



تفريغ عضويه 1

موضوع المحاضرة: aldehydes and ketones

رقم المحاضرة : 1- Final

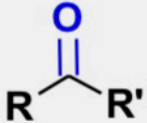
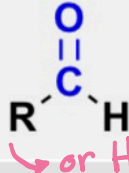
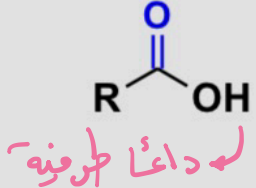
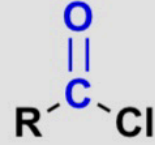
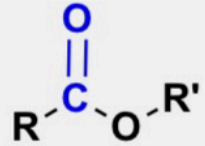
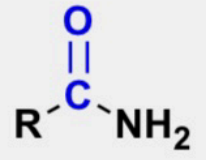
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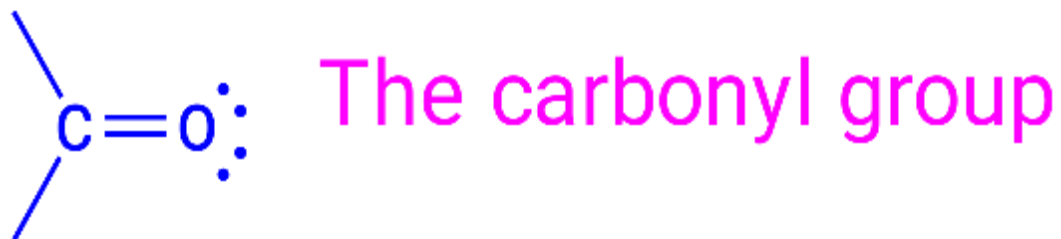
Pharmaceutical Organic Chemistry-1

Chapter-5: Aldehydes & Ketones

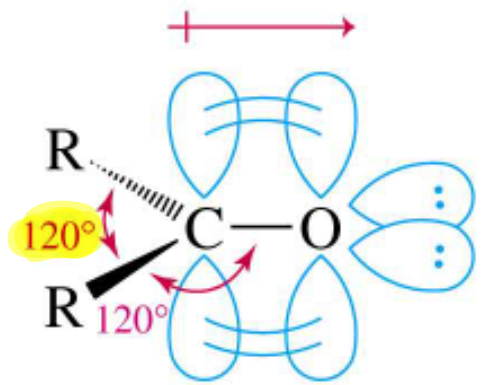
Aldehydes & Ketones

Common Classes of Carbonyl Compounds			
Class	General Formula	Class	General Formula
Ketones		Aldehydes	
Carboxylic acids		Acid Chlorides <i>the most used halide: Cl because good leaving group</i>	
Esters		Amides	

Aldehydes & Ketones

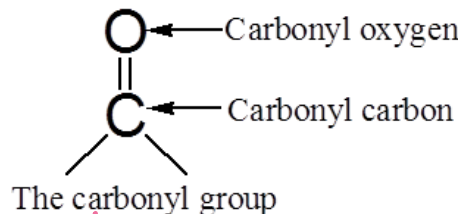


- o Carbon is **sp^2 hybridized**.
- o **C=O bond** is shorter, stronger, and more polar than C=C bond in alkenes.

	<i>length</i>	<i>energy</i>
	ketone C=O bond	1.23 Å 178 kcal/mol (745 kJ/mol)
	alkene C=C bond	1.34 Å 146 kcal/mol (611 kJ/mol)

Structure of Aldehydes and Ketones

- Aldehydes and ketones are characterized by the presence of the carbonyl group.

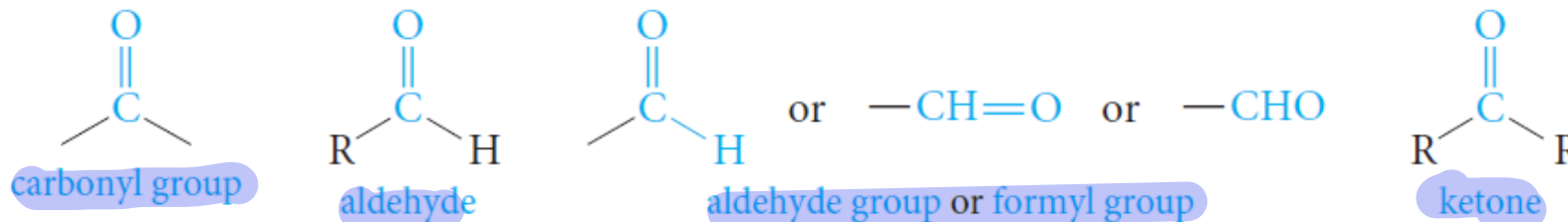


- Aldehydes have at least one hydrogen atom attached to the carbonyl carbon atom.

The remaining group may be another hydrogen atom or any aliphatic or aromatic organic group.

The -CH=O group characteristic of aldehydes is often called a formyl group.

- In ketones, the carbonyl carbon atom is connected to two other carbon atoms.

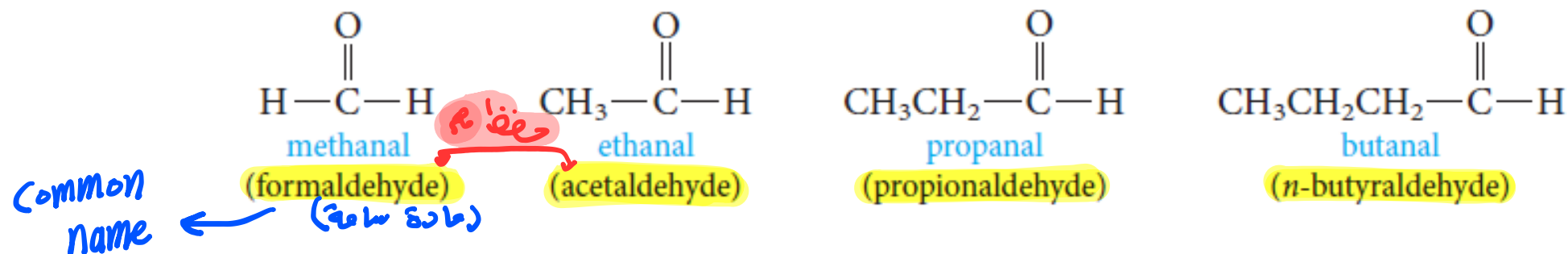


Nomenclature of Aldehydes

IUPAC System

O Aliphatic aldehydes are named by dropping the suffix -e from the name of the hydrocarbon that has the same carbon skeleton as the aldehyde and replacing it with the suffix -al.

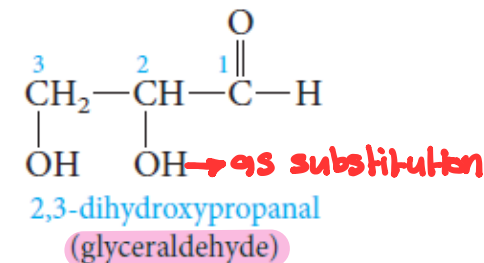
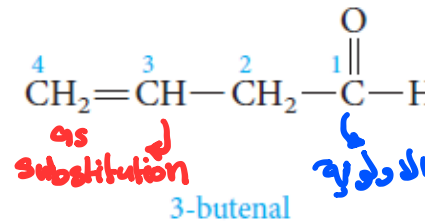
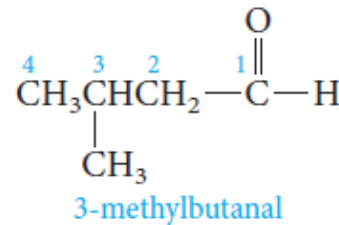
Alkane - e + al = Alkanal



IUPAC System

O Substituted aldehydes, we number the chain starting with the aldehyde carbon.

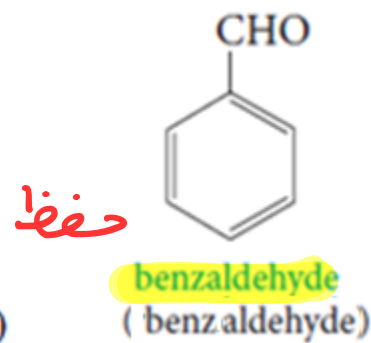
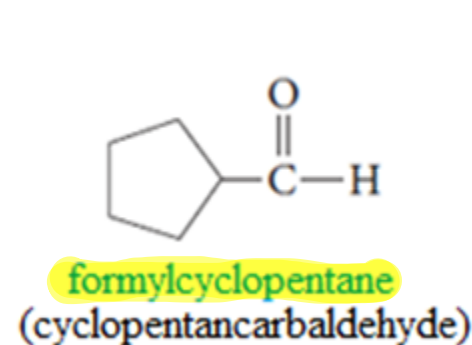
- $-CH=O$ group is assigned the number 1 position.
- Aldehyde group has priority over a double bond or hydroxyl group.



O Cyclic aldehydes, the suffix **-carbaldehyde** is used.

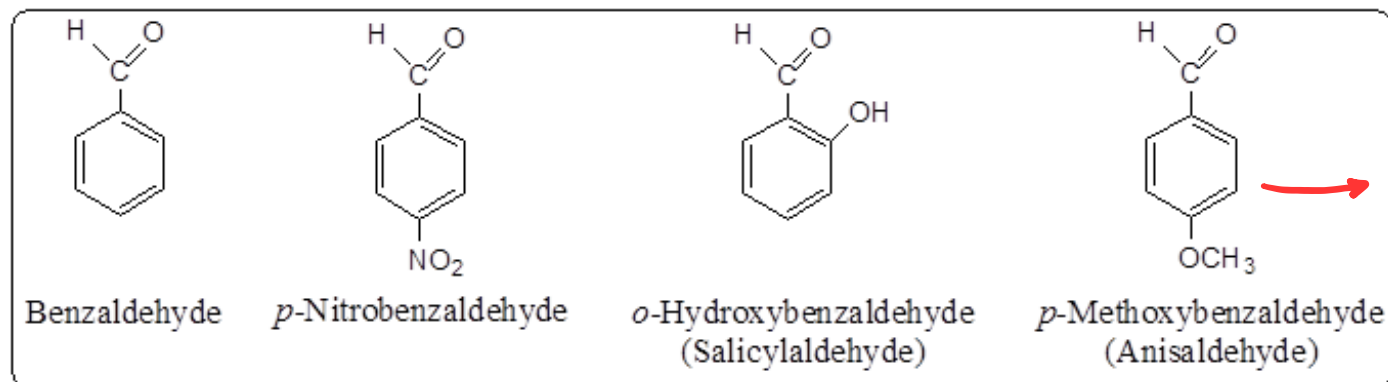
as substitution يكون
وما يكون جزء من ال ring

Common
(IUPAC)



IUPAC System

O **Aromatic aldehydes** are usually designated as derivatives of the simplest aromatic aldehyde, **benzaldehyde**.

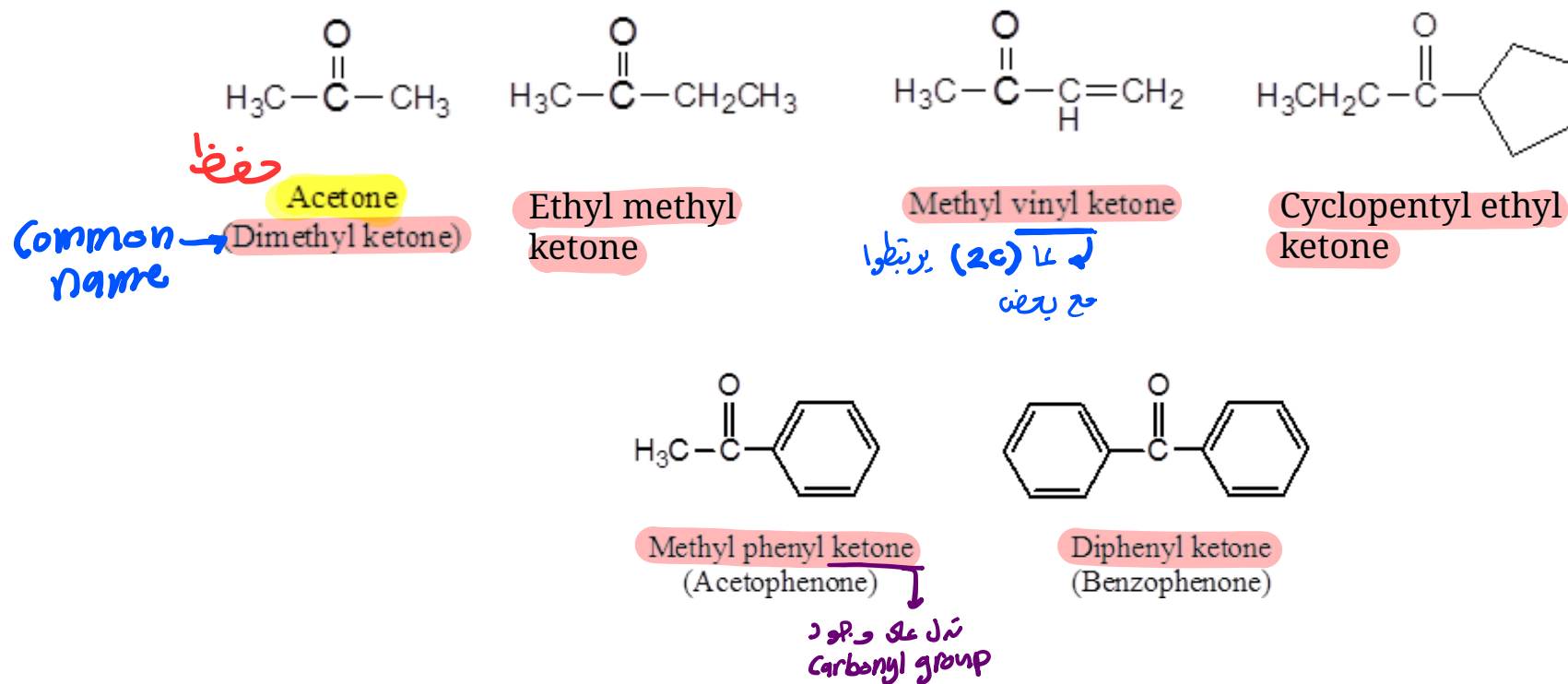


→ ester + aldehyde

Nomenclature of Ketone

Common Names

- Common names of ketones are formed by adding the word *ketone* to the names of the alkyl or aryl groups attached to the carbonyl carbon. **Alkyl ketone.**
- In still other cases, traditional names are used.

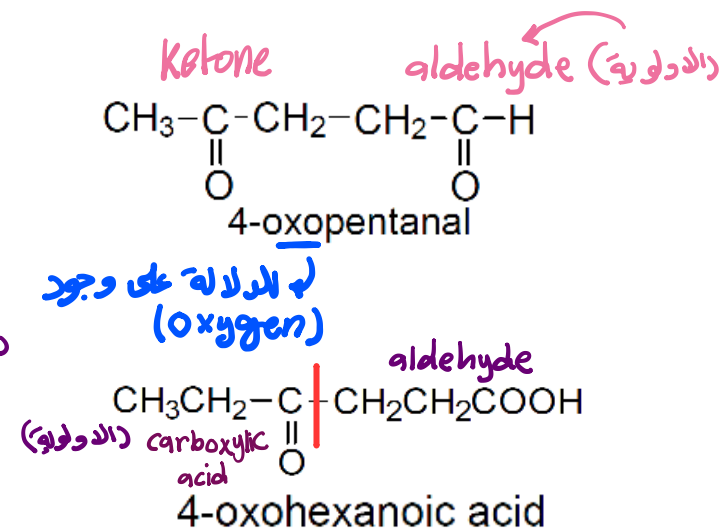
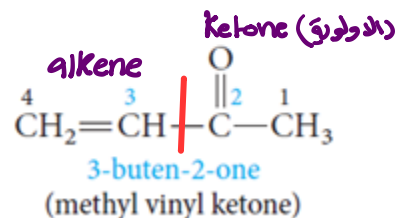
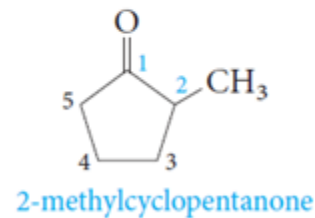
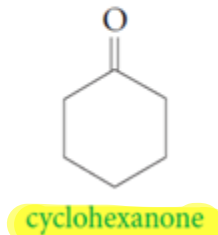
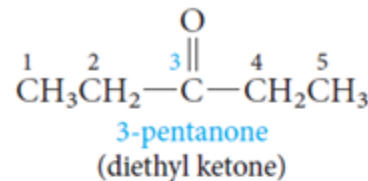
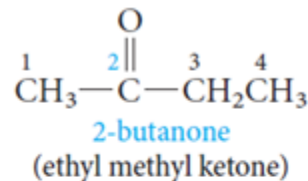
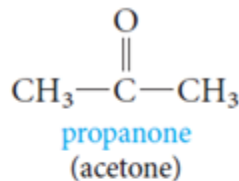


Nomenclature of Aldehydes

IUPAC System

- In the IUPAC system, the ending for ketones is **-one**.
- The chain is numbered so that the **carbonyl carbon has the lowest possible number**.
- For **cyclic ketones**, numbering always starts from the C=O group.
- The prefix "**oxo**" is used when the ketone is not the principal functional group.

IUPAC
(Common)

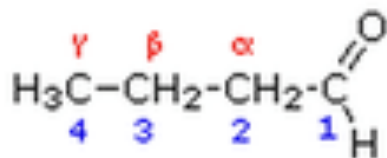


Nomenclature of Aldehydes Ketones

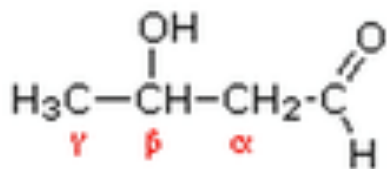
NOTES

- In common names carbon atoms near the carbonyl group are often designated by Greek letters.
- The atom adjacent to the function is *alpha* (α), the next removed is *beta* (β) and so on. Since ketones have two sets of neighboring atoms, one set is labeled α , β etc., and the other α' , β' etc.

Aldehydes

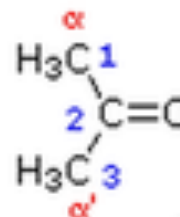


butanal
butyraldehyde

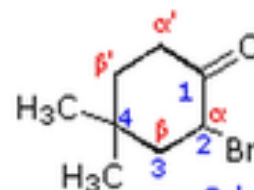


3-hydroxybutanal
 β -hydroxybutyraldehyde
or **aldol**
aldehyde $\leftarrow$ $\rightarrow$ alcohol

Ketones



propanone
acetone



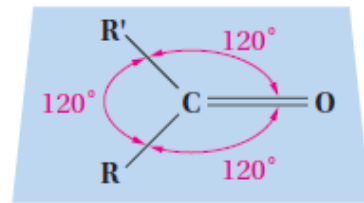
2-bromo-4,4-dimethylcyclohexanone

- The functional group priority order in nomenclature system is as following:
Acid and derivatives > aldehyde > ketone > alcohol > amine > alkene > alkyne > ether

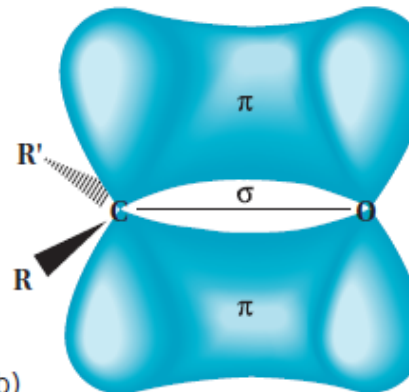
The Carbonyl Group

O The structure and properties of the carbonyl group.

- The carbon–oxygen **double bond** consists of a sigma bond and a pi bond.
- The carbon atom is **sp^2** -hybridized. The three atoms attached to the carbonyl carbon lie in a plane with bond angles of **120°** .
- The **pi bond** is **formed** by **overlap of a p orbital on carbon with an oxygen p orbital**.
- There are also **two unshared electron pairs** on the oxygen atom.
- The C=O bond distance is **1.24Å**, shorter than the **C-O** distance in alcohols and ethers (**1.43Å**) .



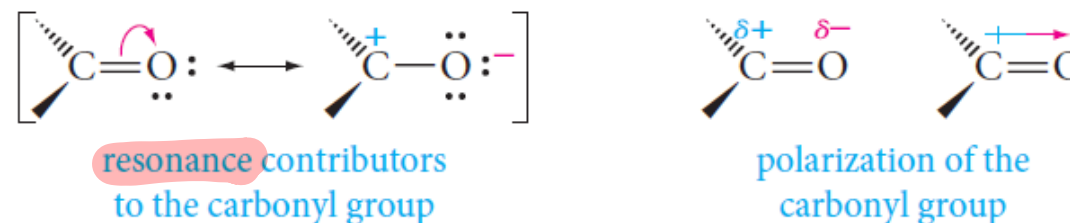
(a)



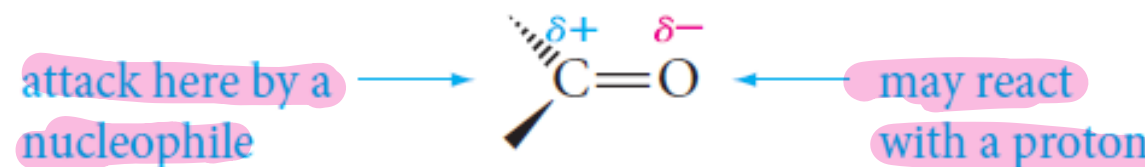
(b)

The Carbonyl Group

- Oxygen is much more electronegative than carbon. Therefore, the electrons in the C=O bond are attracted to the oxygen, producing a highly **polarized bond**.



- As a consequence of this polarization, *most carbonyl reactions involve nucleophilic attack at the carbonyl carbon, often accompanied by addition of a proton to the oxygen (electron rich).*



Physical Properties of Aldehydes and Ketones

Boiling Points

alcohols > Carbonyl > hydrocarbons

- Carbonyl compounds boil at higher temperatures than hydrocarbons, but at lower temperatures than alcohols of comparable molecular weight.

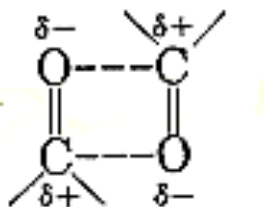
$\text{CH}_3(\text{CH}_2)_4\text{CH}_3$
hexane (bp 69°C)

$\text{CH}_3(\text{CH}_2)_3\text{CH}=\text{O}$
pentanal (bp 102°C)

$\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{OH}$
pentanol (bp 118°C)

justify ↘

- This is due to the intermolecular forces of attraction, called dipole-dipole interactions, which is stronger than van der Waals attractions but not as strong as hydrogen bonds.

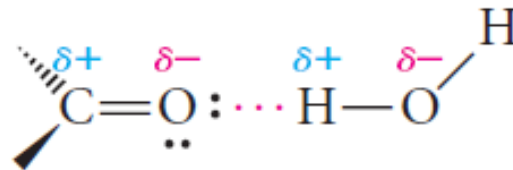


Dipole-dipole attractions among carbonyl compounds

Physical Properties of Aldehydes and Ketones

Solubility

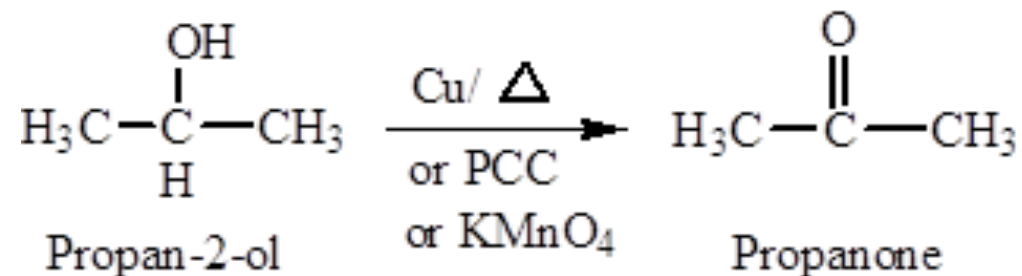
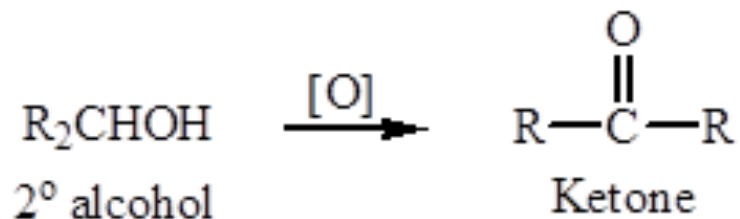
- 0 Carbonyl compounds as aldehydes and ketones have a C=O bond, but no O-H bond, cannot form hydrogen bonds with themselves.
- 0 The polarity of the carbonyl group also affects the solubility properties of aldehydes and ketones.
- 0 Carbonyl compounds with low molecular weights are soluble in water as they can form hydrogen bonds with O-H or N-H compounds.



Preparation of Aldehydes and Ketones

1) Oxidation of Primary and Secondary Alcohols

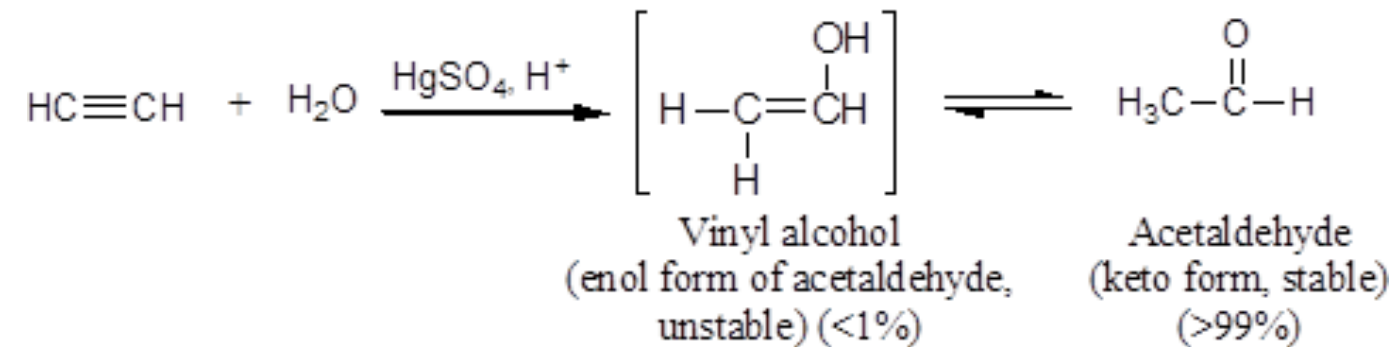
- o Oxidation of secondary alcohols yields ketones.



Preparation of Aldehydes and Ketones

2) Hydration of Alkynes

- Hydration of **acetylene** yields acetaldehyde (catalyzed by acid and mercuric).

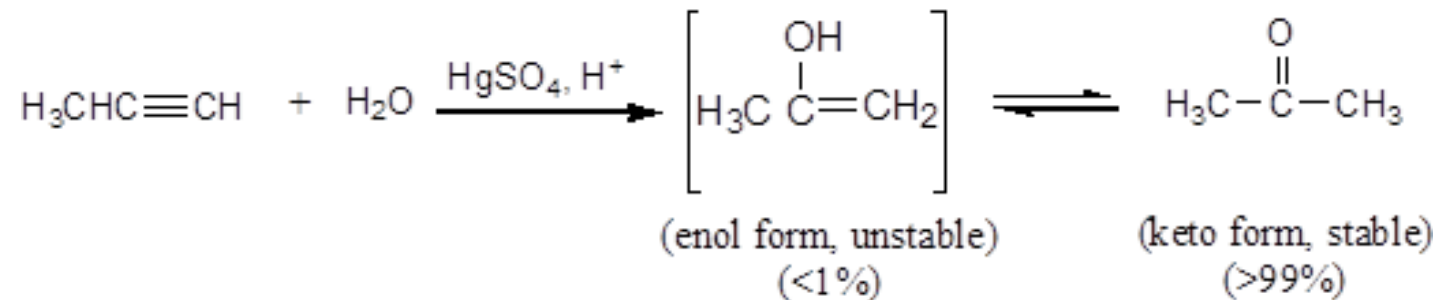
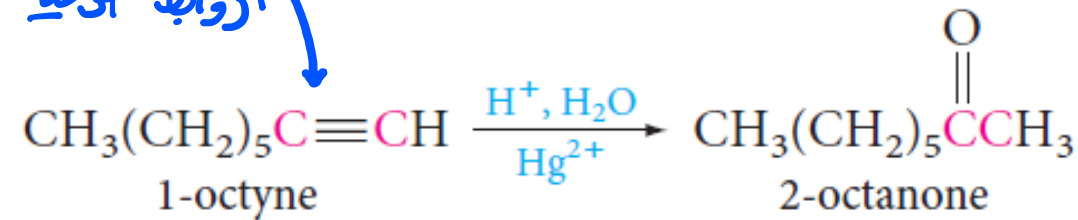


Preparation of Aldehydes and Ketones

2) Hydration of Alkynes

- 0 Hydration of **terminal alkynes** EXCEPT acetylene yields ketones (catalyzed by acid and mercuric).

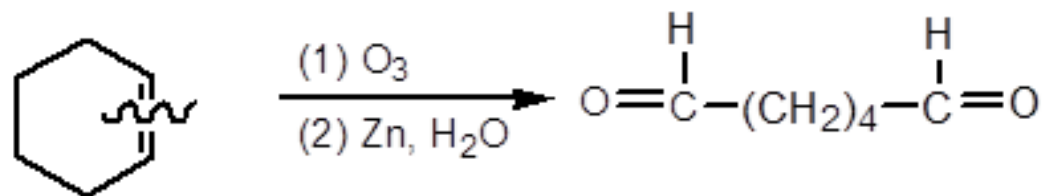
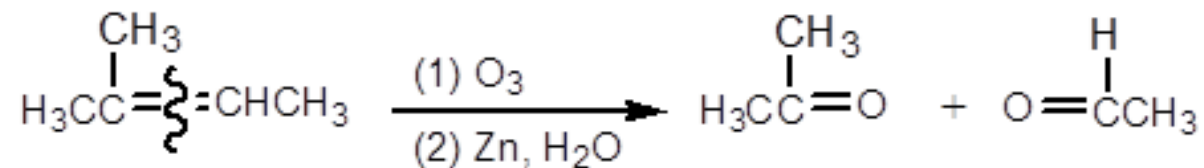
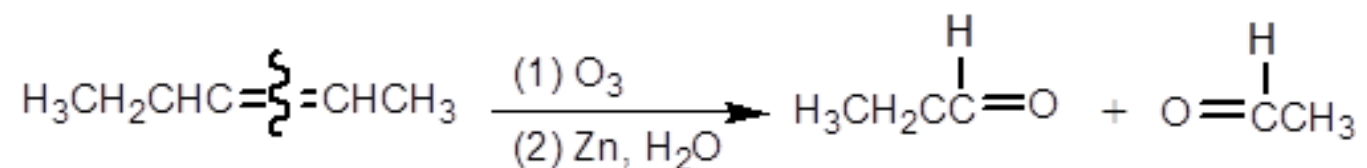
الروابط طرفية



Preparation of Aldehydes and Ketones

3) Ozonolysis of Alkenes

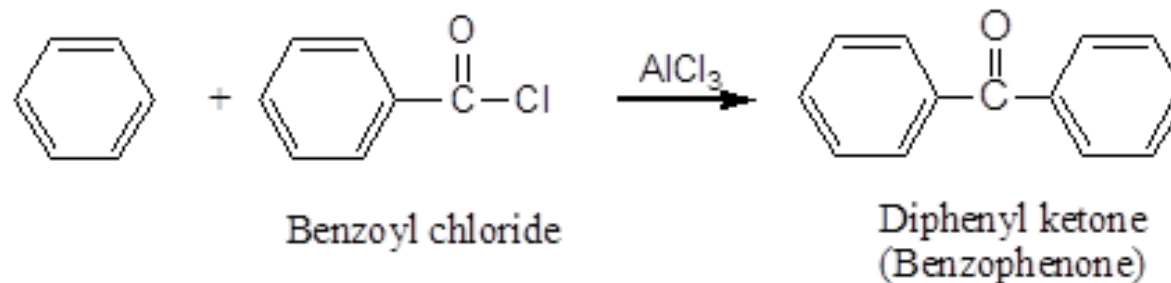
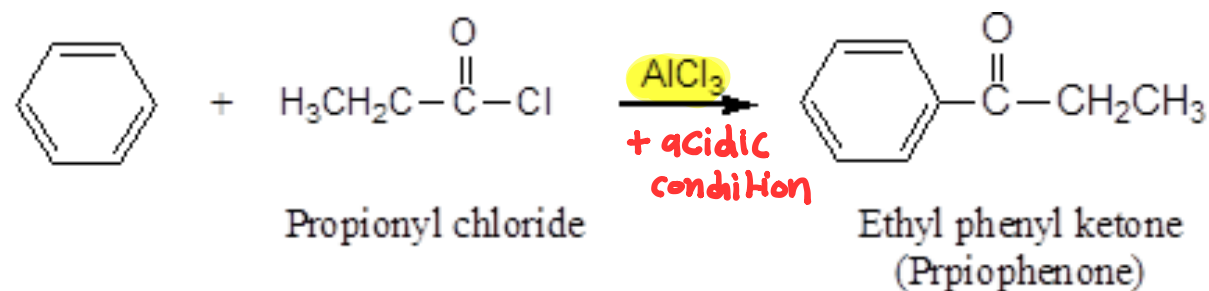
Product (aldehyde or ketone) depends on the **structure of alkene**.



Preparation of Aldehydes and Ketones

4) Friedel-Crafts Acylation

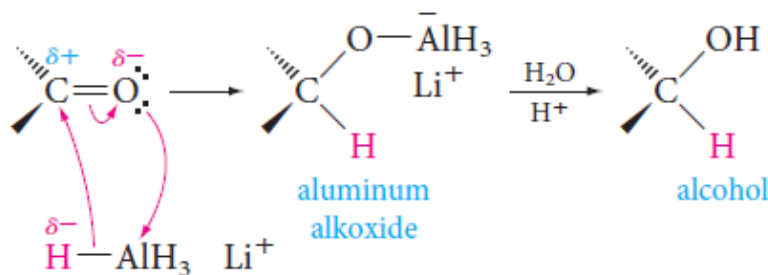
Preparing ketones that contain an aromatic ring.



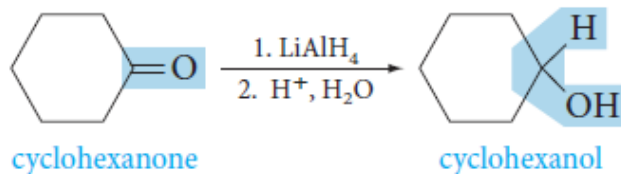
Reactions of Aldehydes and Ketones

A) Reduction of Carbonyl Compounds

- Aldehydes and ketones are easily reduced to primary and secondary alcohols, respectively.
- The most common metal hydrides used to reduce carbonyl compounds are lithium aluminum hydride (LiAlH_4) and sodium borohydride (NaBH_4).



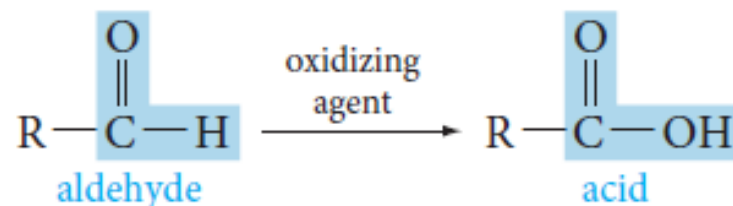
Example:



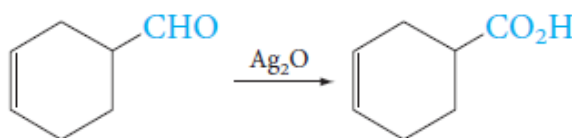
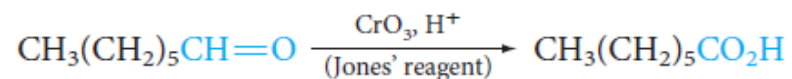
Reactions of Aldehydes and Ketones

B) Oxidation of Carbonyl Compounds

- Oxidation of aldehydes gives a carboxylic acid with the same number of carbon atoms.
- Because the reaction occurs easily, many oxidizing agents, such as KMnO_4 , CrO_3 , Ag_2O and peracids (such as, perchloric acid HClO_4 , and permanganic acid HMnO_4). will work.



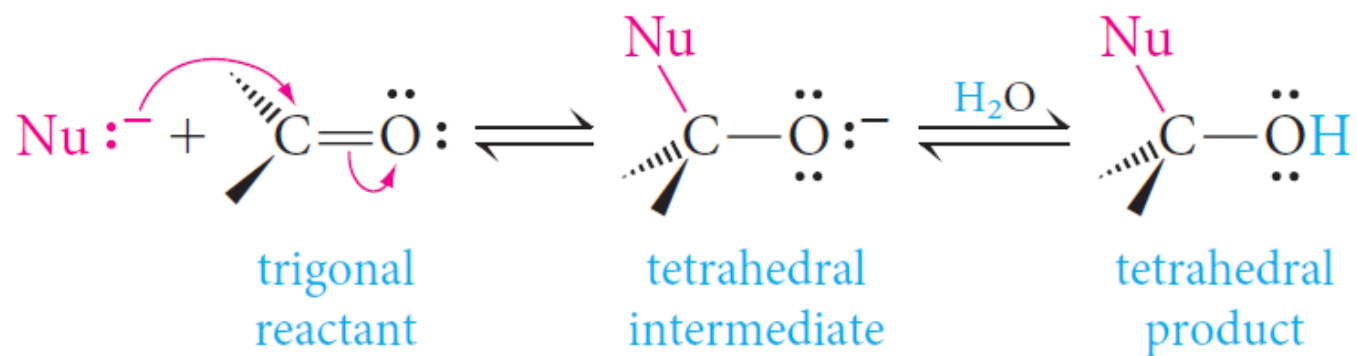
Example:



Reactions of Aldehydes and Ketones

C) Nucleophilic Addition Reactions

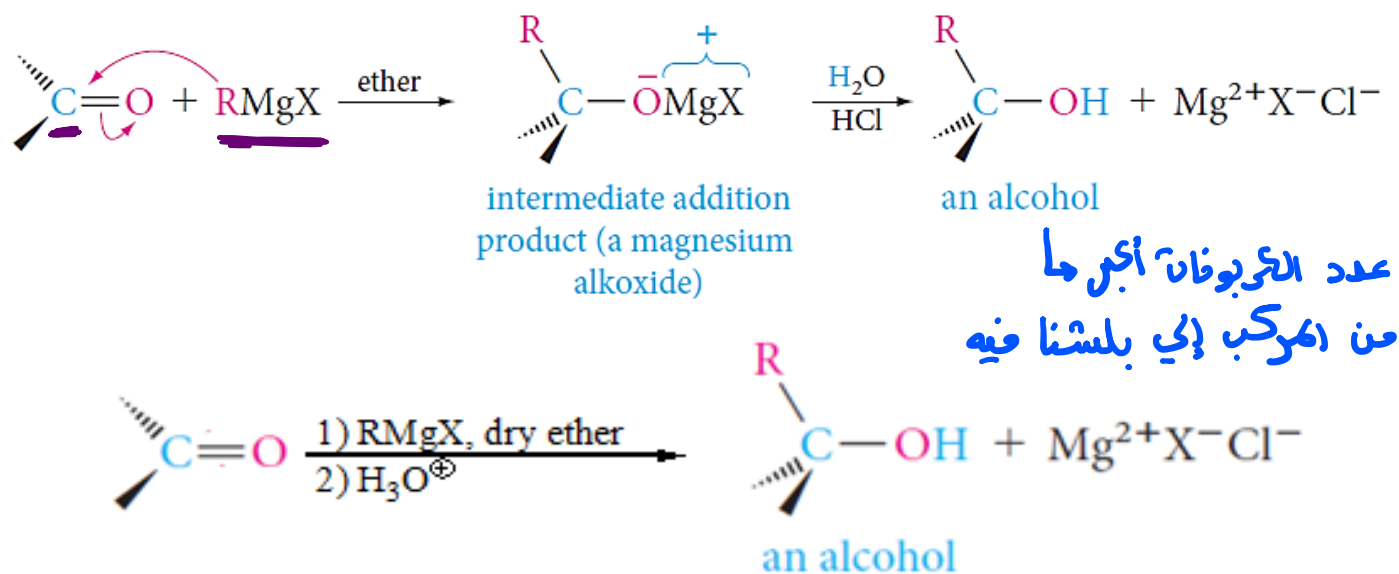
- Nucleophiles attack the carbon atom of a carbon-oxygen double bond because that carbon has a partial positive charge.
- The overall reaction involves addition of a nucleophile and a proton across the pi bond of the carbonyl group (when carried out in alcohol or water).



C) Nucleophilic Addition Reactions

1) Addition of Grignard Reagents: Formation of Alcohols

- Grignard reagents act as carbon nucleophiles toward carbonyl compounds.
- The reaction of a Grignard reagent with a carbonyl compound provides a useful route to alcohols.

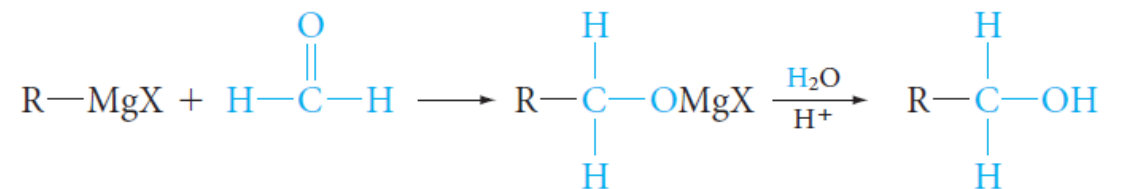


- The type of carbonyl compound chosen determines the class of alcohol produced.

C) Nucleophilic Addition Reactions

1) Addition of Grignard Reagents: Formation of Alcohols

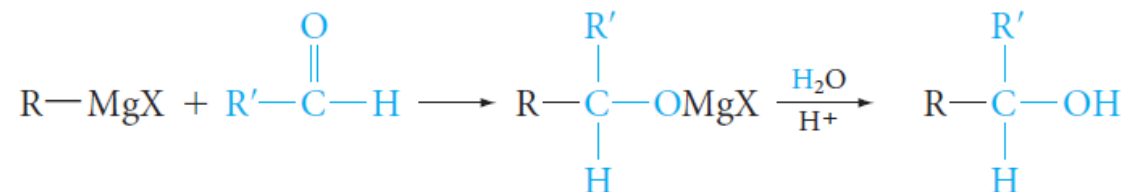
Formaldehyde gives primary alcohols.



formaldehyde

a primary alcohol

Other aldehydes give secondary alcohols

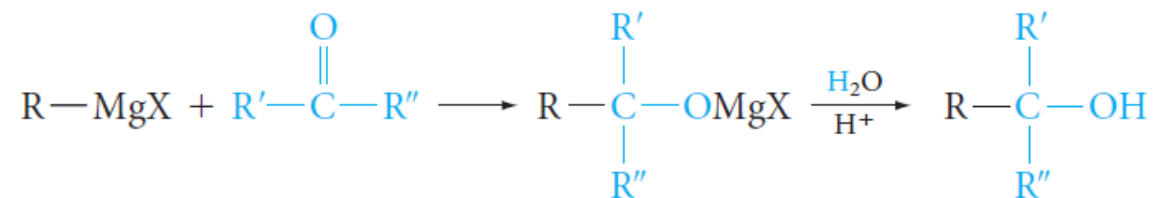


aldehyde

a secondary alcohol

هذه اكرة الوحدية
للحصول على
primary alcohol

Ketones give tertiary alcohols.



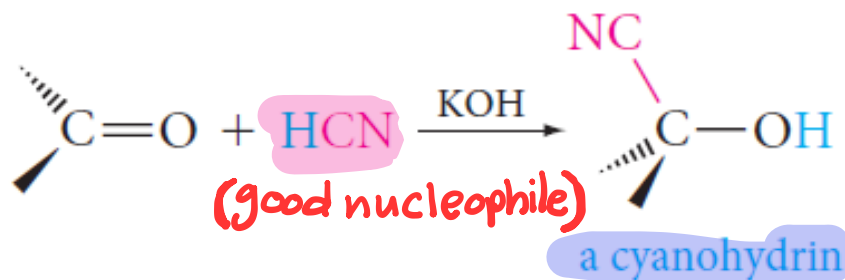
ketone

a tertiary alcohol

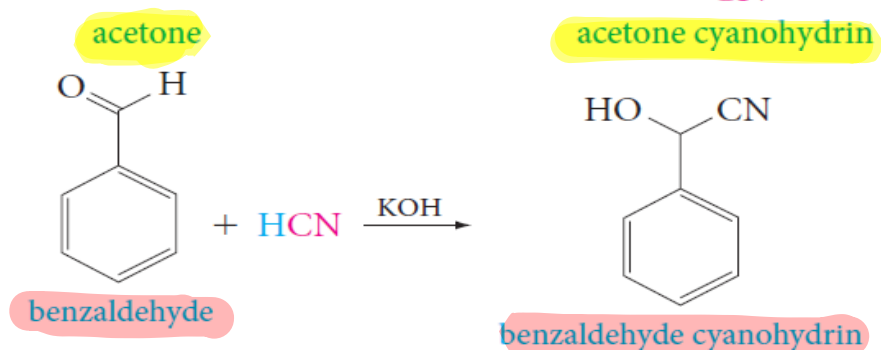
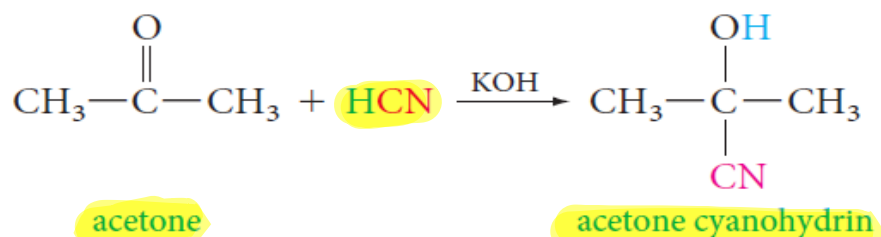
C) Nucleophilic Addition Reactions

2) Addition of Hydrogen Cyanide: Formation of Cyanohydrins

Hydrogen cyanide adds to the carbonyl group of aldehydes and ketones to form **cyanohydrins**, compounds with a hydroxyl and a cyano group attached to the same carbon.



Example



C) Nucleophilic Addition Reactions

3) Addition of Ammonia and Ammonia Derivatives

The addition of nitrogen nucleophile, such as ammonia (NH_3) and substituted ammonia ($\text{NH}_2\text{-Y}$).

