, it obeys the peroxide effect.
From the above data, it is clear that only with HBr , both the steps are exothermic and hence the peroxide effect is observed. With HCl or HF , the peroxide effect is not observed because the second step involving the reaction of carbon radical with HCl or HF is endothermic. Further, the peroxide effect is also not observed with HI because the first step involving the addition of iodine radical to alkenes is endothermic.

Illustration 14: Complete following reactions:
(JEE MAIN)
(a)

(d)
$\mathrm{Br}+\mathrm{HBr} \longrightarrow(\mathrm{D}$
(b)

(e)
$=\mathbb{B}$ $\xrightarrow{1 \mathrm{~mol} \text { of } \mathrm{HBr}}(\mathrm{E})$
(c)


Sol:
(a)

[+I effect of two (me) groups]

2-Methyl but-1-ene
or
Isobutylene

So $\mathrm{Cl}^{\ominus}$ and $\mathrm{I}^{\oplus}$
(Markovnikov's
addition) 2-methyl propane
(A)

2-Chloro-3-iodo-
(b)



(B) 1-Bromo-3,3,3-tri-fluoroprpane
(3,3,3-Trifluoroprop-1-ene)

F is highly electronegative in nature. Due to the -I effect of three F atoms, addition is anti-Markovnikov's (negative part of the reagent is adding to the C atom containing more number of H atoms).

The resonance effect ( $+R$ effect) from the lone pair $e^{-} s$ of $F$ atoms is not operative because lone pair $e^{-} s$ and double bond are not in conjugation.

So the $-I$ effect of $F$ atoms is operative and, hence, the electromeric effect of double bond takes place from $C_{1}$ to $C_{2}$.
(c)


Here, + R effect is more operative than -I effect of $O$ atom, since the lone pair on oxygen and double bond are in conjugation. Moreover, R > I, so Markovnikov's addition takes place.
(d)

(D) 1,1-Dibromethane
(Same explanation as in part (c) above)
(e)



Since, alkyl carbonium ion is more stable than vinyl carbonium ion, alkene reacts at a faster rate than alkynes towards FA.

Illustration 15: Complete the following reaction:
(JEE ADVANCED)
(a)

(A)
(b)

(G)

Sol: It is an example of free radical mechanism.
(a)


(b) $\mathrm{RO}+\mathrm{H}-\mathrm{CCl}_{3} \longrightarrow \mathrm{RO}+\dot{\mathrm{C}} \mathrm{Cl}_{3}$

(G)

Methtylene cyclonpetane

(H)

2,2,2-Trichloroethyl cylonepntane

### 7.5 Addition of HOX

Addition of the elements of hypohalous acids (HOX where $\mathrm{X}=\mathrm{Cl}, \mathrm{Br}$ or I) - Halohydrin formation. Chlorine and bromine in the presence of water readily add to alkenes to form the corresponding halohydrins.


Halohydrin

## Mechanism:



For example,


The overall reaction involves the addition of the elements of hypohalous acid $\left(\stackrel{\delta-}{\mathrm{HO}_{\mathrm{O}}-\mathrm{X}} \mathrm{X}\right)$ in accordance with Markovnikov's rule.


The order of reactivity of different hypohalous acids, i.e., $\mathrm{X}_{2} / \mathrm{H}_{2} \mathrm{O}(\mathrm{HOX})$ is:



### 7.6 Addition of $\mathrm{H}_{2} \mathbf{O}$ (Indirect)

Addition of sulphuric acid - Indirect hydration of alkenes. Cold, conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$ adds to alkenes to form alkyl hydrogen sulphates. In the case of unsymmetrical alkenes, addition occurs in accordance with Markovnikov's rule.



Isopropyl hydrogen sulphate
Importance. Alkyl hydrogen sulphates on boiling with water, undergoes hydrolysis to produce alcohols.
For example,


Thus, alkenes can be converted into alcohols as follows:


Ethene

This overall two-step conversion of an alkene first into alkyl hydrogen sulphate followed by hydrolysis with boiling water to form alcohols is called the indirect hydration of alcohols.

### 7.7 Addition of $\mathrm{H}_{2} \mathbf{O}$ (Direct)

Addition of water - Direct hydration of alkenes. Hydration means the addition of water. Ordinarily, water does not add directly to most of alkenes. However, some reactive alkenes do add water in the presence of mineral acids to form alcohols. The addition occurs in accordance with Markovnikov's rule.

For example,


### 7.8 Allylic Substitution

Alkenes mainly show two types of reactions:

1. Addition reaction (by ionic attack) and 2. Substitution reaction (by free-radical attack).


Substitution by free radical


Addition by ionic

Allylic halogenations can be carried out by the following reagents:

1. $\mathrm{Cl}_{2}$ at high temperature $(773 \mathrm{~K})$ or $\mathrm{Br}_{2}$ at 400 K .
2. NBS ( N -bromosuccinimide) $(\underset{O}{(2)} \mathrm{NBr})+h$
3. With NBS + DMSO (dimethyl sulphoxide)

4. $\mathrm{Cl}_{2} \mathrm{SO}_{2}$ (sulfuryl chloride) $\Rightarrow$ allylic chlorination
5. t-Butyl hypochlorite


The relative rate of abstraction of hydrogen:
$3^{\circ}$ allylic $>2^{\circ}$ allylic $>1^{\circ}$ allylic $>3^{\circ}>2^{\circ}>1^{\circ}>$ methyl $>$ vinylic

Illustration 16: Complete the following reaction:
(JEE MAIN)


Sol:


Cyclopent-1-ene 1-Bromocyclopent-2-ene

Illustration 17: Give the products of the reactions given below
(JEE ADVANCED)
(a)

(c)

Sol:
(a)

(b)

(d)

(b)

(c)



(H)

(F+G)
(cis and trans)

### 7.9 Diels Alder Reaction

A conjugate diene (with S-cis conformation) is treated with alkene or alkyne called a dienophile. The Diels-Alder reaction is insensitive to the presence or absence of solvents and catalysts. For example,


Mechanism: It takes place by concerted mechanism called pericyclic reaction. In a concerted mechanism, breaking and formation of bonds takes place simultaneously.

The reaction proceeds faster when the dienophiles have $\mathrm{e}^{-}$withdrawing groups, such as $(-\mathrm{COOH}),(-\mathrm{COOR}),(-\mathrm{CHO}),\left(-\mathrm{NO}_{2}\right)$, and (-CN), and dienes have $\mathrm{e}^{-}$donating groups such as (Me), (Et), etc. So, 1, 3-butadiene is less reactive than its mono-, di-, and trimethyl derivatives.

These reactions are stereospecific, i.e., maleic acid give cis and fumaric acid gives trans isomers, both by cisaddition.

Mechanisms: cis-additions



Illustration 18: Complete the following reactions:
(a)

(b)

$(\mathrm{D}) \xrightarrow{\Delta}$
(c)
 $+$

 (H) + (I)
(d)
 $+$


Sol: These are examples of Diels alder reaction. Diene and dienophile combine to form new ring. The mechanism is concerted type (Bond breaking and bond making takes place simultaneously)
(a)

(b)




### 7.10 Dimerisation

## Mechanism:


$(B)$ is sterically hindered because the bulky t-butyl group is cis to Me group

Alkene in the presence of HF or conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$ or $\mathrm{H}_{3} \mathrm{PO}_{4}$ gives two alkenes $\left(\mathrm{C}_{8} \mathrm{H}_{16}\right)$.

$\mathrm{HCl}, \mathrm{HBr}$, and HI cannot be used as acid catalysts. An acid catalyst must have a weak nucleophilic conjugate base to avoid addition of HX to the $(\mathrm{C}=\mathrm{C})$ bond. $\mathrm{Cl}^{\ominus}$, $\mathrm{Br}^{\ominus}$ and $\mathrm{I}^{\ominus}$, the conjugate bases of $\mathrm{HCl}, \mathrm{HBr}$, and HI , respectively; are good nucleophiles that bind to alkyl carbocation; whereas $\mathrm{F}^{\ominus}$, the conujugate base of HF , is a weak nucleophile and does not bind to alkyl carbocation.
$\mathrm{BF}_{3}$ with a little amount of $\mathrm{H}_{2} \mathrm{O}$ can also be used to initiate the dimerisation or polymerisation of an alkene only at 73K. It is due to the information of strong Bronsted acid $\left(\mathrm{BF}_{3}+\mathrm{H}_{2} \mathrm{O} \rightleftharpoons \mathrm{HBF}_{3} \cdot \mathrm{OH}\right)$, which provides $\mathrm{H}^{\oplus}$ to alkene to form the $R^{\ominus}$.

Illustration 19: Complete the following reaction:


## Sol:




### 7.11 Alkylation

Alkylation is the addition of alkene to alkane in the presence of $\mathrm{H}_{2} \mathrm{SO}_{4}$ or HF at 273 K .

isobutylene)

Mechanism: $\underset{\mathrm{Me}}{\mathrm{Me}}=\mathrm{CH}_{2} \xrightarrow[\text { Step } 1]{\mathrm{H}^{\oplus}}$




The propagation step is the chain reaction that continues when $\mathrm{Me}_{3} \mathrm{C}^{+}$adds to a molecule of isobutene to form the same $3^{\circ} \mathrm{R}^{+}$, which accepts as $\mathrm{H}^{-}$(hydride ion) from another alkane molecule.

### 7.12 Carbene Insertion to Alkenes

Alkenes react with carbenes in the presence of light to give a cyclic compound.


The carbene is $\mathrm{e}^{-}$-deficient and adds to the $\mathrm{e}^{-}$rich -bond to give a cyclopropane ring.
Stereochemistry of Carbene: The addition of a singlet carbene ( $1 \cdot \mathrm{CH}_{2} \cdot l$ ) is both stereospecific and stereoselective and syn, while the addition of a triplet carbene $\left(1 \cdot \mathrm{CH}_{2} \cdot l\right)$ is non-stereospecific and non-stereoselectively.
Diazomethane generates singlet carbene but in the presence of inert $\mathrm{N}_{2}$, the singlet carbene collides with $\mathrm{N}_{2}$ molecules and loses energy to give the carbene having the lower energy triplet.


Illustration 20: Give the product of the following:
(a) Cyclohexane $+\mathrm{CH}_{2} \mathrm{~N}_{2}+\mathrm{hv}$
(b) Trans-2-Butene $+\mathrm{CHCl}_{3}+\mathrm{KOH}$
(c) Cis-2-Butene $+\mathrm{HCBr}_{2} \mathrm{Cl}+\mathrm{KOH}$
(d) Trans-2-Butene $+\mathrm{HCBr}_{2} \mathrm{Cl}+\mathrm{KOH}$
(e) 3-Methylcyclopentene $+\mathrm{CH}_{2} \mathrm{I}_{2}+\mathrm{Zn} / \mathrm{Cu}$ (couple) (Simmons-Smith reaction)
(f)


## Sol:

(a)

Bicyclo [4.1.0] heptane

(b)

trans-1,2-Dimethyl 3,3-dichloro cyclopropane
(c)

trans-2,3-Dimethyl-1chlorobromo cyclopropane (Each halogen is cis to one Me and trans to other)
(e) Two diastereomers are obtained:

(f) By this method 7- to 10-memberd cyclic ring can be prepared.


### 7.13 Isomerisation and the Shifting of Double Bond

Shifting of double bond (centre of the chain) or migration of methyl group takes place when heated at 770-970K or when heated with a catalyst $\mathrm{Al}_{2}\left(\mathrm{SO}_{4}\right)_{3}$ at $470-570 \mathrm{~K}$.


When either a cis or trans alkene is placed in a strong acid solution, each of them is converted to a mixture of $76 \%$ of trans and $24 \%$ of the cis-isomer. The ratio of isomers is thermodynamically or equilibrium controlled.


### 7.14 Oxidation Reactions

1. Complete oxidation with oxygen or air - Combustion: All combustion reactions are highly exothermic in nature. For example, $\mathrm{CH}_{2}=\mathrm{CH}_{2}+3 \mathrm{O}_{2} \longrightarrow 2 \mathrm{CO}_{2}+2 \mathrm{H}_{2} \mathrm{O}:+\Delta_{\mathrm{C}} \mathrm{H}=-1411 \mathrm{~kJ} \mathrm{~mol}^{-1} \mathrm{k}$.
2. Controlled oxidation with oxidising agent gives different products:
(i) Oxidation with oxygen: Alkenes react with $\mathrm{O}_{2}$ in the presence of silver as a catalyst to form epoxy alkanes or epoxides. For example,



1,2-Epoxypropane

### 7.14.1 Hydroxylation

Alkenes are oxidised to 1,2-diols with any of the following reagents:
(a) Cold alkaline or aqueous $\mathrm{KMnO}_{4}$ solution: It is also called Baeyer's reagent. It is a test for unsaturation for both alkenes and alkynes. The pink colour of $\mathrm{KMnO}_{4}$ is discharged and the black-brown precipitate of $\mathrm{MnO}_{2}$ is obtained, and alkenes are converted to diols.
(b) $\mathrm{OsO}_{4}$ (osmium tetraoxide) in pyridine or ether solution followed by the reaction with $\mathrm{Na}_{2} \mathrm{SO}_{3} / \mathrm{H}_{2} \mathrm{O}$ or $\mathrm{NaHSO}_{3} /$ $\mathrm{H}_{2} \mathrm{O}$ converts alkene to diols and $\mathrm{OsO}_{4}$ to Os metal.
(c) With catalytic amount of $\mathrm{OsO}_{4}$ in $\mathrm{H}_{2} \mathrm{O}_{2}$ : With the reagents (a), (b) and (c), the addition of two (-OH) groups takes place by syn-addition (or cis-addition) and is called syn-hydroxylation.

Anti Hydroxylation with the following reagents :
(d) Peroxy acid followed by the hydrolysis with $\mathrm{H}_{2} \mathrm{O}$ :

For example,

i. $\mathrm{H}-\mathrm{C}$


Peroxybenzoic acid
iii. MCPBA (m-chloroperbenzoic acid)


## Mechanism of syn hydroxylation:

i.


(Meso compound) (O.I.A)
Butan-2,3-diol
ii.


## Mechanism of Anti-hydroxylation

I.
 cis-But-2-ene

II.


For syn hydroxylation,

|  | Reactant | Reagent mechanism | Product |
| :--- | :--- | :--- | :--- |
| 1. | Cis (with two same groups) | Syn (with two same $(-\mathrm{OH})$ groups) | Meso |
| 2. | Trans (with two same groups) | Syn (with two same $(-\mathrm{OH})$ groups) | $( \pm)$ or racemate |
| 3. | Cis or trans (with two different groups) | Syn (with two same $(-\mathrm{OH})$ groups) | $( \pm)$ or racemate |

## For anti-Hydroxylation,

|  | Reactant | Reagent mechanism | Product |
| :--- | :--- | :--- | :--- |
| 1. | Cis (with two same groups) | Anti (with two same $(-\mathrm{OH})$ groups) | $( \pm)$ or racemate |
| 2. | Trans (with two same groups) | Anti (with two same $(-\mathrm{OH})$ groups) | Meso |
| 3. | Cis or trans (with two different groups) | Anti (with two same $(-\mathrm{OH})$ groups) | $( \pm)$ or racemic |

Illustration 21: Convert


Sol: First the compound undergoes dehydrohalogenation reaction to form an unsaturation. This on treatment with aq. $\mathrm{KMnO}_{4}$ gives diol.


### 7.14.2 Cleavage

## A. Oxidative cleavage

Alkenes are oxidatively cleaved by any of the following reagents:

1. Hot alkaline $\mathrm{KMnO}_{4}$ solution followed by acidification
2. Hot acidic $\mathrm{KMnO}_{4}$ solution.
3. Hot acidic $\mathrm{K}_{2} \mathrm{Cr}_{2} \mathrm{O}_{7}$ solution.


With terminal alkenes, one of the products is always methanoic acid (formic acid) with on further oxidation gives $\mathrm{CO}_{2}+\mathrm{H}_{2} \mathrm{O}$. For example,


With non-terminal alkenes, carboxylic or ketones or both of these are obtained depending upon the nature of the alkene. For example.


Rules of oxidation:

| $\mathrm{CH}_{2} \neq \Rightarrow$ | $\mathrm{CO}_{2}$ |
| :--- | :--- |
| $\mathrm{R}-\mathrm{CH} \neq \Rightarrow$ | $\mathrm{R}-\mathrm{COOH}$ |
|  | R |

The oxidative cleavage of alkenes is used to establish the position or location of the double bond and the structure of the alkene chain or ring.



Propan-2-one

Illustration 22: Compound $(A)\left(\mathrm{C}_{6} \mathrm{H}_{8}\right) \xrightarrow{[\mathrm{O}]}$ Malonic acid only. Identify compound (A).
Sol: First, calculate the degree of unsaturation.
D.U. $=\frac{\left(2 n_{C}+2\right)-n_{H}}{2}=\frac{(2 \times 6+2)-8}{2}=3^{\circ}$


It is a 3C-atom dibasic acid, but compound (A) has six C atoms. So, 2 mol are joined with each other.




Cyclohexa-1,4-diene
$2(\mathrm{C}=\mathrm{C})$ bonds
B. Ozonolysis (Oxidation with ozone): When ozone is passed through a solution of an alkene in some inert solvent such as $\mathrm{CH}_{2} \mathrm{Cl}_{2}, \mathrm{CHCl}_{3}$ or $\mathrm{CCl}_{4}$ at a low temperature (196-200K), it oxidises alkenes to ozonides. Ozonides are unstable and explosive compounds. Therefore, they are not usually isolated but are reduced, in situ, with Zn dust and water or $\mathrm{H}_{2} / \mathrm{Pd}$ to give aldehydes or ketones or a mixture of these (reductive cleavage) depending upon the structure of the alkene.



However, if the ozonide are decomposed only with water, the $\mathrm{H}_{2} \mathrm{O}_{2}$ produced during the reaction oxidises the initially formed aldehydes to the corresponding acids (oxidative cleavage).
This two-step conversion of an alkene into an ozonide followed by its reductive cleavage to yield carbonyl compounds is called ozonolysis.
Importance: Ozonolysis is a versatile method for locating the position of a double bond in an unknown alkene gives the same combination of aldehydes and or/ketones. Therefore, this method that has been extensively used in the past for structure elucidation of alkenes.


$\xrightarrow[\text { (ii) } \mathrm{Zn} \text {, } \mathrm{O}_{2} \text { ( } \mathrm{O}_{3} \mathrm{CH}_{2}, 196-200 \mathrm{~K}]{\text { (in }}$
(ii) $\mathrm{Zn} / \mathrm{H}_{2} \mathrm{O}$
$\mathrm{CH}_{3} \mathrm{CHO}+\mathrm{CH}_{3} \mathrm{CHO}$
Ethanal (Two molecules)
But-1-ene


2-Methylpropene




Reagents used for reducive and

(Reductive ozonolysis)

(a) Zn -acid
(b) $\mathrm{H}_{2}+$ Catalyst ( $\mathrm{Ni}, \mathrm{Pt}, \mathrm{Pd}$ )
(c) $\mathrm{Me}_{2} \mathrm{~S}$ (Dimethyl sulphide)
(d) $\mathrm{Ph}_{3} \mathrm{P}$ (Triphenyl phosphine) $\Downarrow$ It gives aldehyde and ketone

Oxidation of ozonide
(Oxidative ozonolysis)
(f) $\mathrm{H}_{2} \mathrm{O}$
(g) $\mathrm{Ag}_{2} \mathrm{O}$
(h) $\mathrm{H}_{2} \mathrm{O}_{2}$
(i) $\mathrm{KMnO}_{4} / \mathrm{NaIO}_{4}$
(j) $\mathrm{H}_{2} \mathrm{O}$
$\left(\mathrm{HCO}_{3} \mathrm{H}, \mathrm{PhCO}_{3} \mathrm{H}\right)$


Aldehyde give acids and ketones do not change

For example:


Illustration 23: Identify the compounds (A), (B), and (C).
(JEE MAIN)


Does not show
Stereoisomerism

$$
\mathrm{O}_{3} / \mathrm{Ph}_{3} \mathrm{P}
$$

Symmetrical diketone
(C) $\left(\mathrm{C}_{8} \mathrm{H}_{12} \mathrm{O}_{2}\right)$

Sol: First calculate degree of unsaturation for $\mathrm{A}, \mathrm{B}$ and C
D.U. in $(A)=\frac{\left(2 n_{C}+2\right)-n_{H}}{2}=\frac{(2 \times 8+2)-12}{2}=3^{\circ}$;
D.U. in $(B)=\frac{\left(2 n_{C}+2\right)-n_{H}}{2}=\frac{(2 \times 8+2)-14}{2}-2^{\circ}$
D.U. in $(C)=\frac{(2 \times 8+2)-12}{2}=3^{\circ}$

In compound (C), Degree of Unsaturation is $3^{\circ}$. Out of three, two are due to diketone.
So, one D.U. in compound $(C)$ is due to either one $(C=C)$ bond or one ring of eight $C$ atoms.
One $(C=C)$ bond is not possible because after ozonolysis of $(A)$, the double bond cannot be left behind. So $(C)$ is 8 C ring with two $(-\stackrel{( }{\mathrm{C}}-)$ groups at the symmetric position.

Proceed reverse,
Remove O atoms and join the double bond to get (A)


Bicyclo [3.3.0] oct-1-ene
Does not show Stereoisomers
Two rings = 2 D.U.
One double bond =1 D.U.
Total D.U. $=3$

| $\mathrm{O}_{3} / \mathrm{Ph}_{3} \mathrm{P}$ |
| :--- | :--- |
| Red. ozonolysis |


(C) Cyclooctane Symmetrical diketone
One Ring = 1 D.U.


Total D.U. $=3$

## 8. USES OF ALKENES

(i) Lower members of the family are used as fuel and illuminants.
(ii) Alkenes and substituted alkenes upon polymerisation form a number of useful polymers such as polythene, PVC, teflon, orlon, etc.
(iii) Ethene is employed for the preparation of ethyl alcohol and ethylene glycol (anti-freeze).
(iv) Ethylene is used for artificial ripening of green fruits.
(v) Ethylene is also used in oxygen ethylene flame for cutting and welding of metals.

## Solved Examples

## JEE Main/Boards

Example 1: Complete the following reactions:
(a)

(A)
(b)

(c)

(B)
(d)
(A)

(A)
(e)


Sol: (a)

(b) (Z) is formed by Saytzeff's elimination, but the product $(\mathrm{Y})$ is formed in major amount because product $(\mathrm{Y})$ is a more stable conjugate diene than $(Z)$, isolated diene


4-Bromo-5-methylhex-1-ene (X)

(c)


Less stable isolated diene
(A)

Cyclohexa-1,4-diene

More stable conjugated diene (B)

Cyclohexa-1,3-diene
(d)


(e)


Example 2: Give the major products (not stereoisomers) of the following:
(a

(b)


1-Ethylcyclopenta-1,3-diene


Sol:
(a)

(A)

All 1,4-addition products


3,5-Dibromo-cyclopent-1-ene


3-Bromo-cyclopent-1-ene


Cyclopent-1-ene-3-thiol
(b)


All 1,4-addition products


Example 3: Write all the possible structures and give the structure of the products that are thermodynamically favoured.

Sol:


Hexa-1,3-5-triene




More stable Products (II and III, conjugate dienes) are thermodynamically favoured products than I and IV (isolated dienes)

Example 4: Distinguish between the given pairs.
(a)

(A)
and

(B)
(b)

(C) and

(D)

Sol: (a) Compounds (A) and (B) are distinguished chemically by quantitative catalytic hydrogenation. Since compound $(A)$ has two double bond whereas compound $(B)$ has only one double bond .Compound $(A)$ will require 2 moles of $\mathrm{H}_{2}$ for 1 mole of $(A)$ while $(B)$ will requires 1 mole of $\mathrm{H}_{2}$ per mole of (B).

(b) Allene (C) with different groups on each of the double bonds is optically active and can be resolved into its enantiomers.
Compound (D) is conjugate diene and does not show optical isomerism.

Example 5: Give the major and minor products.

(A)

3-Methylenecyclohex-1-ene
Sol: At high temperature $\left(45^{\circ} \mathrm{C}\right), 1,4$-addition products is more favourable.


Example 6: Dehydration of (A) with conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$ gives a compound that exists in two isomeric forms. Give the structures of both the isomers.

(A)

Sol:


Example 7:

(A)

Sol: Addition of H atom by Lindlar's catalyst is a syn-addition. It reduces $(\mathrm{C} \equiv \mathrm{C})$ bond to $(C=C)$ bond so the product $(B)$ is


Write all the possible structures of (A), (B) and (C).
Sol: Since the reductive and oxidative ozonolysis products are different, the alkene of the type


If $R=\left(\square-\mathrm{CH}_{2}-\right.$, the possible structures of $(A)$ can be:
$R=\left(D-\mathrm{CH}_{2} \perp\right.$, the possible structures of $(\mathrm{A})$ can be:




(B)
(B)

Optically inactive
Optically inactive

$\left(D_{1}\right)$
(D) is obtained when $\left(C_{1}-C_{3}\right)$ bond breaks
$\left(D_{1}\right)$ is obtained when $\left(C_{2}-C_{3}\right)$ bond
Example 9: Give the number of stereoisomers of (A) in the following reactions.
(a) $(A)\left(\mathrm{C}_{2} \mathrm{H}_{12}\right) \xrightarrow{\mathrm{O}_{3} / \text { oxidation }}$ Acetone + Oxalic acid + Acetic acid
(B)
(C)
(D)
(b) $\quad(\mathrm{A})\left(\mathrm{C}_{8} \mathrm{H}_{14}\right) \xrightarrow{\mathrm{O}_{3} / \text { oxidation }}$ Butane-2-one + Oxalic acid + Acetic acid (B)
(C)
(D)

## Sol:

(a)


(A) $\left(\mathrm{C}_{7} \mathrm{H}_{12}\right)$

Here Compound (A) shows G.I around $\mathrm{C}_{4}-\mathrm{C}_{5}$ double bond (since the two groups around C-4 and C-5 are different) but do not show G.I around $\mathrm{C}_{2}-\mathrm{C}_{3}$ double bond because the two groups around C-2 are the same.

Number of stereoisomers for (A) — 2


4Z-2-Methylhexa-2,4-diene


4E-2-Methylhexa-2,4-diene
(b) Proceed in the same manner as in (a)


(5-Methylhepta-2,4-diene)
(A) shows geometrical isomers around both $\left(C_{2}-C_{3}\right)$ and $\left(C_{4}-C_{5}\right)$ double bonds (two of the groups around these double bonds are different)
Moreover, terminal groups $\left(-\mathrm{CH}_{3}\right)$ and $\left(-\mathrm{C}_{2} \mathrm{H}_{5}\right)$, around $\mathrm{C}_{2}$ and around $\mathrm{C}_{5^{\prime}}$ respectively, are different.
Number of G.I. when the terminal groups are different $=2^{n}$, where $n$ is the number of double bonds (two double bonds).
G.I. $=2^{2}=4$

Number of stereoisomers of $(A)=4$

(I)
$(2 Z, 4 Z)$

(II)
( $2 \mathrm{E}, 4 \mathrm{Z}$ )

(III)
( $2 \mathrm{Z}, 4 \mathrm{E}$ )

(IV)
( $2 \mathrm{E}, 4 \mathrm{E}$ )

Example 10: Explain the formation of the products giving the structures of the intermediates.
(a)



In this reaction presence of double bond influences the reaction pathway and unusual product may also be formed. Due to the presence of $e^{-}$-rich double bond, the intermediate (I) carbocation may also involve intra-molecular rearrangement to form a three membered cyclic ring as shown below:


Other products can be obtained due to carbocation (I) rearrangement.


Example 11: Complete the following reaction:


Sol: It is an example of hydroboration-oxidation reaction, it is used in the formation of alcohol from an alkene. It follows anti-Markovnikov's rule, and the addition of H and OH is syn (cis)(addition of H and $\mathrm{BH}_{2}$ takes place from the same side)

(5-Chloropentane-1-ol)
(B)


## JEE Advanced/Boards

Example 1: Complete the following reaction with appropriate reagents.
(a)

(b)


Sol: (a) Compound (D) Contains a $(C=C)$ bond in the ring, Double bond can be obtained by the dehydration of alcohol.So, $(C)$ is an alcohol.Alcohol $(C)$ is obtained by the reduction of $(>C=0)$ using reducing agent like LAH.
$\therefore$ So $(B)$ contains a keto group $(>C=O),(>C=O)$ group can be obtained from $(A)$ by ozonolysis.
The different steps are as follows:

(b) Proceeding reverse: (Retrosynthetic approach)

Compound (D) contains one Me and one -OH group. It can be obtained by adding $\left(\mathrm{Cl}_{2}+\mathrm{H}_{2} \mathrm{O}\right)(\mathrm{HOCl})$ to (A).
Thus compound $(B)$ will have one OH and one Cl group; now one Me group can be added to (B) by Corey-House synthesis or Wurtz reaction of (B).


Example 2: Complete the following reactions:
(a)

(A)
(b)

(A)
(c)


Sol:
(a)

(b)


(c) It is an example of -elimination.


Example 3: Identify the products in the following reactions giving their stereoisomers (if any).
(a)

(b) (B)(from Problem A) $\xrightarrow[\text { reagent }]{\text { Baeyer's }}(D)+(E)$ (Colour of precipitate)
(c)

(d)


Sol:


2Z-2,3-Di-d-pent-2-ene
2,3-Di-d-pentane
Since Addition of $D_{2}$ by $\left(D_{2}+P d\right)$ is a syn addition Compound $(B)$ is a cis compound and the addition of $\mathrm{H}_{2}+\mathrm{Pt}$ is also syn. But (B) has two different (R) groups (Me and Et), so the product (C) will not be Meso but a racemic compound.
Compound (C) contains two chiral carbon, thus the number of optical isomers will be $2^{2}=4$ (two pairs of enantiomer).
These are:


Racemate



Racemate

Pairs of enantiomers: I and II; III and IV (I, III); (I, IV); (II, III),
Pairs of diastereomers: (II, IV)
(b)

cis-Compound with two different R group

(Brown-black ppt)

2,3-Dideutentropent-2,3-diol
2,3-Di-d-pent-2,3-diol
(E)

$$
+\mathrm{MnO}_{2}
$$

[Two chiral centres; four stereoisomers as in (a)]
(B)
(c) cis-Compound with two different

R group
(d)

or
2E-2,3-Di-d-pent-2-ene


Example 4: Identify (A), (B) and (C) in the following reaction.


(4-Oxohexan-1,6-dioic acid) or 4-Ketoadipic acid)

Sol: First of all we have to calculate degree of unsaturation:
D.U. in $(A)=\frac{\left(2 n_{C}+2\right)-n_{H}}{2}=\frac{(7 \times 2+2)-10}{2}=3^{\circ}$
D.U. in $(B)=\frac{(7 \times 2+2)-14}{2}=1^{\circ}$

Since (A) does not react with P-2 or Brown catalyst $\left(\mathrm{Ni}_{2} \mathrm{~B}\right)$ that reduces alkyne to alkene
Therefore, (A) does not contain ( $C$ ミ $C$ ) bond.
As we have calculated, (B) has 1.D.U., it must be a ring (because after hydrogenation of (A) no double bond will be left)

From the above data it appears that, $(A)$ has one ring and two $(C=C)$ bonds; hence, $(A)$ is a diene.
$\left.\begin{array}{l}\text { Total number of } C \text { atoms in }(A)=7 \\ \text { Total number of } C \text { atoms in }(C)=6\end{array}\right]$ one $C$ atoms is lost as $\mathrm{CO}_{2}$
Possible structure of (A) are as follows:
i.


ii. Other two possibilities of the combination are as follows:




(II)
(Numbering in accordance with naming)
1-(prop-2-enyl)cyclobut-1-ene or 1-Allyl cyclobutene
(b) $\left.\stackrel{6}{6}_{(\mathrm{COOH}}^{\mathrm{CO}}\right)$ Groups combines with $(\mathrm{C}=\mathrm{O})$ group and $\mathrm{CO}_{2}$ is in combination with $(\stackrel{1}{\mathrm{COOH}})$


(Numbering in accordance with problem)

(Numbering in accordance with naming)
(4-(Cyclopropenyl)but-1-ene)
Structure III is not possible, because on hydrogenation (A) gives (B) $\left(\mathrm{C}_{7} \mathrm{H}_{14}\right)$, i.e., it absorbs 2 mol of $\mathrm{H}_{2^{\prime}}$ whereas III will absorb 3 mol of $\mathrm{H}_{2}$.

Because of highly strained ring Structure III is unlikely.
Two possible structures are as :-


Example 5: Complete the following reactions:


Sol: It is an example of Diels Alder reaction. The mechanism is a concerted type (bond breaking and bond making takes place simultaneously)


Example 6: Identify $(\mathrm{A})$ to (H)

(A)


Sol:

(A)

| Reagent I |  | II | III | IV | V |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathrm{H}_{2}+\mathrm{Ni}$ | $\begin{aligned} & 100 \mathrm{~atm}, \Delta \\ & \text { or } \\ & \mathrm{H}_{2}+\mathrm{Pt}+\mathrm{Al}_{2} \mathrm{O}_{3} \\ & 35 \mathrm{~atm}, \Delta \end{aligned}$ | $\mathrm{H}_{2}+\mathrm{Ni}$ | $\mathrm{NH}_{2} \mathrm{NH}_{2}+\mathrm{O}_{2}$ |  |  | $\mathrm{Sia}_{2} \mathrm{BH}+$ $\mathrm{CH}_{3} \mathrm{COOH}$ |
|  | (B) | (C) | (D) $+(\mathrm{E})$ (gas) | $(\mathrm{F})+(\mathrm{G})$ |  |  |




Example 7: Write the product with its mechanism.

(A)

Sol:


## 2,2,3-Trimethyl bicyclo [4.4.0] dec-1-ene

Example 8: Which of the following show G.I.?
(a)

(b)

(c)

(d)

(e)

(f)

(g) $\mathrm{Ph}-\mathrm{N}=\mathrm{N}-\mathrm{Ph}$
(h)


Sol: In order to show geometrical isomerism the alkene must contain different substituents i.e. the substituents on both the carbon bearing double bond must have different substituents
(a) It shows G.I.




(i) Two atoms on the dotted line (......................) means that the groups are below the plane of the ring.
(ii) Two atoms on the bold lines (...........................) means that the groups are above the plane of the ring.
(iii) One atom on the dotted line (......................) means that one group is below and another is above the plane of the ring.
(b) It shows G.I.


(c) It does not show G.I.


Priority of $\mathrm{C}_{2} \mathrm{H}_{5}>\mathrm{CH}_{3}$ and $\mathrm{OH}>$ LP of es

(e) It does not show G.I


Two same groups
(f) It does not show G.I
(g) It shows G.I.


Two different groups

Two different groups

anti or E
(h) It shows G.I.


Cis has a plane of symmetry so O.I.A (Meso from)

(+) or D

(-) or L
Trans is optical active

Example 9: Give IUPAC name to each of the following using E or Z designations.
(A)

(B)

(C)


Sol:
(a)


Two higher priority groups on opposite side, 1 E ,
1-Bromo-2,3-dimethyl pent-2-ene
(b)


Priority of $(-\mathrm{Br})>\left(-\mathrm{CH}_{2} \mathrm{OH}\right)$ and $\left[\left(\mathrm{CH}_{3}\right)_{2} \mathrm{CH}-\right]>\left(-\mathrm{C}_{3} \mathrm{H}_{7}\right)$.
Two higher priory groups on the same side.
2Z,2-Bromo-3-(1-methyl ethyl) hex-2-en-1-ol
(c)


Priority of $\mathrm{HC} \equiv \mathrm{C}->\left(\mathrm{CH}_{3}\right)_{3} \mathrm{C}->\mathrm{CH}_{2}=\mathrm{CH}-\mathrm{B}>\left(\mathrm{CH}_{3}\right)_{2} \mathrm{CH}-$
Two higher priority groups on the same side.
3Z,3-(1-methyl ethyl)-4-(1-1-Dimethyl ethyl)-hexa-1,3-dine-5-yne

## ALKYNES

## 1. INTRODUCTION

### 1.1 Nomenclature

Alkynes are unsaturated hydrocarbons that contain $(\mathrm{C} \equiv \mathrm{C})$ bond. Their general formula is $\mathrm{C}_{n} \mathrm{H}_{2 n-2}$.

| Condensed | Stick | Bond line | IUPAC name | Common or trivial name |
| :---: | :---: | :---: | :---: | :---: |
| $\mathrm{C}_{2} \mathrm{H}_{2}$ | $\mathrm{H}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}$ | $\stackrel{2}{\overline{1}}$ | Ethyne | Acetylene |
| $\mathrm{C}_{3} \mathrm{H}_{4}$ | $\mathrm{H}_{3} \mathrm{C}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}$ | $3 \underline{-}^{2} \equiv$ | Propyne | Methylacetylene |
| $\mathrm{C}_{4} \mathrm{H}_{6}$ | $\begin{aligned} & \mathrm{CH}_{3}-\mathrm{CH}_{2}-\mathrm{C} \equiv \mathrm{CH} \\ & \mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3} \end{aligned}$ | $\begin{aligned} & 4_{3}^{2}=1 \\ & 4-3{ }^{2}=1 \end{aligned}$ | But-1-yne <br> But-2-yne | Ethylacetylene <br> Dimethy lacetylene |
| $\mathrm{C}_{5} \mathrm{H}_{8}$ | $\mathrm{CH}_{3}-\mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}$ |  | 3-Methyl-but-1-yne | Propylacety lene |
|  | or | $\rangle^{4}{ }^{2}=1$ | 3-Methyl-but-1-yne | Isopropyl-acetylene |
|  | $\stackrel{\text { or }}{\mathrm{CH}_{3}-\mathrm{CH}_{2}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3}}$ | $\underline{5}_{4}^{3}=2$ | Pent-2-yne | Ethyl methylacetylene |

### 1.2 Structure

Each carbon atom of ethyne is sp-hybridized. One sp-hybridized orbital of each carbon undergoes head on overlap with sp-hybrisized orbital of another carbon to form a sp-sp, C-C, $\sigma$-bond. The second sp-hybridized orbital of each carbon overlaps along the internuclear axis with 1 s -orbital of each of the two hydrogen atoms forming two $\mathrm{sp}-\mathrm{s}$, $\mathrm{C}-\mathrm{H}, \sigma$-bonds. Each carbon is now left with two unhybridized p -orbitals ( $2 \mathrm{p}_{\mathrm{x}}$ and $2 \mathrm{p}_{\mathrm{y}}$ ) which are perpendicular to each other as well as to the plane of the C-C sigma bond.

The two $2 p_{x}$ - orbitals, one on each carbon, are parallel to each other and hence overlap sideways to form a $\pi$-bond. Similar overlap between $2 p_{y}$ - orbitals, one on each carbon, results in the formation of a second $\pi$-bond as shown in figure.
(a)

(b)


Carbon-carbon triple bond consists of one strong $\sigma$-bond and two weak $\pi$-bonds. The total strength of $C \equiv C$ bond in ethyne is $823 \mathrm{~kJ} \mathrm{~mol}^{-1}$. It is stronger than the $\mathrm{C}=\mathrm{C}$ bond of ethane ( $599 \mathrm{~kJ} \mathrm{~mol}^{-1}$ ) and $\mathrm{C}-\mathrm{H}$ bond of ethane ( 348 kJ mole ${ }^{-1}$ ). Further, due to the smaller size of sp-orbitals (as compared to $\mathrm{sp}^{2}$ and $\mathrm{sp}^{3-}$ ) and sideways overlap of p -orbitals, the carbon-carbon bond length in ethyne is shorter $(120 \mathrm{pm})$ than those of $C=C(134 \mathrm{pm})$ and $\mathrm{C}-\mathrm{C}$ (154 pm).

### 1.3 Isomerism in Alkynes

(i) Position isomerism. The first two members, i.e., ethyne and propyne exist in one form only. However, butyne and higher alkynes exhibit position isomerism due to the different position of the triple bond on the carbon chain. For example,



But-2-yne
(ii) Chain isomerism. Alkynes having five or more carbon atoms show chain isomerism.

(iii) Functional isomerism: Alkynes are functional isomers of dienes



But-1-yne
Buta-1,3-diene
(iv) Ring chain isomerism: Alkynes show ring chain isomerism with cycloalkenes. For example,


Propyne

## 2. PREPARATION OF ALKYNES

### 2.1 From Calcium Carbide

Alkynes are prepared by the following general methods.

1. By the action of water on calcium carbide: Ethyne (acetylene) is prepared in the laboratory as well as on a commercial scale by the action of water on calcium carbide.

$$
\mathrm{CaC}_{2}+2 \mathrm{H}_{2} \mathrm{O} \rightarrow \mathrm{HC} \equiv \mathrm{CH}+\mathrm{Ca}(\mathrm{OH})_{2}
$$

Calcium carbide Ethyne (Acetylene)
Calcium carbide needed for the purpose is manufactured by heating limestone (calcium carbonate) with coke in an electric furnace at 2275 K .
$\mathrm{CaCO}_{3} \xrightarrow{2275 \mathrm{~K}} \mathrm{CaO}+\mathrm{CO}_{2}$
$\mathrm{CaO}+3 \mathrm{C} \xrightarrow{2275 \mathrm{~K}} \mathrm{CaC}_{2}+\mathrm{CO}$
2. Procedure: Lumps of calcium carbide are placed on a layer of sand in a conical flask fitted with a dropping funnel and a delivery tube. The air present in the flask is replaced by oil gas since acetylene forms an explosive mixture with air. Water is now dropped from the dropping funnel and the acetylene gas thus formed is collected over water.
3. Purification: Acetylene gas prepared by the above method contains impurities of hydrogen sulphide and phosphine due to the contamination of CaS and calcium phosphide in calcium carbide. Phosphine is removed by passing the gas through a suspension of bleaching powder. Pure acetylene is finally collected over water.

### 2.2 Double Dehydrohalogenation of Dihalides

Double dehydrohalogenation of a geminal dihalide


Geminal dihalide Sodium amide Alkyne Ammonia Sodium halide

## Double dehydrohalogenation of a vicinal dihalide



Vicinal dihalide Sodium amide Alkyne Ammonia Sodium halide
The most frequent application of these procedures are in the preparation of terminal alkynes. Since the terminal alkyne product is acidic enough to transfer a proton to the amide anion. One equivalent of base in addition to the two equivalents required for double dehydrohalogenation is needed. Adding water or acid after the reaction is complete converts the sodium salt to the corresponding alkyne.


1,1-Dichloro-3,3- Sodium salt of alkyne 3,3-Dimethyl-
Dimethylbutane product (not isolated) 1-butyne (56-60\%)


1,2-Dibromodecane $\quad$ Sodium salt of alkyne 1-Decyne (54\%) Product (not isolated)

Double dehydrohalogenation to form terminal alkynes may also be carried out by heating geminal and vicinal dihalides with potassium ter-butoxide in dimethyl sulfoxide. By heating with alcoholic solutio`n of KOH


## PLANCESS CONCEPTS

Misconception: The reaction, in fact, occurs in two steps and each step involves the loss of a molecule of HBr as shown below:


Ethylene dibromide Vinyl bromide
$\mathrm{CH}_{2}=\mathrm{CH}-\mathrm{Br}+\mathrm{KOH}$ (alc.) $\xrightarrow{\Delta} \mathrm{CH} \equiv \mathrm{CH}+\mathrm{KBr}+\mathrm{H}_{2} \mathrm{O}$
Vinyl bromide Acetylene
Aman Gour (JEE 2012, AIR 230)

Explanation: In ethylene dibromide, Br is present on a saturated carbon atom. Therefore. Like alkyl halides, it is a reactive molecule. Consequently, on heating with alcoholic KOH , it readily eliminates a molecule of HBr to form vinyl bromide in good yield. In contrast, due to the presence of Br on a doubly bonded carbon atom, vinyl bromide is a highly unreactive molecule and hence on heating with alcoholic KOH , it does not easily lose a molecule of HBr to form acetylene. Thus, with alcoholic KOH , the yield of acetylene is low. Therefore, to obtain acetylene in fairly good yield from vinyl bromide, a much stronger base than alcoholic KOH such as $\mathrm{NaNH}_{2}$ in liquid $\mathrm{NH}_{2}$ is usually used. Thus, dehydrohalogeneation of ethylene dibromide to acetylene is preferably carried out in the following two stages.

$\Theta$
With a strong base ( $\mathrm{NH}_{2}$ ), isomerisation also taken place to give terminal alkyne.


## Dehalogenation of tri/tetra halides




3. By dehalogenation of tetrahalides.: Tetrahaloalkanes when heated with zinc dust in methanol undergo dehalogenation to yield alkynes. For example


1,1,2,2-Tetrabromoethane

### 2.4 Dehalogenation of Haloforms

By dehalogenation of haloforms: Chloroform and iodoform on heating with silver powder undergo dehalogenation to form ethyne.



### 2.5 Kolbe's Electrolytic Reaction

Kolbe's electrolytic reaction: Acetylene can be prepared by electrolysis of a concentrated solution of sodium or potassium salt of maleic acid or fumaric acid. Thus,


This reaction is called Kolbe's electrolytic reaction and is believed to occur by the following steps:




At cathode: $2 \mathrm{H}^{+}+2 \mathrm{e}^{-} \longrightarrow[2 \mathrm{H}] \longrightarrow \mathrm{H}_{2}$

### 2.6 From $\alpha$-diketo

$\alpha$ - Diketone reacts with hydrazine $\left(\mathrm{NH}_{2}-\mathrm{NH}_{2}\right)$ to give bis-hydrazone which on oxidation with HgO gives unstable bis(diazo) compound which decomposes to give alkyne.


### 2.7 Industrial Method

$2 \mathrm{CH}_{4}(\mathrm{~g}) \xrightarrow{1773 \mathrm{~K}} \mathrm{CH} \equiv \mathrm{CH}(\mathrm{g})+3 \mathrm{H}_{2}(\mathrm{~g})$
The reaction is highly endothermic, yet the optimum time for this reaction is 0.01 second.
$\Delta n=n_{P}-n_{R}=4-2=2$ mol, and this causes a significant increase in $\Delta S$. At this high temperature, the $T \Delta S$ term in the equation $\Delta \mathrm{G}=\Delta \mathrm{H}-\mathrm{T} \Delta \mathrm{S}$ predominates, making $\Delta \mathrm{G}=-\mathrm{ve}$, although $\Delta \mathrm{H}=+\mathrm{ve}$.
But $\Delta \mathrm{H}_{\mathrm{f}}^{\circ}$ for $\mathrm{C}_{2} \mathrm{H}_{2}=+227 \mathrm{k} \mathrm{mol}^{-1}$ shows that acetylene is thermodynamically unstable, explodes readily, and gets converted into its elements.
$\mathrm{C}_{2} \mathrm{H}_{2}(\mathrm{~g}) \longrightarrow 2 \mathrm{C}(\mathrm{s})+\mathrm{H}_{2}(\mathrm{~g}), \Delta \mathrm{H}^{\circ}=-227 \mathrm{~kJ} \mathrm{~mole}^{-1}$

### 2.8 Berthelot Synthesis

Mearcellin Berthelot synthesized acetylene from its elements (carbon and hydrogen) by striking an electric arc between two electrodes in an atmosphere of $\mathrm{H}_{2}$ gas.


### 2.9 Higher Alkynes from Acetylene

Synthesis of higher alkynes from acetylene: Acetylene is first treated with sodium metal at 475K or with sodamide in liquid ammonia at 196 K to form sodium acetylide. This upon treatment with alkyl halides gives alkyne.

For example:





An analogous sequence usually terminal alkynes as starting material yields alkynes of the type $R C=C R^{\prime}$.


Dialkylation of acetylene can be achieved by carrying out the sequence twice.


## 3. PROPERTIES OF ALKYNES

(i) No cis-trans isomerism: The sp hybrid orbitals are linear, ruling out cis-trans stereoisomers in which substituents must be on different sides of the multiple bond.
(ii) Bonds: Bond length decreases in the following order: Alkane > Alkene > Alkyne

Bond length order: $-\underset{s p^{3}}{-\mathrm{C}}-\mathrm{H}>\underset{\mathrm{sp}^{2}}{\mathrm{C}}-\mathrm{H}>\equiv \underset{\mathrm{sp}}{\mathrm{C}}-\mathrm{H}$
Bond Strength order: $\equiv \underset{\mathrm{sp}}{\mathrm{C}}-\mathrm{H}>\underset{\mathrm{sp}^{2}}{\mathrm{C}}-\mathrm{H}>\underset{\mathrm{sp}}{ }{ }^{3}-\mathrm{H}$
As a general rule 'more the s - character of hybrid orbitals used by an atom, closer are the bonding electrons to the atom and shorter and stronger is any of its $\sigma$-bonds'.
(iii) Solubility and Dipole moment

Decreasing solubility in $\mathbf{H}_{\mathbf{2}} \mathbf{O}$ : Alkynes > Alkenes > Alkanes
Alkynes are slightly more soluble in $\mathrm{H}_{2} \mathrm{O}$ because they are somewhat more polar.
The dipole moment of alkyne is greater than those in alkene because $(C-C)$ sp bond is more polarized than $(\mathrm{C}-\mathrm{C}) \mathrm{sp}^{2}$ bond: this is because C with more s character is more electronegative.
Dipole moment of but-1-yne is 0.8 D and that of but-1-ene is 0.3 D .
Due to a slightly high polarity and dipole moment of alkynes than those of alkenes (and of alkenes higher than alkane) boiling point of alkynes > alkenes > alkanes, with the same number of $C$ atoms.
They go from gases to liquids to solids with increasing molecular weights. There is a little difference in boiling points of these hydrocarbons with similar C skeletons.
(iv) Smell: All alkynes are odourless except acetylene which has a garlic smell due to the presence of impurity, phosphine.
(v) Physical state: The first three members are gases, the next eight are liquids, and the rest are solids.
(vi) Melting and Boiling points: The m.p. and b.p. of alkynes are slightly higher than those of the corresponding alkenes and alkanes. This is probably due to the presence of a triple bond, alkynes have linear structures and hence their molecules can be more closed packed in the crystal lattice as compared to those of corresponding alkenes and alkanes.
(vii) Densities: Densities of alkynes like those of alkenes and alkanes increase as the molecular size increases. However, they are all lighter than water, since densities lie in the range $0.69-0.77 \mathrm{~g} / \mathrm{cm}^{2}$.
(viii) Stability: Alkynes are less stable than isomeric dienes as is evident from their heat of formation ( $\Delta \mathrm{H}_{\mathrm{f}}^{\circ}$ ) values.

(1,4-Pentadiene) $\quad+106.0 \mathrm{~kJ}$ mol-1

(ix) Reactivity: However, alkynes are less reactive than the corresponding alkenes towards electrophilic addition (EA) reaction (except catalytic hydrogenation), even though they contain two $\pi$-bonds. This is due to:
(a) Greater EN of sp-hybridised C atom of alkynes than $\mathrm{sp}^{2}$-hybridised C atom of alkenes which hold the $\pi$ - electrons of alkynes more tightly.
(b) Greater delocalisation of $\pi$-electrons in alkynes (because of cylindrical nature of their $\pi$-electrons cloud) than in alkenes. As a result, $\pi$-electrons of alkynes are less easily available for addition reaction than those of alkenes. So alkynes are less reactive than alkenes towards electrophilic addition reaction.

Catalytic hydrogenation, however, is an exception: Alkenes are adsorbed on the surface of catalyst only when the plane of $\pi$-bond approaches perpendicularly. In alkynes because of the cylindrical nature of $\pi$-bonds, approach by hydrogen along the axis of the cylinder is more effective and thus transition state in alkynes is less strained. So alkynes react faster than alkenes with $\mathrm{H}_{2}$.

## (x) Acidity of alkynes:

Terminal alkynes are more acidic than alkenes because alkyne C-atoms are sp hybridized and electrons in an s-orbital are more tightly held than in a p-orbital also s-electrons are closer to the nucleus.

Furthermore, since the $s p$ orbital has more s character ( $50 \% \mathrm{~s}$ ) than $\mathrm{sp}^{2}(33 \% \mathrm{~s})$ or $\mathrm{sp}^{2}$ orbital ( $25 \% \mathrm{~s}$ ), the electrons in sp orbital are more tightly held by the nucleus than electrons in $\mathrm{sp}^{2}$ or $\mathrm{sp}^{3}$ orbital.

In another words, sp -hybridised C is more EN than $\mathrm{sp}^{2}$, or $\mathrm{sp}^{3}$-hybridised C atom. Due to this greater EN , the electrons of $(\mathrm{C}-\mathrm{H})$ bonds are displaced more towards the C atom than towards the H atom. Therefore, the H -atom is less tightly held by the C atom and hence can be removed as a proton ( $\mathrm{H}^{+}$ion) by a strong base and consequently terminal alkynes behave as acids.


As a rule, as the $s$ character decreases from $s p$ to $s p^{2}$ to $s p^{3}$ hybridised $C$ atom, the acidic character of hydrocarbons decreases in the following order.

$$
\mathrm{Me}-\mathrm{HC}=\mathrm{O} \longleftrightarrow \mathrm{Me}-\mathrm{C}^{\oplus} \mathrm{H}-\mathrm{O}^{\Theta}
$$

## 4. REACTIONS OF ALKYNES

### 4.1 Acidic Character

(i) Formation of alkali metal acetylides: Ethyne and other terminal alkynes (Alkynes in which the triple bond is at the end of the carbon chain) or 1-alkynes react with strong bases such as sodium metal at 475 K or sodamide in liquid ammonia at 196 K to form sodium acetylides with evolution of $\mathrm{H}_{2}$ gas.



During these reactions, the acetylenic hydrogen is removed as a proton to form stable carbanions (acetylide ions) Sodium acetylide is decomposed by water regenerating acetylene. This shows that water is a stronger acid than acetylene and thus displaces acetylene from sodium acetylide.

(ii) Formation of heavy metal acetylides: Acetylenic hydrogens of alkynes can also be replaced by heavy metal ions such as $\mathrm{Ag}^{+}$and $\mathrm{Cu}^{+}$ions. For example. When treated with ammoniacal silver nitrate solution (Tollen's reagent), alkynes form a white precipitate of silver acetylides.



Similarly, with ammoniacal cuprous chloride solution, terminal alkynes form red ppt. of copper acetylides.


Unlike alkali metal acetylides are not decomposed by water. They can however, be decomposed with dilute mineral acids to regenerate the original alkynes.

$$
\begin{aligned}
& \underset{\text { (Disilver acetylide) }}{\mathrm{AgC} \equiv \mathrm{CAg}}+2 \mathrm{HNO}_{3} \longrightarrow \underset{\text { Acetylene }}{\mathrm{HC}} \equiv \mathrm{CH}+2 \mathrm{AgNO}_{3} \\
& \underset{\text { (Dicopper acetylide) }}{\mathrm{CuC} \equiv \mathrm{CCu}}+2 \mathrm{HCl} \longrightarrow \underset{\text { Acetylene }}{\mathrm{HC}}+2 \mathrm{CuCl}
\end{aligned}
$$

(iii) Formation of alkynyl Grignard reagents: Acetylne and other terminal alkynes react with Grignard reagents to form the corresponding alkynyl Grignard reagents. For example.


Alkynyl Grignard reagents like usual Grignard reagents can be used to prepare a variety of organic compounds.
Importance: The formation of metal acetylides can be used:
(a) For the separation and purification of terminal alkynes from non-terminal alkynes, alkanes and alkenes.
(b) To distinguish terminal alkynes from non-terminal alkynes or alkenes.

### 4.2 Hydrogenation / Reduction



## Alkene is an intermediate



The general case


The reaction takes place on the surface of Pt or Pd. Since the addition is twice, it is hard to see the syn addition.
But the reaction can be stopped at the alkene stage by using the following reagents.
(a) $\mathrm{H}_{2}+\mathrm{Pd}+\mathrm{BasO}_{4}+\mathrm{S}$ or quinoline in boiling xylene (called Lindlar's catalyst). In this case, reaction proceeds via syn addition of $\mathrm{H}_{2}$.
(b) $\mathrm{H}_{2}+\mathrm{P}-2$ or Brown catalyst $\left(\mathrm{H}_{2}+\mathrm{Ni}+\mathrm{B}\right)$ or $\left(\mathrm{H}_{2}+\mathrm{Ni}_{2} B\right)$. It also proceeds via syn addition of $\mathrm{H}_{2}$.
(c) Alkali metals $(\mathrm{Na}, \mathrm{K}, \mathrm{Cs})+$ liq. $\mathrm{NH}_{3}$ and $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{OH}$ (It is called Birch reduction). It proceeds via anti-addition of $\mathrm{H}_{2}$.
(d) $\mathrm{LiAlH}_{4}(\mathrm{LAH})$. It reduces $(\mathrm{C} \equiv \mathrm{C})$ to $(\mathrm{C}=\mathrm{C})$ via anti addition of $\mathrm{H}_{2}$; does not reduce $(\mathrm{C}=\mathrm{C})$ to $(\mathrm{C}-\mathrm{C})$ bond; reduces $(C=C)$ to $(C-C)$ only when the phenyl group is attached to the $-C$ atom of the double bond.
$\beta \quad \alpha$
(E.g. $\mathrm{Ph}-\mathrm{CH}=\mathrm{CH}-\mathrm{CH}_{3}$ to $\mathrm{Ph}-\mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{CH}_{3}$ )

## Reduction of alkynes to (Z) - alkenes

Lindlar's catalyst: Pd , quinoline, Pb and $\mathrm{CaCO}_{3}$ poisons the metal catalyst. So that the $\mathrm{H}_{2}$ adds only to the alkyne. Not reactive enough for alkenes.



Reduction of alkynes to (E) - alkenes. Reaction via Na radical dissolved in liquid $\mathrm{NH}_{3}$.
$\mathrm{Na} \cdot+\mathrm{NH}_{3} \rightarrow \mathrm{Na}^{+}+\mathrm{e}^{-}\left[\mathrm{NH}_{3}\right]_{n}$
Solvated electron
Mechanism of $\mathrm{Na} / \mathrm{NH}_{3}$ reductions: $\quad \mathrm{Na}^{+} \xrightarrow{-33^{\circ} \mathrm{C}} \mathrm{Na}^{+}+\mathrm{NH}_{3}\left(\mathrm{e}^{-}\right)$


### 4.3 Addition of $\mathbf{X}_{2}$

Like alkenes, 1 mol of alkynes adds 2 moles of halogens and proceeds via anti-addition of halogens, and orange or brown colour of $\mathrm{Br}_{2}$ is discharged (test for unsaturation)
(I) HC


## Mechanism:

Like alkenes it proceed via the formation of intermediate cyclic bromonium ion. First, electrophile $(\operatorname{Br} \oplus)$ add and then the addition of nucleophile $\left(\mathrm{Br}^{\ominus}\right)$ from anti-position take place. That is why this reaction is called EA (electrophilic addition) reaction. The reaction is stereospecific (attack by electrophile in a specific manner).


### 4.4 Addition of Water

Hydration of alkynes


(Oxymercuration-demercuriation of triple bond)



### 4.5 Addition of HX


(i)
(I) $\mathrm{HC} \equiv \mathrm{CH} \xrightarrow{\mathrm{HBr}} \mathrm{HC=C} \underset{\substack{\mathrm{CBr}}}{\mathrm{M}^{+} \mathrm{Br} \mathrm{Br}^{-}}$(EA) $)$

11.72
(II)



1-Bromobut-1-ene
1,1-Dibromobutane
(Butylidene bromide)

## PLANCESS CONCEPTS

The order of reactivity of HX is $\mathrm{HI}>\mathrm{HBr}>\mathrm{HCl}>\mathrm{HF}$
Aishwarya Karnawat (JEE 2012 AIR 839)

### 4.6 Addition of Acids

Addition of carboxylic acids: When acetylene is passed into warm acetic acid in presence of mercury salts, first vinyl acetate and then ethylidene diacetate is formed.


Vinyl acetate is used for the manufacture of vinyl resin. Ethylidene diacetate, when heated rapidly to 573-673K, gives acetic anhydride and acetaldehyde.


### 4.7 Addition of HCN

The reaction is nucleophilic addition (NA) initiated by $\mathrm{CN}^{-}$from NaCN .


### 4.8 Addition to Other Compounds






### 4.9 Ethyne + Methanal

The addition of ethyne to unsaturated link like ( $>\mathrm{C}=\mathrm{O}$ ) is called ethinylation. Acetylene or terminal alkyne containing $\equiv \mathrm{CH}$ (a methine H -atom) reacts with a carbonyl group ( $>\mathrm{C}=\mathrm{O}$ ) in the presence of sodium or potassium alkoxide (RONa or ROK) or $\mathrm{NaNH}_{2}$ to give alkyndiol with a small amount of alkyneol.




But-2-yn-1,4-diol

### 4.10 Nucleophilic Addition

Because of the greater electronegativity of the sp-hybridized carbons as compared to $\mathrm{sp}^{2}$ hybridized carbons, alkynes are more susceptible to nucleophilic addition reactions than alkenes.

For example, when acetylene is passed into methanol at $433-473 \mathrm{~K}$ in presence of a small amount ( $1-2 \%$ ) of potassium methoxide under pressure, methyl vinyl ether is formed.


Methyl vinyl ether is used for making polyvinyl ether plastics.

### 4.11 Propargylic Halogenation

Alkenes undergo allylic substitution with NBS, whereas alkynes undergo propargylic halogenation with $\mathrm{NBS}, \mathrm{Cl}_{2} \mathrm{SO}_{2}$ (sulfuryl chloride). $\mathrm{Me}_{3} \mathrm{COCl}\left(\mathrm{t}\right.$-butyl hypchlorite), and $\mathrm{Cl}_{2}$ at $500^{\circ} \mathrm{C}$.


Reaction proceeds via free radical mechanism to give propargylic radical, which is stabilized by resonance via extended $\pi$-bonding.

### 4.12 Hydroxylation of Alkynes

Hydroxylation of alkyne with aqueous or neutral $\mathrm{KMnO}_{4}$ solution (Baeyer's reagent) test for unsaturation)
Pink colour of $\mathrm{KMnO}_{4}$ is discharged and brown black precipitate of $\mathrm{MnO}_{2}$ is obtained. This reaction converts alkynes first to enediols and then further gives tetraols, which being unstable lose $\mathrm{H}_{2} \mathrm{O}$ to give diketones.


(Break two bonds and add
O atom on both sides)


But-1-en-3-yne
3,4-Dihydroxy-2-oxo butan-1-al

### 4.13 Oxidation Reactions

### 4.13.1 Oxidative Cleavage

(a) Alkynes are oxidatively cleaved in alkaline or acidic conditions at higher temperature, as mentioned.

$$
\begin{gathered}
\mathrm{R}-\mathrm{C} \not \equiv \Longrightarrow \mathrm{R}-\mathrm{COOH} \\
\mathrm{H}-\mathrm{C} \not \equiv \equiv[\mathrm{H}-\mathrm{COOH}] \Longrightarrow \mathrm{CO}_{2}
\end{gathered}
$$

(i)

(ii)

(iii)

$\mathrm{CrO}_{3}$ in $\mathrm{CH}_{3} \mathrm{COOH}$ oxidizes alkenes but does not affect alkynes

### 4.14 Ozonolysis of Alkynes

Alkyne form ozonides with $\mathrm{O}_{3}$ and are decomposed by $\mathrm{H}_{2} \mathrm{O}$ to give diketones, which are then oxidised by $\mathrm{H}_{2} \mathrm{O}_{2}$ or $\mathrm{KMnO}_{4} / \mathrm{NaIO}_{4}$ or peracids; given acids, and on reduction with metal / acid, LAH , or $\mathrm{NaBH}_{4}$ give diols.
(i)


Terminal alkynes give HCOOH as one of the products which is further oxidized to $\mathrm{CO}_{2}$
These oxidative cleavage reactions are rarely used for synthesis because alkynes are not readily available. They are used to locate the position of a triple bond in an alkyne.

### 4.15 Hydroboration of Alkynes

- Terminal alkynes of HBO reaction give aldehyde, whereas internal alkynes give ketone.
- Alkynes react with $\mathrm{BH}_{3}$ or $\mathrm{B}_{2} \mathrm{H}_{6}$ complexed with THF (tetrahydrofuran $\quad$ ) to give trivinyl borane, which upon subsequent treatment with alkaline $\mathrm{H}_{2} \mathrm{O}_{2}$ gives alcohols corresponding to anti-Markovnikov's addition of $\mathrm{H}_{2} \mathrm{O}$ to alkynes, which on tautomerisation give corresponding aldehydes or Ketones.
- If trivinyl borane is treated with acid $\left(\mathrm{CH}_{3} \mathrm{COOH}\right.$ ), it gives cis-alkene (proceeds via syn-addition of $\mathrm{H}_{2}$ ).



Butan-2-one

### 4.16 Polymerisation

## 1. Linear Polymerisation



## 2. Cyclic Polymerisation



3. Propyne



1,3,5-Trimethyl benzene or Mesitylene

### 4.17 Isomerization

When alkynes are heated with $\mathrm{NaNH}_{2}$ in an inert solvent such as kerosene oil or paraffin oil, they undergo isomerization. i.e. 2 -alkynes isomerize to 1 -alkynes and vice versa.


## Uses of alkynes

(i) Acetylene and its derivatives are widely used in synthetic organic chemistry for synthesis of cis-and transalkenes, methyl ketones etc.
(ii) Oxyacetylene flame is used for cutting and welding of metals.
(iii) Acetylene is used as illuminant in hawker's lamp and in light houses.
(iv) Acetylene is used for the ripening of fruits and vegetables.
(v) Acetylene is used for manufacture of ethyl alcohol, acetaldehyde, acetic acid, vinyl plastics, synthetic rubbers such as Buna N and synthetic fibers such as Orlon.

## PLANCESS CONCEPTS

Difference between Alkenes and Alkynes

|  | Test | Alkene | Alkyne |
| :---: | :--- | :--- | :--- |
| 1. | Flame observed on comnustion | Luminous | Smoky |
| 2. | $\mathrm{Br}_{2} / \mathrm{CCl}_{4}$ solution | Orange colour is discharged | Orange colour is discharged |
| 3. | Cold aq. or alkaline $\mathrm{KMnO}_{4}$ <br> solution (Baeyer's reagent) | Pink colour is discharged with the <br> formation of diol compound | Pink colour is discharged with the <br> formation of dicarbonyl compound |
| 4. | Ammoniacal AgNO3 solution <br> $[\mathrm{Ag}(\mathrm{NH} 3) 2]+($ Tollens reagent $)$ | No action | Whote ppt. of silver alkynide (given <br> by only terminal alkynes) |
| 5. | Ammoniacal CuCl solution | No action | Red ppt. of copper acetylide (given <br> by only terminal alkynes) |
| 6. | Reactivity towards electrophilic <br> addition reaction (e.g. $\mathrm{HX}, \mathrm{X}_{2}$ ) | Alkenes are more reactive than alkynes |  |
| 7. | Reactivity towards cataytic | Alkynes are more reactive than alkenes |  |
| 8. | Reactivity towards nuclephilic <br> addition reaction (e.g. $\mathrm{CN}^{+}$) | Alkynes are more reactive than alkenes |  |

## POINTS TO REMEMBER




## Solved Examples

## JEE Main/Boards

Example 1: Identify A to C .


Sol: It is an example of ethinylation (addition of ethyne to unsaturated link like $\geq \mathrm{c}$ group).
First step is the reaction with the sodium metal to form disodium acetylide with evolution of hydrogen gas.
This reaction shows the acidic character of alkynes.
During this step acetylenic hydrogens are removed as a proton to form stable carbanion.
Now the carbanion reacts with two moles of carbon dioxide and forms an adduct which on treatment with acid yields acid.

Next step is reduction using Lindlar Catalyst $\left(\mathrm{H}_{2} / \mathrm{Pd}-\mathrm{CaCO}_{3}\right)$ the addition is syn.
On treatment with $\left(\mathrm{NH}_{3}+\mathrm{EtOH}\right)$ trans product is formed thus the addition is anti.


Example 2: Identify B to G.


Sol:

- Addition of $\mathrm{Br}_{2}$ across double bond forms 1,2-dibromo propane.
- This on treatment with soda lime undergoes double dehalogenation to form an alkyne
- Now alkyne on the treatment with a brominating agent (NBS- N -Bromosuccinimide) and $\mathrm{CCl}_{4}$ as a solvent in presence of light.
- On the other side, the first step is Allylic chlorination using $\mathrm{Cl}_{2}$ at $500^{\circ} \mathrm{C}$.
- The reaction proceeds through radical formation.
- Now Allyl chloride undergoes chain lengthening reaction to form Hex-1-en-5-yne.
- The reaction is known as Corey house synthesis



1,2-Dibromopropane Prop-1-yne

$\left\lvert\,$| $\mathrm{Cl}_{2}$ at $500^{\circ} \mathrm{C}$ | Propar- |
| :--- | :---: |
| Allylic | gylic |
| radical | radical |
| chlorination | brominating |$\downarrow \mathrm{NBS}+\mathrm{CCl}_{4}+\mathrm{hv}\right.$


$\left(\begin{array}{c}\text { Allyl chloride } \\ \text { or } \\ \text { 3-Chloro prop-1-ene }\end{array}\right)$

| Coerey | 1. Li |
| :--- | :--- |
| -House |  |
|  | 2. Cul |


(F)

Diallyl lithium cuperate

(D)
$\left(\begin{array}{c}\text { 3-Bromoprop-1-yne } \\ \text { or } \\ \text { propargylic } \\ \text { bromide }\end{array}\right)$

Example 4: There are two path (a) and (b) For the preparation of a compound (a) $\left(\begin{array}{ll}123 \\ \mathrm{Me}\end{array} \mathrm{M}^{4} \mathrm{Me}\right)$ (2-methylpent-1-en-3-yne), which path is correct and why? Also name the paths (a) and (b). Path (a)

Sol: Path (b) is correct because in path (a) compound $(\underset{\mathrm{Br}}{\stackrel{\mathrm{Me}}{\sim}})$ vinyl halide which is not reactive, so the reaction does not occur.
Path (a) is the alkylation of alkyne, whereas path (b) is propinylation (type of ethinylation), i.e., addition of propyne to unsaturated link like (> $\mathrm{C}=\mathrm{O}$ ) group.


## JEE Advanced/Boards

Example 1: Complete the following missing reagents


Sol: (i)
(a) $\mathrm{HBr}+$ Peroxide

First step is addition of HBr , it takes place according to Markovnikov's rule.
(b) (1) Li (2) CuI (Corey - House): Second step is Corey House synthesis.
(c)
 Third step is the reaction with alkyl halide.
(d) $\mathrm{Br}_{2} / \mathrm{CCl}_{4}$

Bromination using $\mathrm{Br}_{2}$ and $\mathrm{CCl}_{4}$ as a solvent
(e) $\mathrm{KOH}(\mathrm{s})$, at 473 K

Last step is double dehalogenation using base like KOH at 473 K
(ii) Cis compound on addition with $\mathrm{Br}_{2}$ (Anti addition) forms a racemic mixture or enantiomer. Trans compound on addition with $\mathrm{Br}_{2}$ (Anti addition) produces Meso compound (it contains plane of symmetry)


Example 2: Convert the following


Sol:
(a)

(b)



Example 3: Convert the following.


Write the structure of (E).

Sol: The structure of $E$ is


First prepare $3 C$ - atom alkyne $(\mathrm{Me}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H})$ from $(A)$ and then react 4 C - atom $R X$ with alkynide ion to get ( E ).

## Reaction:


$2 \mathrm{NaNH}_{2}$


(D)
(E)

Example 4: Complete the following:


Sol:

(A)

Sodium ethynide
(B)

But-1-yne
(C)


(D)

Sodium butynide

## JEE Main/Boards

## Exercise 1

Q. 1 What are alkenes? Discuss briefly the various methods used for the preparation of alkenes. Describe with a labelled diagram the laboratory preparation of ethane from ethanol.
Q. 2 Write the major products of the hydrocarbon and oxidation of:
(A) 1-Ethyl cyclopentene
(B) Methylene cyclopentene
Q. 3 Give the missing compounds in the following:
(A)

(B)

(C)

(D)

Q. 4 Complete the following equation:
(A) $\mathrm{CH}_{3} \mathrm{C} \equiv \mathrm{CCH}_{3} \xrightarrow[\mathrm{Pt}_{2}]{\mathrm{H}_{2}}$ (A)

$$
\xrightarrow[\text { Pt }]{\mathrm{H}_{2}} \mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{3}
$$

(B)


$$
\text { (B) }+\mathrm{NaOC}_{2} \mathrm{H}_{5}
$$

Q. 5 Give an account of physical and chemical reactions of alkenes.
Q. 6 What are alkynes? How are they prepared in the laboratory?
Q. 7 Give an account of physical properties and chemical reactions of alkynes.
Q. 8 (A) Starting with any alkyne, prepare ethyl cyclohexyl ethyne.
(B) Starting with cyclohexyl ethyne, prepare acetyl cyclohexane.
Q. 9 (A) Complete the following reactions:


(B) Complete the following reactions:


$$
\xrightarrow[\text { Lindlar's catalyst }]{\mathrm{H}_{2}} \mathrm{~B} \xrightarrow[\text { Zn(Cu) }]{\mathrm{CH}_{2} \mathrm{I}_{2}} \mathrm{C}
$$

Q. 10 Predict the product of the following:
(A)

(B)

(C)

Q. 11 (A) Give reasons for the following:

When 1-penten-4-yne is treated with HBr in equimolecular proportion, the addition takes place on double bond and not on triple bond yielding thereby the product

## $\mathrm{CH}_{3} \mathrm{CH}(\mathrm{Br}) \mathrm{CH}_{2} \mathrm{C} \equiv \mathrm{CH}$.

(B) Provide a suitable mechanism for the following reaction.

Q. 12 Three isomeric alkenes $A, B$ and $C,\left(C_{5} H_{10}\right)$ are hydrogenated to yield 2-methylbutane $A$ and $B$ gave the same $3^{\circ} \mathrm{ROH}$ on oxymercuration-demercuration $B$ and $C$ give different $1^{\circ}$ ROH's on hydroborationoxidation supply the structures of $\mathrm{A}, \mathrm{B}$ and C .
Q. 13 3,3-dimethyl-1-butene and HI react to give two products, $\mathrm{C}_{6} \mathrm{H}_{13}$ I. On reaction with alc. KOH one isomer, (1) gives back 3,3-dimethyl-1-butene the other (J) gives an alkene that is reductively ozonized to give $\mathrm{Me}_{2} \mathrm{C}=\mathrm{O}$. Give the structures of (I) and (J) and explain the formation of the latter.
Q. 14 Identify $X, Y$ and $Z$ in the following sequence of reaction giving stereo chemical structure wherever possible.


## Exercise 2

## Single Correct Choice Type

Q. 1 On heating $\mathrm{CH}_{3} \mathrm{COONa}$ with soda lime the gas evolved will be
(A) $\mathrm{C}_{2} \mathrm{H}_{2}$
(B) $\mathrm{CH}_{4}$
(C) $\mathrm{C}_{2} \mathrm{H}_{6}$
(D) $\mathrm{C}_{2} \mathrm{H}_{4}$
Q. 2 The addition of $\mathrm{Br}_{2}$ to trans-2-butene produces
(A) (+) 2, 3-dibromobutane
(B) (-) 2, 3-dibromobutane
(C) rac -2,3-dibromobutane
(D) meso -2,3-dibromobutane
Q. 3 Isomers which can be interconverted through rotation around a single bond are
(A) Conformers
(B) Diastereomers
(C) Enantiomers
(D) Positional isomers
Q. 4 The olefin, which on ozonolysis gives $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CHO}$ and $\mathrm{CH}_{3} \mathrm{CHO}$
(A) But-1-ene
(B) But-2-ene
(C) Pent-1-ene
(D) Pent-2-ene
Q. 5 Which of the C C bond is strongest?
(A) Formed by $\mathrm{sp}^{3}-\mathrm{sp}^{3}$ hybridised carbon atoms (as in alkanes)
(B) Formed by $\mathrm{sp}^{2}-\mathrm{sp}^{2}$ hybridised carbon (as in alkenes)
(C) Formed by $\mathrm{sp}-\mathrm{sp}$ hybridised carbon atoms (as in alkynes)
(D) None of these
Q. 6 End product of the following sequence is:

(A) Ethanol
(B) Ethyl hydrogen sulphate
(C) Ethanal
(D) Ethylene glycol
Q. 7 The treatment of $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{MgI}$ with water produces
(A) Methane
(B) Ethane
(C) Ethanal
(D) Ethanol
Q. 8 When isobutene is brominated, the percentage of

(A) $0 \%$
(B) $83 \%$
(C) $10 \%$
(D) $100 \%$
Q. 9 Propene can be converted into 1-propanol by oxidation. Which set of the reagents is used to effect the conversion?
(A) $\mathrm{OsO}_{4}-\mathrm{CHCl}_{3}$
(B) $\mathrm{O}_{3} / \mathrm{Zn}-\mathrm{H}_{2} \mathrm{O}$
(C) Alkaline and cold $\mathrm{KMnO}_{4}$
(D) $\mathrm{B}_{2} \mathrm{H}_{6}$ and alk. $\mathrm{H}_{2} \mathrm{O}_{2}$
Q. 10 In the following sequence of reactions, identify the product ( d )
$\mathrm{CH} \equiv \mathrm{CH} \xrightarrow{\mathrm{HBr}}(A) \xrightarrow{\mathrm{HBr}}(B) \xrightarrow{\text { alk. } \mathrm{KOH}}(C)$ $\xrightarrow{\mathrm{NaNH}_{2}}(\mathrm{D})$
(A) Ethanol
(B) Ethyne
(C) Ethanal
(D) Ethene
Q. 11 Which one is highly unstable?
(A) $\mathrm{CH}_{2}=\mathrm{CH}-\mathrm{OH}$
(B) $\mathrm{CH}_{2}=\mathrm{CH}-\mathrm{Cl}$
(C) $\mathrm{CH}_{2}=\mathrm{CH}-\mathrm{CH}_{2}-\mathrm{Cl}$
(D) $\mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{CH}$

## Previous Years' Questions

Q. 1 Marsh gas mainly contains
(1980)
(A) $\mathrm{C}_{2} \mathrm{H}_{2}$
(B) $\mathrm{CH}_{4}$
(C) $\mathrm{H}_{2} \mathrm{~S}$
(D) CO
Q. 2 Which of the following will decolourise alkaline $\mathrm{KMnO}_{4}$ solution?
(1980)
(A) $\mathrm{C}_{3} \mathrm{H}_{8}$
(B) $\mathrm{CH}_{4}$
(C) $\mathrm{CCl}_{4}$
(D) $\mathrm{C}_{2} \mathrm{H}_{4}$
Q. 3 The compound 1,2-butadiene has
(1983)
(A) Only sp-hybridised carbon atoms
(B) Only $\mathrm{sp}^{2}$-hybridised carbon atoms
(C) Both sp and $\mathrm{sp}^{2}$-hybridised carbon atoms
(D) $\mathrm{sp}, \mathrm{sp}^{2}$ and $\mathrm{sp}^{3}$-hybridised carbon atoms
Q. 4 When propyne in treated with aqueous $\mathrm{H}_{2} \mathrm{SO}_{4}$ in presence of $\mathrm{HgSO}_{4}$, the major product is
(1983)
(A) Propanal
(B) Propyl hydrogen sulphate
(C) Acetone
(D) Propanol
Q. 5 Bayer's reagent is
(1984)
(A) Alkaline permanganate solution
(B) Acidified permanganate solution
(C) Neutral permanganate solution
(D) Aqueous bromine solution
Q. 6 Acidic hydrogen is present in
(1985)
(A) Ethyne
(B) Ethene
(C) Benzene
(D) Ethane
Q. 7 Hydrogenation of the adjoining compound in the presence of poisoned palladium catalyst gives (2001)

(A) An optically active compound
(B) An optically inactive compound
(C) A racemic mixture
(D) A diastereomeric mixture
Q. 8 The reaction of propene with HOCl proceeds via the addition of
(2001)
(A) $\mathrm{H}^{+}$in the first step
(B) $\mathrm{Cl}^{+}$in the first step
(C) $\mathrm{OH}^{-}$in the first step
(D) $\mathrm{Cl}^{+}$and $\mathrm{OH}^{-}$single step
Q. 9 In the presence of peroxide, hydrogen chloride and hydrogen iodide do not give anti Markownikoff's addition to alkenes because
(2001)
(A) Both are highly ionic
(B) One is oxidising and the other is reducing
(C) One of the steps is endothermic in both the cases
(D) All the steps are exothermic in both the cases
Q. 10 Which of the following compound will exhibit geometrical isomerism?
(2015)
(A) 1-Phenyl-2-butene
(B) 3-Phenyl-1-butene
(C) 2-Phenyl-1-butene
(D) 1,1-Diphenyl-1-propane
Q. 11 The reaction of propene with $\mathrm{HOCl}\left(\mathrm{Cl}_{2}+\mathrm{H}_{2} \mathrm{O}\right)$ proceeds through the intermediate :
(2016)
(A) $\mathrm{CH}_{3}-\mathrm{CH}^{+}-\mathrm{CH}_{2}-\mathrm{Cl}$
(B) $\mathrm{CH}_{3}-\mathrm{CH}(\mathrm{OH})-\mathrm{CH}_{2}+$
(C) $\mathrm{CH}_{3}-\mathrm{CHCl}-\mathrm{CH}_{2}$
(D) $\mathrm{CH}_{3}-\mathrm{CH}^{+}-\mathrm{CH}_{2}-\mathrm{OH}$
Q. 12 The major organic compound formed by the reaction of 1, 1, 1-trichloroethane with silver powder is
(2014)
(A) Acetylene
(B) Ethene
(C) 2-Butyne
(D) 2-Butene
Q. 13 In the reaction,

the product $C$ is
(2014)
(A) Acetaldehyde
(B) Acetylene
(C) Ethylene
(D) Acetyl chloride
Q. 14 In the following sequence of reactions, the alkene affords the compound ' B '


The compound $B$ is
(2008)
(A) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CHO}$
(B) $\mathrm{CH}_{3} \mathrm{COCH}_{3}$
(C) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{COCH}_{3}$
(D) $\mathrm{CH}_{3} \mathrm{CHO}$
Q. 15 The hydrocarbon which can react with sodium in liquid ammonia is
(2008)
(A) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{C} \equiv \mathrm{CCH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{3}$
(B) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{C} \equiv \mathrm{CH}$
(C) $\mathrm{CH}_{3} \mathrm{CH} \equiv \mathrm{CHCH}_{3}$
(D) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{C} \equiv \mathrm{CCH}_{2} \mathrm{CH}_{3}$
Q. 16 The treatment of $\mathrm{CH}_{3} \mathrm{MgX}$ with $\mathrm{CH}_{3} \mathrm{C} \equiv \mathrm{C}-\mathrm{H}$ produces
(2008)
(A) $\mathrm{CH}_{3}-\mathrm{CH} \equiv \mathrm{CH}_{2}$
(B) $\mathrm{CH}_{3} \mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3}$
(C)

(D) $\mathrm{CH}_{4}$
Q. 17 The alkene that exhibits geometrical isomerism is:
(2009)
(A) Propene
(B) 2-methyl propene
(C) 2-butene
(D) 2-methyl -2- butane
Q. 18 One mole of a symmetrical alkene on ozonolysis gives two moles of an aldehyde having a molecular mass of 44 u . The alkene is
(2010)
(A) Propene
(B) 1-butene
(C) 2-butene
(D) Ethene

## Assertion and Reason Type

Read the following questions and answer as per the direction given below:
(A) Statement-I is true; statement-II is true;
statement-II is a correct explanation of statement-I.
(B) Statement-I is true; statement-II is true; statement-II is not the correct explanation of statement-I.
(C) Statement-I is true; statement-II is false.
(D) Statement-I is false; statement-II is true.
Q. 19 Statement-I: Addition of $\mathrm{Br}_{2}$ to 1-butene gives two optical isomers.
Statement-II: The product contains one asymmetric carbon.
(1998)
Q. 20 Statement-I: 1-butene on reaction with HBr in the presence of a peroxide produces 1 bromobutane.

Statement-II: It involves the formation of a primary radical.
(2000)
Q. 21 Statement-I: Dimethyl sulphide is commonly used for the reduction of an ozonide of an alkene to get the carbonyl compound.

Statement-II: It reduces the ozonide giving water soluble dimethyl sulphoxide and excess of it evaporates.
(2000)
Q. 22 Statement-I: Addition of bromine to trans-2butene yields meso-2, 3-dibromo butane.
Statement-II: Bromine addition to an alkene is an electrophilic addition.

## JEE Advanced/Boards

## Exercise 1

Q. 1 Compound ( X ) $\left(\mathrm{C}_{5} \mathrm{H}_{8} \mathrm{O}\right)$ does not appreciably react with Lucas reagent at room temperature but gives a precipitate with silver nitrate (ammoniacal). With Me $\mathrm{Mg} \mathrm{Br}, 0.42 \mathrm{gms}$ of $(\mathrm{X})$ gives 22400 mL of $\mathrm{CH}_{4}$ at STP. Treatment of $\mathrm{H}_{2}$ in the presence of Pt catalyst followed by boiling with excess HI, gives n-pentane. Suggest the structure for $(X)$ and write equations involved.
Q. 2 Compound (A) $\mathrm{C}_{5} \mathrm{H}_{10}$ decolourizes $\mathrm{Br}_{2}$. When (A) is treated with hot $\mathrm{KMnO}_{4}$, two acids (B) and (C) are formed. When $(B)$ is treated with soda lime, methane is formed. When $(C)$ is heated with soda lime ethane is formed. What are the structure of (A), (B) and (C).
Q. 3 An organic compound (A) $\mathrm{C}_{6} \mathrm{H}_{12}$ is treated with $\mathrm{Cl}_{2}$ in the presence of $\mathrm{CCl}_{4}$ gives $(B) \mathrm{C}_{6} \mathrm{H}_{12} \mathrm{Cl}_{2}$. Compound
(B) on treatment with alcoholic KOH followed by $\mathrm{NaNH}_{2}$ resulting in the formation of compound (C) $\mathrm{C}_{6} \mathrm{H}_{10}$. Compound (C) on treatment with $\mathrm{H}_{2}$ (gas) over platinum forms 2-methyl pentane. Compound (C) gives ammoniacal $\mathrm{AgNO}_{3}$ test. Ozonolysis of (A) gives two aldehydes (D) and (E) where E is acetaldehyde. Identify $A, B, C, D$ and $E$.
Q. 4 A hydrocarbon $(A)$ is treated with excess of HCl . $A$ dihalogen derivative ( $B$ ) is formed. Compound (B) on treatment with aqueous KOH gives C . Compound (C) can be obtained by heating calcium salt of 2-methyl propanoic acid with calcium acetate what are $A, B$ and $C$ ?

Q. 6 An organic compound (A) $\mathrm{C}_{9} \mathrm{H}_{12}$ gave (B) $\mathrm{C}_{8} \mathrm{H}_{6} \mathrm{O}_{4}$ on oxidation by alkaline $\mathrm{KMnO}_{4}$. (B) on heating does not
form anhydride. Also (B) reacts with $\mathrm{Br}_{2}$ in the presence of iron to give only one monobromo derivative (C). $\mathrm{C}_{8} \mathrm{H}_{5} \mathrm{BrO}_{4}$. What are $\mathrm{A}, \mathrm{B}$ and C .
Q. 7 A hydrocarbon (A) $\mathrm{C}_{9} \mathrm{H}_{10}$ adds $\mathrm{Cl}_{2}$ to give (B) $\mathrm{C}_{9} \mathrm{H}_{10} \mathrm{Cl}_{2}$. Hydrolysis of (B) gives (C) $\mathrm{C}_{9} \mathrm{H}_{12} \mathrm{O}_{2}$. Oxidation of (A) gave two acids identified as benzoic acid and acetic acids. What are $A, B$ and $C$ ?
Q. 8 An organic compound (A) $\mathrm{C}_{9} \mathrm{H}_{10} \mathrm{Br}_{2}$ forms $\mathrm{C}_{9} \mathrm{H}_{11} \mathrm{OBr}$ with caustic soda solution the later is resistant to further action of alkali. (B) on treating with $\mathrm{H}_{2} \mathrm{SO}_{4}$ converts into (C) with on ozonolysis gives ethanal and another compound (D). Identify (A) to (D) and give reactions.
Q. 9 An organic compound (A) $\mathrm{C}_{6} \mathrm{H}_{10^{\prime}}$ on reduction first gives (B) $\mathrm{C}_{6} \mathrm{H}_{12}$ and finally (C) $\mathrm{C}_{6} \mathrm{H}_{14}$. (A) on ozonolysis followed by hydrolysis gives two aldehydes (D) $\mathrm{C}_{2} \mathrm{H}_{4} \mathrm{O}$ and ( E ), $\mathrm{C}_{2} \mathrm{H}_{2} \mathrm{O}_{2}$. Oxidation of (B) with $\mathrm{KMnO}_{4}$ gives the acid (F) $\mathrm{C}_{4} \mathrm{C}_{8} \mathrm{O}_{2}$. Determine the structures of (A) to (F) with proper reasoning.
Q. 10 An organic compound (A) of molecular formula $\mathrm{C}_{5} \mathrm{H}_{8}$ when treated with Na in liquid ammonia followed by reaction with n-propyl iodide yields (B) $\mathrm{C}_{8} \mathrm{H}_{14}$. (A) gives a ketone $\mathrm{C}_{5} \mathrm{H}_{10} \mathrm{O}$ when treated with dil. $\mathrm{H}_{2} \mathrm{SO}_{4}$ and $\mathrm{HgSO}_{4}$. (B) on oxidation with alkaline. $\mathrm{KMnO}_{4}$ gives two isomeric acids (D) and (E) $\mathrm{C}_{4} \mathrm{H}_{8} \mathrm{O}_{2}$. Give structures of compounds $(A)$ to $(D)$ with proper reasoning.
Q. 11 An organic compound (A) $\mathrm{C}_{8} \mathrm{H}_{6}$, on treatment with dil. $\mathrm{H}_{2} \mathrm{SO}_{4}$ containing mercuric sulphate gives a compound ( $B$ ), which can also be obtained from a reaction of benzene with acid chloride in the presence of anhydrous $\mathrm{AlCl}_{3}$. The compound (B) when treated with iodine in aq. KOH , yields C and a yellow compound (D). Identify $A, B, C$ and $D$ with justification.
Q. 12 An organic compound $A\left(\mathrm{C}_{5} \mathrm{H}_{8}\right)$ on hydrogenation gives compound $B\left(C_{5} H_{12}\right)$. Compound $A$ on ozonolysis gives formaldehyde and 2-ketopropanal. Deduce structure of compound $A$.
Q. 13 What are the possible products from 1,4-addition of HBr on state which carbocation is more stable


A Predict the structure of A and explain.
Q. 15 A hydrocarbon ( P ) of the formula $\mathrm{C}_{8} \mathrm{H}_{10^{\prime}}$ on ozonolysis, gives compound $\mathrm{Q}\left(\mathrm{C}_{4} \mathrm{H}_{6} \mathrm{O}_{2}\right)$ only. The compound ( Q ) can also be obtained from an alkyl bromide, $\mathrm{R}\left(\mathrm{C}_{3} \mathrm{H}_{5} \mathrm{Br}\right)$ upon treatment with Mg in dry ether, followed by $\mathrm{CO}_{2}$ and acidification. Identify (P), (Q) and $(R)$ and give equations for the reactions.
Q. 16 Two isomeric alkyl bromides (A) and (B) have molecular formula $\mathrm{C}_{5} \mathrm{H}_{11} \mathrm{Br}$. (A) on treatment with alcoholic KOH gives two isomers (C) and (D) of formula $\mathrm{C}_{5} \mathrm{H}_{10}$. (C) on ozonolysis gives formaldehyde and 2-methylpropanal. (B) with alcoholic KOH gives (D) and (E). Catalytic hydrogenation of (C), (D) and (E) gives (F) $\mathrm{C}_{5} \mathrm{H}_{12}$. Identify (A) to (F).
Q. 17 How would you bring about the following conversion.
(i) 1, 3-Butadiene to racemic - 1, 4-dibromo -2, 3-dideutrobutene
(ii)

(iii)

(iv) Z-2, 3-Dideutero-2-butene to racemic-erythro-2-Bromo-2,3-Didenterobutane
(v) 1-Methyl cyclopentene to trans-2-Methyl cyclopentane
(vi)

to (S, S) - Glycol + R, S Glycol
Q. 18 A compound ( A ) has $\mathrm{C}=85.7 \%, \mathrm{H}=14.3 \%$. Its molecular weight is 70 . It does not react with $\mathrm{Cl}_{2}$ in dark but gave a substitution product $\mathrm{C}_{5} \mathrm{H}_{9} \mathrm{Cl}$ in presence of light. What is (A)?
Q. 19 An organic compound (A) $\mathrm{C}_{6} \mathrm{H}_{10}$ does not react with ammoniacal $\mathrm{AgNO}_{3}$ but on reduction first gives (B) $\mathrm{C}_{6} \mathrm{H}_{12}$ and then (C) $\mathrm{C}_{6} \mathrm{H}_{14}$. (A) on ozonolysis gives two aldehydes $\mathrm{C}_{2} \mathrm{H}_{4} \mathrm{O}(\mathrm{D})$ and $\mathrm{C}_{2} \mathrm{H}_{2} \mathrm{O}_{2}(\mathrm{E})$. Oxidation of (B) with alkaline $\mathrm{KMnO}_{4}$ gives acid (F) $\mathrm{C}_{3} \mathrm{H}_{6} \mathrm{O}_{2}$. What are (A) of (F)?
Q. 20 Two isomeric organic compounds $(A)$ and $(B)$ on reduction with Zn -Cu couple give n -butane. On treating
(A) and (B) with Na and ether separately, n -octane (C) and 3, 4-diemthylhexane (D) are formed but if a mixture of (A) and (B) reacts with Na and ether, one more product ( E ) in addition to (C) and (D) is

## Exercise 2

## Single Correct Choice Type

Q.1 A sample of 1.79 mg of a compound of molar mass $90 \mathrm{~g} \mathrm{~mol}^{-1}$ when treated with $\mathrm{CH}_{3} \mathrm{MgI}$ releases 1.34 mL of a gas at STP. The number of active hydrogen in the molecule is
(A) 1
(B) 2
(C) 3
(D) 4
Q. 2 Reaction of $\mathrm{Br}_{2}$ on ethylene in presence of NaCl gives
(A) $\mathrm{BrCH}_{2}-\mathrm{CH}_{2} \mathrm{Br}$
(B) $\mathrm{ClCH}_{2}-\mathrm{CH}_{2} \mathrm{Br}$
(C) Both of these
(D) None of these
Q. 3 The reaction conditions leading to the best yield of $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{Cl}$ are
(A) $\mathrm{C}_{2} \mathrm{H}_{6}$ (excess) $+\mathrm{Cl}_{2} \xrightarrow[\text { light }]{\text { UV }}$
(B) $\mathrm{C}_{2} \mathrm{H}_{6}+\mathrm{Cl}_{2} \xrightarrow[\text { room temp. }]{\text { dark }}$
(C) $\mathrm{C}_{2} \mathrm{H}_{6}+\mathrm{Cl}_{2}$ (excess) $\xrightarrow[\text { light }]{\text { UV }}$
(D) $\mathrm{C}_{2} \mathrm{H}_{6}+\mathrm{Cl}_{2} \xrightarrow[\text { light }]{\text { UV }}$
Q. 4 The treatment of $\mathrm{CH}_{3} \mathrm{C}\left(\mathrm{CH}_{3}\right)=\mathrm{CHCH}_{3}$ with $\mathrm{NaIO}_{4}$ or boiling $\mathrm{KMnO}_{4^{\prime}}$ produces
(A) $\mathrm{CH}_{3} \mathrm{COCH}_{3}+\mathrm{CH}_{2} \mathrm{O}$
(B) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CHO}+\mathrm{CH}_{3} \mathrm{CHO}$
(C) $\mathrm{CH}_{3} \mathrm{COCH}_{3}+\mathrm{CO}_{2}$
(D) $\mathrm{CH}_{3} \mathrm{COCH}_{3}+\mathrm{HCOOH}$
Q. 5 What is the chief product obtained when n-butane is treated with bromine in the presence of light at $130^{\circ} \mathrm{C}$ ?
(A)

(B)

(C)

(D)

(A) $\mathrm{CH}_{2}=\mathrm{CHCl}$
(B) $\mathrm{CH}_{2}=\mathrm{CH}_{2}$
(C) $\mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2}$
(D) $\left(\mathrm{CH}_{3}\right)_{2} \mathrm{C}=\mathrm{CH}_{2}$
Q. 7 Which of the following alkene will react fastest with $\mathrm{H}_{2}$ under catalytic hydrogenation condition?
(A)

(B)

(C)

(D)


(A)

(B)

(C)

(D)

Q. 9 The $\mathrm{C}-\mathrm{H}$ bond length is longest in
(A) $\mathrm{C}_{2} \mathrm{H}_{2}$
(B) $\mathrm{C}_{2} \mathrm{H}_{4}$
(C) $\mathrm{C}_{2} \mathrm{H}_{6}$
(D) $\mathrm{C}_{2} \mathrm{H}_{2} \mathrm{Br}_{2}$
Q. 10 Point out (A) in the given reaction sequence:
(A) $\xrightarrow{\mathrm{O}_{3} / \mathrm{H}_{2} \mathrm{O}}(\mathrm{B}) \xrightarrow{\Delta} 2 \mathrm{CH}_{3} \mathrm{COOH}+\mathrm{CO}_{2}$
Q. 11 Hydrogenation of the compound


In the presence of poisoned palladium catalyst gives:
(A) An optically active compound
(B) An optically inactive compound
(C) A racemic mixture
(D) A diastereomeric mixture
Q. 12

(Predominant). A is
(A)

(B)

(C)

(D) Non of these
Q. 13 Identify $(B)$ in the following sequence of reactions $\mathrm{CH} \equiv \mathrm{CH} \xrightarrow{\mathrm{Cu}_{2} \mathrm{Cl}_{2}+\mathrm{NH}_{4} \mathrm{Cl}}(A) \xrightarrow{\mathrm{HCl}}(B)$
(A) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{Cl}$
(B) $\mathrm{CH}_{2}=\mathrm{CH}-\mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{Cl}$

(D) $\mathrm{H}_{2} \mathrm{C}=\stackrel{\text { Cl }}{=}=\mathrm{CH}_{2}$
Q. $14 \mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2} \xrightarrow{\mathrm{Cl}_{2}-\mathrm{H}_{2} \mathrm{O}} \mathrm{Z}$. Z may be

(B)

(C)

(D)

Q. 15

(A)

(B)

(C)

(D) Non of these

(A) $\mathrm{CH}_{3}-\mathrm{CHO}$
(B) $\mathrm{CH}_{3}-\mathrm{COOH}$

(D) None of these
Q. 17 Which one of the following heptanols can be dehydrated to hept-3-ene only?
(A) Heptan-3-ol
(B) Heptan-4-ol
(C) Heptan-2-ol
(D) Heptan-1-ol
Q. 18 Aqueous solution of potassium propanoate is electrolysed. Possible organic products are:
(A) n-Butane
(B) $\mathrm{C}_{2} \mathrm{H}_{5} \mathrm{COOC}_{2} \mathrm{H}_{5}$
(C) $\mathrm{CH}_{3}-\mathrm{CH}_{3}$
(D) $\mathrm{CH}_{2}=\mathrm{CH}_{2}$
Q. $19 \underbrace{\text { N }}_{N+}$ 入 $\bar{O}$

The above compound undergoes elimination on heating to yield which of the following products?
(A)

(B)

(C)

(D)


## Multiple Correct Choice Type


(A)

(B) $\mathrm{CH}_{2}=\mathrm{CHCH}=\mathrm{CH}-\mathrm{CH}_{2} \mathrm{Br}$
(C) $\mathrm{CH}_{2}=\mathrm{CHCH}_{2} \mathrm{CH}=\mathrm{CHBr}$

Q. 21 Which of the following elimination reactions will occur to give but-1-ene as the major product?
(A) $\mathrm{CH}_{3} \cdot \mathrm{CHCl} \cdot \mathrm{CH}_{2} \cdot \mathrm{CH}_{3}+\mathrm{KOH} \xrightarrow{\mathrm{EtOH}}$

(C) $\mathrm{CH}_{3} \cdot \mathrm{CH}_{2} \cdot \mathrm{CHCl}^{2} \cdot \mathrm{CH}_{3}+\mathrm{Me}_{3} \mathrm{CO}^{-} \mathrm{K}^{+} \xrightarrow{\Delta}$
(D) $\mathrm{CH}_{3} \cdot \mathrm{CH}_{2} \cdot \mathrm{CHOH} \cdot \mathrm{CH}_{3}+$ conc. $\mathrm{H}_{2} \mathrm{SO}_{4}$

## Q. 22 Select true statement(s):

(A) $I_{2}$ does not react with ethane at room temperature even though $\mathrm{I}_{2}$ is more easily cleaved homolytically than the other halogens.
(B) Regiochemical outcome of a free radical addition and an electrophilic addition reaction on propene is identical.
(C) The rate of bromination of methane is decreased if HBr is added to the reaction mixture.
(D) Allylic chloride adds halogens faster than the corresponding vinylic chloride.
Q. 23 An alkene on ozonolysis yields only ethanal. There is an isomer of this which on ozonolysis yields:
(A) Propanone
(B) Ethanal
(C) Methanal
(D) Only propanal
Q. $24 \mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}-\mathrm{CH}_{3}+\mathrm{CH}_{2} \mathrm{~N}_{2} \longrightarrow \Delta \mathrm{~A}$

A can be
(A)

(B)

(C)

(D)

Q. 25 Which of the following will give same product with HBr in presence or absence of peroxide.
(A) Cyclohexene
(B) 1-methylcyclohexene
(C) 1, 2-dimethylcyclohexene
(D) 1-butene
Q. 26 The ionic addition of HCl to which of the following compounds will produces a compound having Cl on carbon next to terminal.
(A) $\mathrm{CH}_{3} \cdot(\mathrm{CH} 2)_{3} \cdot \mathrm{CH}=\mathrm{CH}_{2}$
(B) $\mathrm{CH}_{3} \cdot \mathrm{CH}=\mathrm{CH}_{2}$
(C) $\mathrm{CF}_{3} \cdot \mathrm{CH}=\mathrm{CH}_{2}$
(D) $\mathrm{CF}_{3} \cdot \mathrm{CH}=\mathrm{CH}_{2}$
Q. 27 Which is/are true statements/reactions?
(A) $\mathrm{Al}_{4} \mathrm{C}_{3}+\mathrm{H}_{2} \mathrm{O} \longrightarrow \mathrm{CH}_{4}$
(B) $\mathrm{CaC}_{2}+\mathrm{H}_{2} \mathrm{O} \longrightarrow \mathrm{C}_{2} \mathrm{H}_{2}$
(C) $\mathrm{Mg} 2 \mathrm{C}_{3}+\mathrm{H}_{2} \mathrm{O} \longrightarrow \mathrm{CH}_{3} \mathrm{C} \equiv \mathrm{CH}$
(D) $\mathrm{Me}_{3} \mathrm{C}-\mathrm{H}+\mathrm{KMnO}_{4} \longrightarrow \mathrm{Me}_{3} \mathrm{C}-\mathrm{OH}$
Q. $28 \mathrm{C} \xrightarrow[\mathrm{H}_{2} \mathrm{O}_{2} / \mathrm{OH}^{-}]{\mathrm{BH}_{3} / \mathrm{THF}} \mathrm{A} \xrightarrow[\mathrm{H}_{2} \mathrm{SO}_{4}]{\mathrm{HgSO}_{4}} \mathrm{~B}$
$B$ and $C$ are identical when $A$ is -
(A) $\mathrm{H}=\mathrm{H}$
(B) $二 \mathrm{H}$
(С) $二$
(D) $=\mathrm{H}$

## Assertion Reasoning Type

Each of the questions given below consists of two statements, an assertion (A) and reason (R). Select the number corresponding to the appropriate alternative as follows:
(A) If both Assertion and Reason are true and Reason is the correct explanation of Assertion .
(B) If both Assertion and Reason are true but Reason is not the correct explanation of Assertion .
(C) If Assertion is true but Reason is false.
(D) If Assertion is false but Reason is true.
Q. 29 Assertion: In $\alpha, \beta$-unsaturated compounds with $\mathrm{C}=\mathrm{C}$ and $\mathrm{C}=\mathrm{O}$ conjugated, attack of nucleophile takes place on $\mathrm{C}=\mathrm{C}$.

Reason: The $\mathrm{C}=\mathrm{O}$ bond is stronger than $\mathrm{C}=\mathrm{C}$.
Q. 30 Assertion: Alkyl iodides are more reactive than alkyl chlorides for elimination reactions.

Reason: I- is a better leaving group than Cl-
Q. 31 Assertion: $\mathrm{CH}_{3} \mathrm{CH}_{2}^{-}$is a stronger base than $\mathrm{F}-$.

Reason: The negative charge density on carbon is greater than the negative charge density of $\mathrm{F}-$.
Q. 32 Assertion: $\mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3}$ is more reactive than $\mathrm{CH} \equiv \mathrm{CH}$ towards HCl .

Reason: The carbocation formed is more stable in the case of $\mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3}$ than $\mathrm{CH} \equiv \mathrm{CH}$.
Q. 33 Assertion: $\mathrm{CH} \equiv \mathrm{CH}$ reacts with HCl in the presence of $\mathrm{HgCl}_{2}$ while $\mathrm{CH}_{2}=\mathrm{CH}_{2}$ does not.
Reason: There is more unsaturation in $\mathrm{CH} \equiv \mathrm{CH}$ than in $\mathrm{CH}_{2}=\mathrm{CH}_{2}$.

## Comprehension Type (34-36)

n-Butane is produced by mono bromination of ethane followed by Wurtz reaction, as follows
(i) $\mathrm{CH}_{3} \mathrm{CH}_{3} \xrightarrow[h v]{\mathrm{Br}_{2}} \mathrm{CH}_{3} \mathrm{CH}_{2}-\mathrm{Br}$


Mechanism: $\mathrm{Na} \rightarrow \mathrm{Na}^{+}+\mathrm{e}^{-}$;
$\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{Na}+\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{Br} \rightarrow$
$\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{3}+\mathrm{CH}_{2}=\mathrm{CH}_{2}+\mathrm{CH}_{3}-\mathrm{CH}_{3}$
$\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{Br} \rightarrow \mathrm{CH}_{3} \mathrm{CH}_{2}+\mathrm{Br}$
$\mathrm{CH}_{3} \mathrm{CH}_{2}+\mathrm{e}^{-} \rightarrow \mathrm{CH}_{3} \mathrm{CH}_{2} \cdot$
Q. 34 The amount of ethyl bromide required to prepare 55 g of butane would be
(A) 106.72
(B) 206.72
(C) 20.67
(D) 2.067
Q. 35 If the yield of n-butane was 85\%, then the actual amount of ethyl required to produce 55 g of n -butane would be
(A) 2.432 g
(B) 24.32 g
(C) 243.2 g
(D) 0.2432 g
Q. 36 The other products which could be formed
(A) $\mathrm{CH}_{2}=\mathrm{CH}_{2}$
(B) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{3}$
(C) $\mathrm{CH}_{4}$
(D) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{CH}_{3}$

## Comprehension Type (36-38)

The functional group in alkenes is $\mathrm{C}-\mathrm{C}$ double bond which is made of one $\sigma$-bond and one $\pi$-bond. The weaker $\pi$-bond would tend to attract the electrophile and convert itself to a stable carbonium ion which reacts with nucleophile remainder to form the addition product. The addition reaction of alkenes is stereo specific as well as stereoselective. i.e., the isomeric alkenes react differently with same addendum to give stereochemically different products. Where addition of HBr proceeds through carbocation formation and hence racemisation is expected. With low concentration of $\mathrm{Br}_{2}$ in vapour phase substitution predominates over addition.
Q. 37 Reaction of 1,2-dimethylcyclohexene with bromine water gives the product
(A)

(B)

(C)

(D)

Q. $38{ }^{14} \mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2} \xrightarrow[\text { high temperature }]{\mathrm{Br}_{2} \text { /low conc. }}$

Product of the above reaction is
(A) $\mathrm{Br}-{ }^{14} \mathrm{CH}_{2}-\mathrm{CH}=\mathrm{CH}_{2}$
(B) ${ }^{14} \mathrm{CH}_{2}=\mathrm{CH}-\mathrm{CH}_{2}-\mathrm{Br}$
(C) Both (a) and (b)
(D) None of these
Q. 39 Trans-2-butene when reacts with $\mathrm{Br}_{2}$, the product formed is
(A) Racemic mixture
(B) Meso form
(C) Optically pure
(D) No specific stereochemistry

## Match the Columns

Q. 40 Match the entries in column I with entries in column II

| Column I | Column II |
| :--- | :--- |
| (A) $\mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3}$ cis-2-butene | (p) $\mathrm{Na} / \mathrm{NH}_{3}(\mathrm{l})$ |
| $(\mathrm{B}) \mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3} \longrightarrow$ trans-2-butene | (q) $\mathrm{H}_{2} / \mathrm{Pd} / \mathrm{BaSO}_{4}$ |
| $(\mathrm{C}) \mathrm{CH}_{3} \mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3} \longrightarrow$ 1-Butyne | (r) Alc. $\mathrm{KOH}, \Delta$ |
| (D) $\mathrm{CH}_{3}-\mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{CH} \longrightarrow$ 2-Butyne | (s) $\mathrm{NaNH}_{2}, \Delta$ |

Q. 41 Match List-I with List-II and select the correct answer

| List I (Reaction) | List II (Reagents) |
| :--- | :--- |
| (A) $\mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2} \rightarrow \mathrm{CH}_{3}-\mathrm{CHBr}-\mathrm{CH}_{3}$ | (p) HBr |
| (B) $\mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2} \rightarrow \mathrm{CH}_{3}-\mathrm{CH}_{2}-\mathrm{CH}_{2} \mathrm{Br}$ | (q) Br |
| (C) $\mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2} \rightarrow \mathrm{BrCH}_{2}-\mathrm{CH}=\mathrm{CH}_{2}$ | (r) $\mathrm{HBr} /$ Peroxide |
| (D) $\mathrm{CH}_{3}-\mathrm{CH}=\mathrm{CH}_{2} \rightarrow \mathrm{CH}_{3}-\mathrm{CHBr}-$ <br> $\mathrm{CH}_{2} \mathrm{Br}$ | (s) NBS |

Q. 42 Match the column

| Column I | Column II |
| :--- | :--- |
| (A) $\mathrm{RCOONa} \xrightarrow{\text { Electrolysis }} \mathrm{R}-\mathrm{R}$ | (p) Corey-House <br> reaction |
| (B) $\mathrm{R}-\mathrm{CH}_{2}-\mathrm{COOH} \xrightarrow[\Delta]{\text { Soda Lime }}$ <br> $\mathrm{R}-\mathrm{CH}_{3}$ | (q) Kolbe electrolysis |
| (C) $\mathrm{RCOOH} \xrightarrow[\text { (i) } \mathrm{Cl}_{2} / \Delta]{\text { (i) } \mathrm{AgNO}_{3}} \mathrm{R}-\mathrm{Cl}$ | (r) Oakwood <br> degradation |

## Previous Years' Questions

Q. $1\left(\mathrm{CH}_{3}\right)_{3} \mathrm{CMgCl}$ on reaction with $\mathrm{D}_{2} \mathrm{O}$ produces (1997)
(A) $\left(\mathrm{CH}_{3}\right)_{3} \mathrm{CD}$
(B) $\left(\mathrm{CH}_{3}\right)_{3} \mathrm{OD}$
(C) $\left(\mathrm{CD}_{3}\right)_{3} \mathrm{CD}$
(D) $\left(\mathrm{CD}_{3}\right)_{3} \mathrm{OD}$
Q. 2 In the compound, of the type
(1999)
(A) $\mathrm{sp}-\mathrm{sp}^{2}$
(B) $\mathrm{sp}^{3}-\mathrm{sp}^{3}$
(C) $\mathrm{sp}-\mathrm{sp}^{3}$
(D) $\mathrm{sp}^{2}-\mathrm{sp}^{3}$
Q. 3 The product(s) obtained via oxymercuration $\left(\mathrm{HgSO}_{4}\right.$ $+\mathrm{H}_{2} \mathrm{SO}_{4}$ ) of 1-butyne would be
(A) $\mathrm{CH}_{3}-\mathrm{CH}_{2}-\stackrel{\mathrm{O}}{\mathrm{C}}-\mathrm{CH}_{3}$
(B) $\mathrm{CH}_{3}-\mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{CHO}$
(C) $\mathrm{CH}_{3}-\mathrm{CH}_{2}-\mathrm{CHO}+\mathrm{HCHO}$
Q. 4 Identify a reagent from the following list which can easily distinguish between 1-butyne and 2-butyne.
(2002)
(A) Bromine, $\mathrm{CCl}_{4}$
(B) $\mathrm{H}_{2}$, Lindlar catalyst
(C) Dilute $\mathrm{H}_{2} \mathrm{SO}_{4}, \mathrm{HgSO}_{4}$
(D) Ammoniacal $\mathrm{CuCl}_{2}$ solution
Q. 5 Consider the following reaction


Identify the structure of the major product X
(2002)
(A)

(B)

(C)

(D)

Q. $6 \mathrm{Ph}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{3} \xrightarrow[\mathrm{H}^{+}]{\mathrm{Hg}^{2+}} \mathrm{A}$; A is
(2003)
(A)

(B) P

(C)

(D) Ph

## Comprehension Based Questions (Q. 7 \& 8)

An acyclic hydrocarbon $P$, having molecular formula $\mathrm{C}_{6} \mathrm{H}_{10^{\prime}}$ gave acetone as the only organic product
through the following sequence of reactions, in which $Q$ is an intermediate organic compound.
Q. 7 The structure of compound $P$ is
(A) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{2}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}$
(B) $\mathrm{H}_{3} \mathrm{CH}_{2} \mathrm{C}-\mathrm{C} \equiv \mathrm{C}-\mathrm{CH}_{2} \mathrm{CH}_{3}$
(C)

(D)

Q. 8 The structure of the compound Q is


(2011)
(A)

(B)

(C)

(D) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{CH}(\mathrm{OH}) \mathrm{CH}_{2} \mathrm{CH}_{3}$
Q. 9 Give reasons for the following in one or two sentences:
(1983)
(i) Methane does not react with chlorine in dark.
(ii) Propene reacts with HBr to give isopropyl bromide but does not give n-propyl bromide.
Q. 10 Following statements are true, only under some specific conditions. Write the condition for each sub question in not more than two sentences:
(i) 2-methyl propene can be converted into isobutyl bromide by hydrogen bromide
(ii) Ethyne and its derivatives will give white precipitate with ammoniacal silver nitrate solution.
(1984)

## Paragraph for Questions 11 and 12

Schemes 1 and 2 describe sequential transformation of alkynes M and N . Consider only the major products formed in each step for both schemes.


Q. 11 The product $X$ is
(2014)
(A) $\mathrm{H}_{3} \mathrm{CO}$
(C)

(B)

(D)

Q. 12 The correct statement with respect to product Y is
(2014)
(A) It gives a positive Tollens test and is a functional isomer of X .
(B) It gives a positive Tollens test and is a geometrical isomer of $X$.
(C) It gives a positive iodoform test and is a functional isomer of $X$.
(D) It gives a positive iodoform test and is a geometrical isomer of X .

# PlancEssential Questions 

## JEE Main/Boards

## Exercise 1

$\begin{array}{llll}\text { Q. } 3 & \mathrm{Q} .8 & \mathrm{Q} .13 & \mathrm{Q} .15 \text { (b) } \\ \text { Q. } 20 \text { (a) } & & & \end{array}$

## Exercise 2

Q. 1
Q. 7
Q. 9
Q. 14
Q. 28

Previous Years' Questions
Q. 10
Q. 13

## JEE Advanced/Boards

## Exercise 1

Q. 1
Q. 8
Q. 13
Q. 18
Q. 21

## Exercise 2

Q. 5
Q. 9
Q. 14
Q. 22
Q. 26

## Previous Years' Questions

Q. $6 \quad$ Q. 8

## Answer Key

## JEE Main/Boards

## Exercise 2

Single Correct Choice Type
Q. 1 B
Q. 2 C
Q. 3 A
Q. 4 D
Q. 5 C
Q. 6 C
Q. 7 B
Q. 8 D
Q. 9 D
Q. 10 B
Q. 11 A

## Previous Years' Questions

| Q. 1 B | Q.2.D | Q. 3 D | Q. 4 C | Q. 5 A | Q. 6 A | Q 7 B |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Q. 8 B | Q. 9 C | Q. 10 A | Q. 11 A | Q. 12 D | Q. 13 C | Q. 14 D |
| Q. 15 B | Q. 16 D | Q. 17 C | Q. 18 C | Q. 19 A | Q. 20 B | Q. 21 A |

Q. 22 B

## JEE Advanced/Boards

## Exercise 2

Single Correct Choice Type

| Q. 1 C | Q. 2 C | Q. 3 A | Q. 4 B | Q. 5 B, C | Q. 6 B | Q. 7 D |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Q. 8 B | Q. 9 C | Q. 10 C | Q. 11 B | Q. 12 C | Q. 13 D | Q. 14 D |
| Q. 15 C | Q. 16 A | Q. 17 B | Q. 18 A | Q. 19 B |  |  |

Multiple Correct Choice Type
Q. 20 A, B
Q. 21 B, C
Q. 22 A, C, D
Q. 23 A, C
Q. 24 A, B, C, D
Q. 25 A, C
Q. 26 A,B,D
Q. 27 A,B,C,D Q. 28 A,C

Assertion Reasoning Type
Q. 29 B
Q. 30 A
Q. 31 A
Q. 32 A
Q. 33 B

Comprehension Type
Q. 34 A
Q. 35 A
Q. 36 A
Q. 37 D
Q. 38 C
Q. 39 B

Match the Columns
Q. $40 \mathrm{~A} \rightarrow \mathrm{q} ; \mathrm{B} \rightarrow \mathrm{p} ; \mathrm{C} \rightarrow \mathrm{s} ; \mathrm{D} \rightarrow \mathrm{r}$
Q. $41 \mathrm{~A} \rightarrow \mathrm{p} ; \mathrm{B} \rightarrow \mathrm{r} ; \mathrm{C} \rightarrow \mathrm{s} ; \mathrm{D} \rightarrow \mathrm{q}$
Q. $42 \mathrm{~A} \rightarrow \mathrm{q} ; \mathrm{B} \rightarrow \mathrm{r} ; \mathrm{C} \rightarrow \mathrm{s} ; \mathrm{D} \rightarrow \mathrm{p}$

## Previous Years Questions

Q. 1 A
Q. 2 D
Q. 3 A
Q. 4 D
Q. 5 B
Q. 6 A
Q. 7 D
Q. 8 B
Q. 11 A
Q. 12 C

## Solutions

## JEE Main/Boards

## Exercise 1

Sol 1: Alkenes are substances which contain less hydrogen than the maximum quantity of hydrogen which a carbon atom can have

## Structure:-



Preparation of Alkenes:-
(1) Dehydrohalogenation of Alkyl halide
(2) Dehyration of Alcohols
(3) Dehalogenation of vicinal dihalides
(4) Dehalogenation of geminal dihalide

Sol 2: (A)
 trans additon
$\therefore$ Recemic mixture
(B)


Antimarkov's addition

Sol 3: (A)

(B)

(C)

(D)


## Sol 4

(A)

(B)


## Sol

5: Physical properties: Refer theory pg. 26
Chemical properties: Refer theory pg. 29

## Sol 6: Alkynes:-

In the family of hydrocarbon alkynes are the most unsaturated one with $\mathrm{C} \equiv \mathrm{C}$ bond and are highly reactive.

## Preparation:-

(1) Dehydrohalogenation
(2) From tetradehalogenation
(3) From haloform
(4) Kolbe's electrolytic decarboxylation
(5) Berthlot's reaction

Sol 7: Physical properties: Refer theory pg. 42
Chemical properties: Refer theory pg. 44

## Sol 8:

(A)


(B)


Sol 9:
(A)

(B)


Sol 10: (A)

(B)

(C)


Doubled bonds are more electron donating than triple bond because of difference in electronegativity and because of which first addition will takes place on double bond rather than triple bond.
(B)


Sol 12:



Sol 13:

$\downarrow \downarrow 3^{\circ} \mathrm{ROH}$


(I)




Sol 14

(X)

$\underbrace{\mathrm{Cl}}_{\text {(1) } \mathrm{CH}_{3} \mathrm{CO}} \mathrm{H}^{+} / \mathrm{H}_{2} \mathrm{O}$


## Exercise 2

## Single Correct Choice Type

Sol 1: (B) Dicarboxylation of acid ion


Sol 2: (C)


Sol 3: (A) Definition of conformers.

Sol 4: (D)


Sol 5: (C) $C \equiv C$ is strongest because it is shortest and more orbitals are overlapping.

Sol 6: (C)


Sol 7: (B) $\mathrm{C}_{2} \mathrm{H}_{3} \mathrm{MgI} \xrightarrow{\mathrm{H}_{2} \mathrm{O}} \mathrm{C}_{2} \mathrm{H}_{6}$
Abstracts acidic hydrogen.

Sol 8: (D)


Sol 9: (D)


Anti Markonikov's addition
Sol 10: (B)


Sol 11: (A) Enol form is not net stable as keto is much more stable.

## Previous Years' Questions

Sol 1: (B) Methane is produced due to the decay of vegetables or animal organisms present in swamps and marsh, by the action of bacteria. Due to this method of formation, methane is also known as marsh gas.

Sol 2: (D) Unsaturated compounds which contain $C=C$ or $C \equiv C$, decolourises the purple colour of alkaline $\mathrm{KMnO}_{4}$ solution.


Sol 3: (D) Structural formula of 1, 2-butadiene is:


Sol 4: (C) Alkynes undergo Markownikoff's addition of water in presence of $\mathrm{H}_{2} \mathrm{SO}_{4} / \mathrm{HgSO}_{4}$ :


Sol 5: (A) Bayer's reagent is cold, dilute, alkaline perman-ganate solution, used to detect presence of olefinic bonds.

Sol 6: (A) Terminal alkynes are slightly acidic, forms salt with very strong base like $\mathrm{Na}, \mathrm{NaNH}_{2}$ etc.

$$
\mathrm{H}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}+\mathrm{Na} \xrightarrow{\square} \mathrm{C} \text { ethyne } \mathrm{H}-\mathrm{C} \equiv \mathrm{C}^{-} \mathrm{Na}^{+}+\frac{1}{2} \mathrm{H}_{2} \uparrow
$$

Sol 7: (B) Hydrogenation with poisoned palladium brings about cis hydrogenation of alkyne and does not affect double bonds:

Sol 8: (B)

i.e. Reaction is initiated by $\mathrm{Cl}^{+}($Chloronium ion electrophile).

Sol 9: (C) In addition of HBr to an alkene, in presence of peroxide, both the propagation steps are exothermic:
$\mathrm{HBr}+\mathrm{HO}^{\bullet} \rightarrow \mathrm{H}_{2} \mathrm{O}+\mathrm{Br}^{\bullet}$
Propagation


In case of addition of HCl and HI , one of the propagation step is endothermic, reaction fail to occur.

## Sol 10: (A)

$\mathrm{C}_{6} \mathrm{H}_{5} \mathrm{CH}_{2}=\mathrm{CHCH}_{3}$ has two geometrical isomers


So it is 1-Phenyl-2-butene
Sol.11: (D) $\mathrm{CH}_{3}-\mathrm{CH}^{+}-\mathrm{CH}_{2}-\mathrm{Cl}$

## Sol 12: (A)



## 1, 1, 1- Trichloroethane

## Sol 13: (C)

Sol 14: (D)


Sol 15: (B)


It is a terminal alkyne, having acidic hydrogen.
Note: Solve it as a case of terminal alkynes, otherwise all alkynes react with Na in liq. $\mathrm{NH}_{3}$

Sol 16: (D) $\mathrm{CH}_{3}-\mathrm{MgX}+\mathrm{CH}_{3}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H} \rightarrow \mathrm{CH}_{4}$

Sol 17: (C)


Cis


Trans

Sol 18: (C) 2-butene is symmetrical alkene


Molar mass of $\mathrm{CH}_{3} \mathrm{CHO}$ is 44 u .

Sol 19: (A)


Therefore, Statement I is correct but Statement II is incorrect.

Sol 20: (B) Both Statement I and Statement II are correct and Statement II is correct explanation of Statement I.

## Sol 21: (A)



Both Statement I and Statement II are correct and Statement II is correct explanation of Statement I.

Sol 22: (B) Statement I is correct. Statement II is also correct. Meso form of the product is due to anti addition of Bron cyclic bromonium ion intermediate, hence Statement II is not correct explanation of Statement II.


## JEE Advanced/Boards

## Exercise 1

Sol 1: $(X) \longrightarrow \not \longrightarrow$ Lucas reagent
$\mathrm{C}_{5} \mathrm{H}_{8} \mathrm{O}$


Degree of unsaturation: 2



Sol 2: (A)
 (B) $+(C)$

(B)

(C) $\rightarrow \mathrm{C}_{2} \mathrm{H}_{5}-\stackrel{\text { II }}{\mathrm{C}}$
$(\mathrm{A}) \rightarrow \sim$

Sol 3:

(B)


(C) terminal alkyne
$(D)+(E)$


Sol 4:


$$
+\left(\mathrm{CH}_{3}-\stackrel{\mathrm{O}}{\mathrm{C}}-\mathrm{O}^{-}\right)_{2} \mathrm{Ca}^{2+}
$$

Sol 5:


Sol 6:

$$
\begin{gathered}
\text { (A) } \xrightarrow[\mathrm{C}_{9} \mathrm{H}_{12}]{\text { KinO }} \xrightarrow{\text { alk. }} \underset{\mathrm{C}_{8} \mathrm{H}_{6} \mathrm{O}_{4}}{(\mathrm{BM}) \xrightarrow{\Delta}} \begin{array}{l}
\text { does not } \\
\text { form anhydride }
\end{array} \\
\text { Fe } \mathrm{FBr}_{2} \\
\text { only one monobromo } \\
\text { derivative }
\end{gathered}
$$

Degree of unsaturation of $(A)=4$
So it should contain benzene ring
In the compound (B) both -COOH should be at para position to give only one bromination product
(A)

(B)
(C)


(A)

(B)

(C)


Sol 8:

$\mathrm{C}_{9} \mathrm{H}_{10} \mathrm{Br}_{2}$

(C)

Ozonolysis


Since only one -Br is hydrolysed
$\therefore$ One is attached to alkyl group and other one to aryl group.
(A)





Sol 9:

(D) + (E) Aldehydes
$2 \mathrm{CH}_{2} \mathrm{OC}_{2} \mathrm{H}_{2} \mathrm{O}_{2}$


So (A) is

$(B)$ is 1-4 addition
(C)


(F)


Sol 10:

$\mathrm{C}_{5} \mathrm{H}_{10} \mathrm{O}$ (C)
Ketone
(D) $+(E) \mathrm{C}_{4} \mathrm{H}_{8}$ isomeric acids

Degree of unsaturation of $(A)=2$
$\therefore$ Alkyne. Terminal alkyne because formation of (B) and branched because formation of ketone (C)
(A)

(C)

(D), (E)


Sol 11:


For reaction $(B) \rightarrow(C)$ it is iodoform so (D) $\rightarrow \mathrm{CHI}_{3}$ yellow precipitate and $(\mathrm{B})$ should have acetyl ketone.
(B)


(C)

(A)


OMDM for alkyne give ketone

Sol 12: $\quad(A) \xrightarrow{\text { Hydrogenation }}(B)$

(A)



Sol 13:





Compound (A) carbocation is more stable resonance is in both the compounds but
(A) have $3 \alpha-H$
(B) have $0 \alpha-H$

Sol 14:

(B) $\xrightarrow[\mathrm{KOH}]{\text { Alc. }}$
(D) $+(E)$
(C), (D), (E)

$\downarrow$| Catalytic |
| :--- |
| hydrogenation |

$\mathrm{C}_{5} \mathrm{H}_{12}$
$\therefore$ (D) is isomer of (C)
$\therefore(\mathrm{D}) \longrightarrow$

(B)

(D)


Sol 15:
「_symmetric compound

Q is an acid because after reaction with $\mathrm{CO}_{2}, \mathrm{H}^{+}$of RMgBr we get a carboxylic acid. If

$\therefore(P)$

to satisfy the degree of unsaturation of (P) alkyl group must be cyclic.

Sol 16: (A), (B) $\mathrm{C}_{5} \mathrm{H}_{11} \mathrm{Br}$
isomeric alkyl bromides $(A) \xrightarrow[K O H]{\text { alc. }}(C)+(D)$ isomers $\mathrm{C}_{5} \mathrm{H}_{10}$

Sol 17:
(i)

(ii)

(iii)

(iv)


Anti-addition
(v)

(vi)


Sol 18: (A) $\xrightarrow[h v]{\mathrm{Cl}_{2}} \mathrm{C}_{5} \mathrm{H}_{9} \mathrm{Cl}$
C $=85.7 \% ; h=14.3 \%$
total molecular weight $=70$


Sol 19:


(A)

(B)

(C)

(F)


Sol 20: (A) + (B)
isomers

$(\mathrm{A}) \xrightarrow{\mathrm{Na} / \text { Ether }}$
(C)
(B)



But mixture gives ( E )
Na /Ether is Wurtz reaction and dimerise the alkyl bromide so they are symmetric.
(A)

(B)



## Exercise 2

## Single Correct Choice Type

Sol 1: (C) Releases 1.34 mL of a gas at STP
$\therefore 3$ moles of $\mathrm{CH}_{4}$
$\therefore 3$ active $\alpha-\mathrm{H}$ hydrogen.
Sol 2: (C) $\mathrm{CH}_{2}=\mathrm{CH}_{2} \xrightarrow[\mathrm{NaCl}]{\mathrm{Nr}_{2}} \mathrm{Br}-\mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{Br}$


Sol 3: (A) $\underset{\substack{2 \\(\times 5)}}{\mathrm{C}_{6}}+\mathrm{Cl}_{2} \xrightarrow{h \nu} \mathrm{C}_{2} \mathrm{H}_{5} \mathrm{Cl}$

Sol 4: (B)


Sol 5: (B,C)


Sol 6: (B) Carbon is least stable in $\begin{array}{r}\mathrm{CH}_{3}-\stackrel{\oplus}{\mathrm{CH}} \mathrm{H}_{2} \\ \mathrm{CH}_{2}=\mathrm{CH}_{2}\end{array}$
$\therefore$ Addition will be least $\mathrm{CH}_{2}=\mathrm{CH}_{2}$
Sol 7: (D) Stability of complex that forms during reaction with $\mathrm{H}_{2}$


Sol 8: (B)


Sol 9: (C) C-C-H is sp ${ }^{3}$ hybridised, hence ' $s$ ' character is less, So bond length in C-H is longest

Sol 10: (C)


Sol 11: (B) Hydrogenate product

$\therefore$ optically inactive compound.

Sol 12: (C)



Sol 13: (D)


Sol 14: (D)


Sol 15: (C)


Sol 16: (A)
-

Sol 17: (B)


Sol 18: (A)


Sol 19: (B)


## Multiple Correct Choice Type

Sol 20: $(A, B)$


Sol 21: (B,C)
(A)

(B)

(C)

(D)


Sol 22: (A,C,D) but the reaction will be reversible since C-I is very week.
(B) $\times$
(C) $\checkmark$
(D) more stable system

Sol 23: (A,C)


## Sol 24: (A,B,C,D)



Same for trans form.

Sol 25: (A,C)
(A)

(B)
 absence

(C)

(D)


Sol 26: (A,B,D)
(A) $\mathrm{CF}_{3}-\mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{CH}_{2}-\mathrm{CH}=\mathrm{CH}_{2}$

(B)

(C)

(D)



Sol 27: (A,B,C,D)
(A) $\mathrm{Al}_{4} \mathrm{C}_{3} \xrightarrow{\mathrm{H}_{2} \mathrm{O}} \mathrm{Al}(\mathrm{OH})_{3}+3 \mathrm{CH}_{4} 4 \mathrm{Al}^{3+}+3 \mathrm{C}^{4-}$
(B) $\mathrm{Ca}^{2+}{ }^{-} \mathrm{C} \equiv \mathrm{C}^{-}+\mathrm{H}_{2} \mathrm{O} \longrightarrow \mathrm{Ca}(\mathrm{OH})_{2}+\mathrm{HC} \equiv \mathrm{CH}$
(C) $2 \mathrm{Mg}^{2+}+\mathrm{C}_{3}^{4-}+\mathrm{H}_{2} \mathrm{O} \longrightarrow \mathrm{Mg}(\mathrm{OH})_{2}+\mathrm{H}_{3} \mathrm{C}-\mathrm{C} \equiv \mathrm{CH}$
(D) $\mathrm{Me}_{3} \mathrm{C}-\mathrm{H}+\mathrm{KMnO}_{4} \longrightarrow \mathrm{Me}_{3} \mathrm{C}-\mathrm{OH}$

Sol 28: (A,C)
(A)

(B)

(C)

(D)


## Assertion Reasoning Type

Sol 29: (B) When the $C=C$ bond is conjugated with a $\mathrm{C}=\mathrm{O}$ group, the electron withdrawing nature of carbonyl group polarizes the $C=C$ bond so the nucleophilic attack can take place at the $C=C$ carbon.

Sol 30: (A) Alkyl iodides are more reactive than alkyl chlorides for elimination reactions as I- is a better leaving group than $\mathrm{Cl}-$

Sol 31: (A) is a stronger base than F-because the negative charge density on carbon is greater than the negative charge density of F -.

Sol 32: (A) CH3-C $=\mathrm{C}-\mathrm{CH} 3$ is more reactive than
$\mathrm{CH} \equiv \mathrm{CH}$ towards HCl because the carbocation formed is more stable in thecase of $\mathrm{CH} 3 \mathrm{C} \equiv \mathrm{C}-\mathrm{CH} 3$ than $\mathrm{CH} \equiv \mathrm{CH}$.

Sol 33: (B) Alkynes are electron-rich nucleophiles with a cylindrical electron cloud formed of two bonds around a carbon-carbon bond. An electrophilic reagent can therefore easily react with the relatively weak alkyne bond.

## Comprehension Type

Sol 34: (A)

$\frac{m}{208}=\frac{55}{58}$

Sol 35: (A) 2.432 g

Sol 36: (A) Alkenes generally forms by elimination.

Sol 37: (D)
 Anti addition


Sol 38: (C)


Sol 39: (B)


(B) Anti Addition

## Match the Columns

## Sol 40:

(A) $=\longrightarrow \longrightarrow$
cis addition
$\mathrm{H}_{2} / \mathrm{Pd} / \mathrm{BaSO}_{4}$
$(\mathrm{B}) \longrightarrow \underset{\begin{array}{l}\text { Birch } \\ \text { reduction }\end{array}}{ } \longrightarrow$
(C) $\longrightarrow \underset{\mathrm{NaNH}_{2, \Delta}}{ } \equiv$

Isomerism
(D) $\rightleftharpoons \underset{\text { Alc. } \mathrm{KOH}, \Delta}{ }=$ Isomerism
$A \rightarrow q ; B \rightarrow p ; C \rightarrow s ; D \rightarrow r$

Sol 41: $(A) \rightarrow(p)$ Markonikov
(B) $\rightarrow$ (r) Anti-markonikov
$(C) \rightarrow(s)$ Alkylic carbon get brominated
(D) $\rightarrow$ (q) $\mathrm{Br}_{2}$ addition

Sol 42: $(A) \rightarrow(q)$ Free radical mechanism
$(B) \rightarrow(r)$ Decarboxylation
$(C) \rightarrow(s)$ Free radical mechanism
$(D) \rightarrow(p)$ Same as grignard

## Previous Years' Questions

Sol 1: (A)


Sol 2: (D) According to the IUPAC conventions, compound can be numbered as:


Here, $\mathrm{C}-2$ is $s p^{2}$ and $\mathrm{C}-3$ is $s p^{3}$ hybridised.

Sol 3: (A) Oxymercuration - demercuration brings about Markownikoff's addition of water as


Sol 4: (B) Ammonical $\mathrm{CuCl}_{2}$ forms red precipitate with terminal alkynes, can be used to distinguish terminal alkynes from internal alkynes:


Sol 5: (B) Bromination is highly selective, occur at the carbon where the most stable free radical is formed:


Sol 6: (A) Reaction proceeds through carbocation intermediate:


Sol 7, 8: (D, B) The final ozonolysis product indicates that the alkene before ozonolysis is


Also $\mathrm{P}\left(\mathrm{C}_{6} \mathrm{H}_{10}\right)$ has two degree of unsaturation and oxymercuration demercuration hydration indicates that it is an alkyne. As alkyne, on hydration, gives a carbonyl compound which on reduction with $\mathrm{NaBH}_{4}$ gives a $2^{\circ}$ alcohol.


The secondary alcohol that can give above shown alkene on acid catalyzed dehydration is



Sol 9: (i) Free radical chorination of alkane require energy which is supplied either in the form of heat or radiation.

(ii) Addition of HBr proceeds through carbocation intermediates.

Sol 10: (i)


In the absence of peroxide, HBr would be added giving tertiary butyl bromide.
(ii) Tertiary alkynes are slightly acidic, forms silver salt with ammoniacal solution of silver nitrate:
$\mathrm{R}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}+\mathrm{AgNO}_{3} \xrightarrow{\mathrm{NH}_{3}(\mathrm{aq})} \mathrm{R}-\underset{\text { white ppt. }}{\mathrm{C}} \equiv \mathrm{C} \downarrow$

Solution for the Q. No. 11 and 12: (A, C)




