

Chapter 12



Reactions of Aldehydes and Ketones

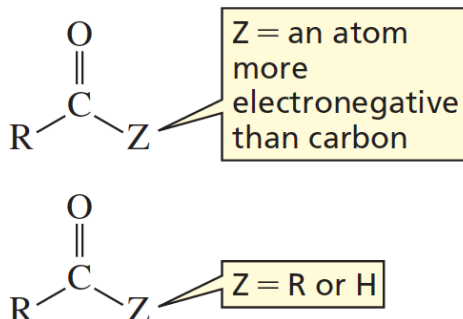
More Reactions of Carboxylic Acid Derivatives

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Santa Barbara

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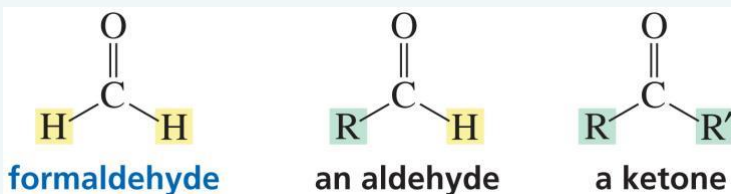
This chapter continues the discussion of the families of compounds in Group IV

Group IV



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Aldehydes and Ketones

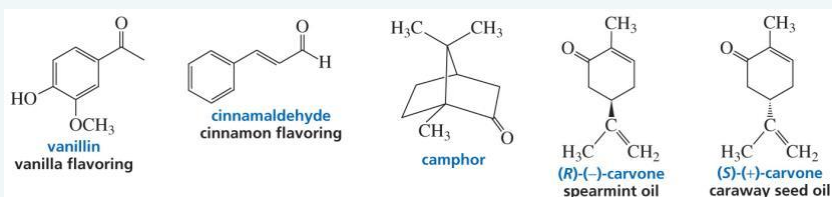


Carbonyl compounds with hydrogen and alkyl groups

- Formaldehyde: **two hydrogens**
- Aldehyde: a **hydrogen** and an **alkyl** group
- Ketone: **two alkyl** groups

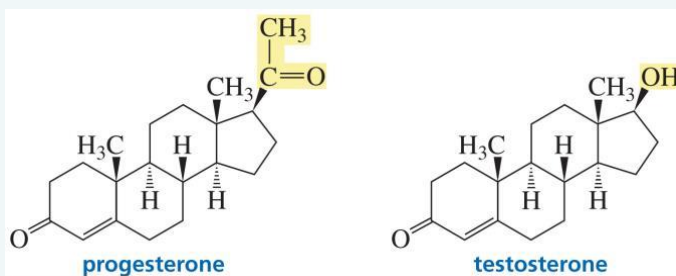
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Some Aldehydes and Ketones found in Nature



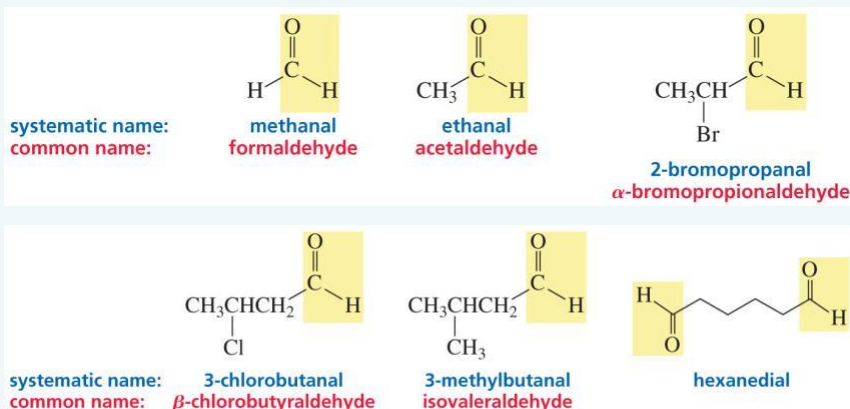
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Biologically Important Ketones



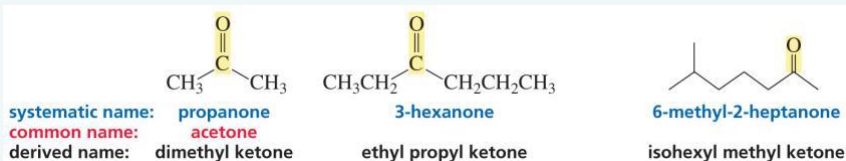
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Naming Aldehydes

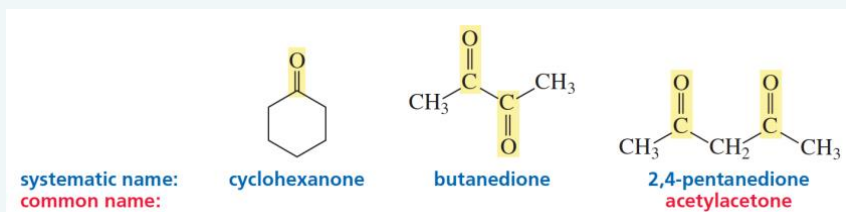


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Naming Ketones



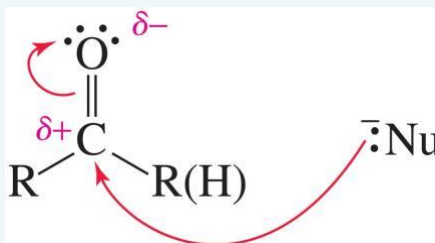
the carbonyl is assumed to be at the **1-position** in **cyclic ketones**



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Aldehydes and Ketones are attacked by Nucleophiles

The **partial positive charge** on the **carbonyl carbon** causes it to be attacked by **nucleophiles**:



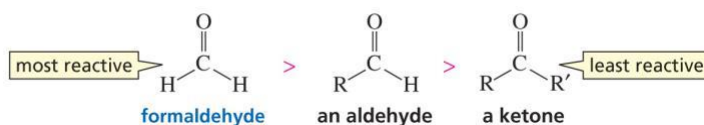
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Aldehydes are more reactive than Ketones

Electronic Reason:

An **aldehyde** has a **greater partial positive** charge on its carbonyl carbon than does a ketone because a **hydrogen is more electron withdrawing** than an alkyl group.

relative reactivities



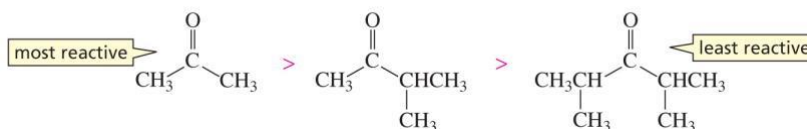
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Aldehydes are more reactive than Ketones

Steric Reason:

The carbonyl carbon of an aldehyde is **more accessible to the nucleophile**.

relative reactivities



ketones with larger R groups are even less reactive

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Aldehydes and Ketones are moderately reactive Carbonyl Compounds

relative reactivities of carbonyl compounds

acyl halide > acid anhydride > aldehyde > ketone > ester ~ carboxylic acid > amide > carboxylate ion

most reactive

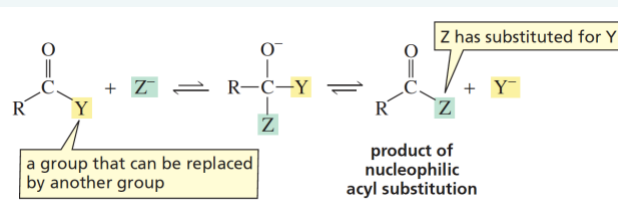
least reactive

Aldehydes and ketones are **less reactive** than acyl halides and acid anhydrides.

Aldehydes and ketones are **more reactive** than esters, carboxylic acids, amides, and carboxylate ions.

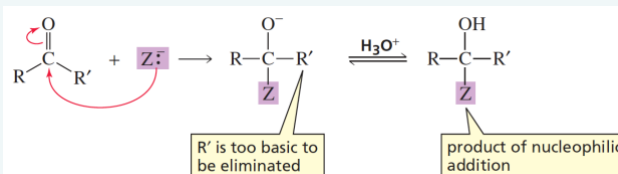
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Aldehydes and Ketones react differently than do Carboxylic Acid Derivatives



In a **carboxylic acid derivative**, Y^- can leave.

Nucleophilic acyl substitution is observed.

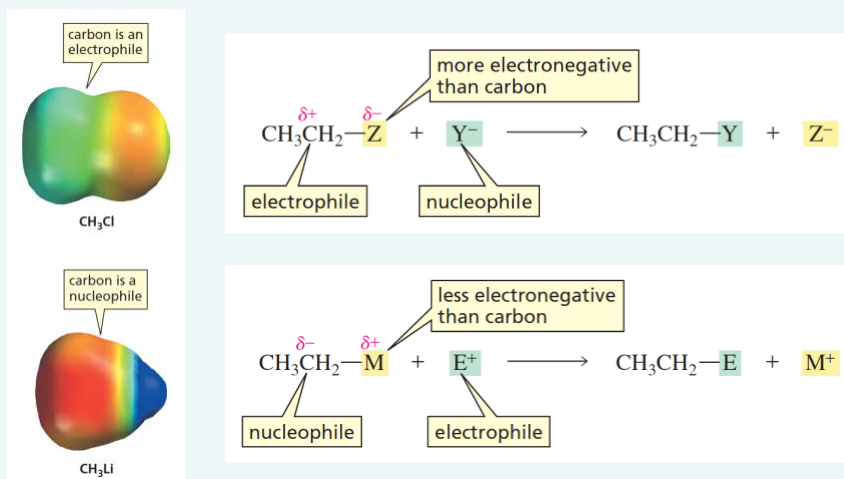


In **aldehydes and ketones**, neither alkyl nor hydride groups can leave.

Nucleophilic addition is observed.

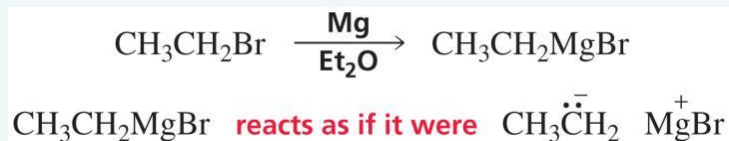
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Organometallic Compounds

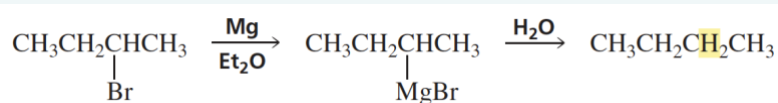


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Organometallic Compounds: Grignard Reagents



Grignard reagents react as carbanions.



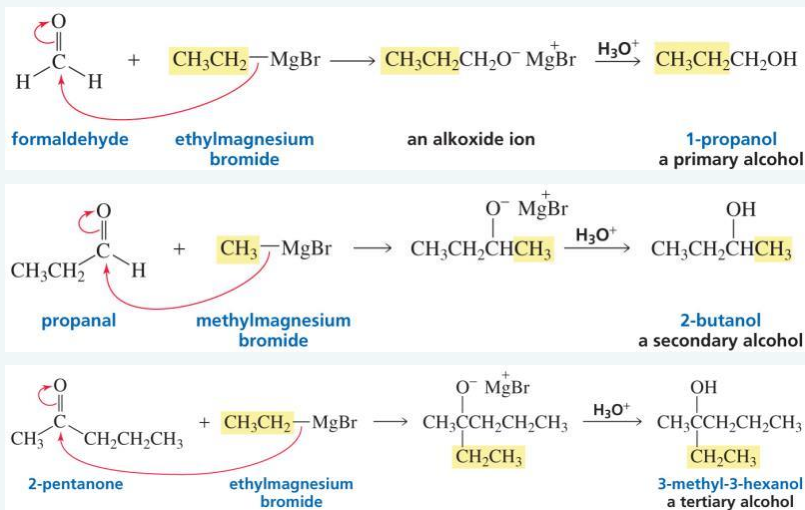
They are strong bases and react vigorously with water.

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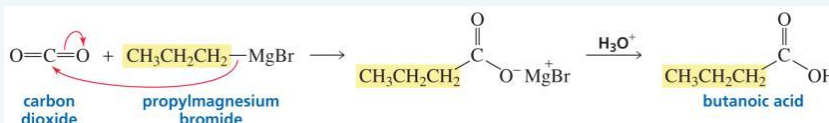
$$\text{R}-\overset{\text{O}}{\underset{\text{R}(\text{H})}{\text{C}}} + \text{R}'-\text{MgBr} \longrightarrow \text{R}-\overset{\text{O}^- \text{ Mg}^+\text{Br}}{\underset{\text{R}'}{\text{C}}}-\text{R}(\text{H}) \xrightarrow{\text{H}_3\text{O}^+} \text{R}-\overset{\text{OH}}{\underset{\text{R}'}{\text{C}}}-\text{R}(\text{H})$$

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Grignard Reagents are used to prepare alcohols



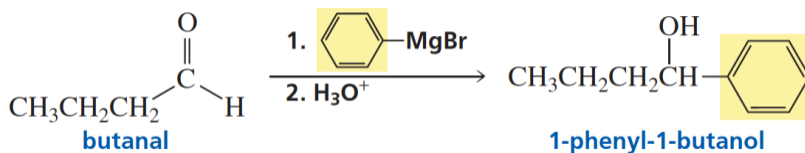
Grignard Reagents form Carboxylic Acids by reaction with Carbon Dioxide



The carboxylic acid has **one more carbon** than the Grignard reagent.

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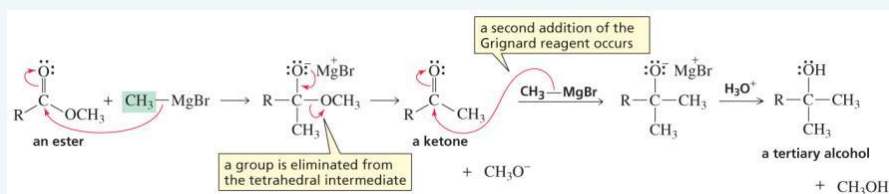
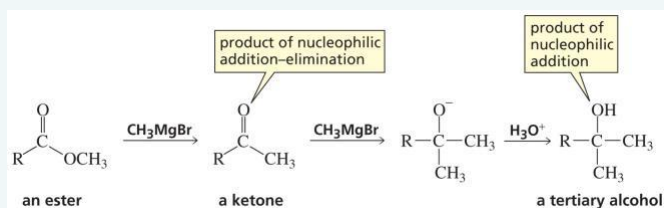
Examples of reactions with Grignard Reagents



- The numbers above and below the arrows mean the acid is not added until the Grignard reagent has reacted with the carbonyl compound.
- The product of this reaction is a **racemic mixture** because a compound **with** an asymmetric center was created from a reactant **without** an asymmetric center.

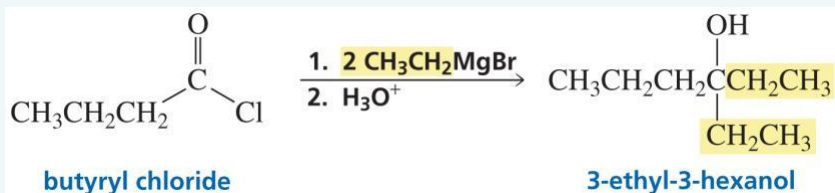
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Esters react with Grignard Reagents to form alcohols



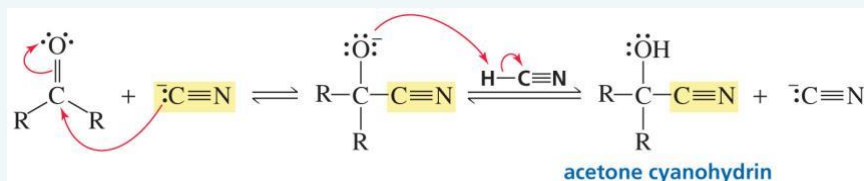
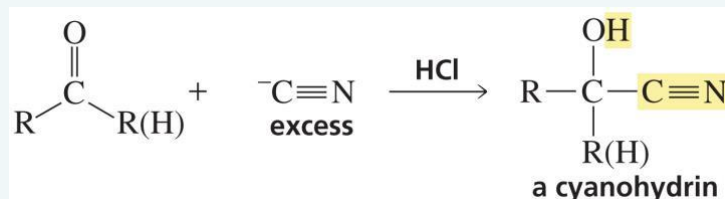
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Acyl Chlorides react with Grignard Reagents to form alcohols



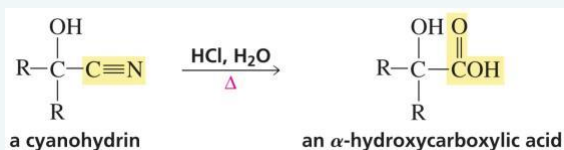
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Reaction of Aldehydes and Ketones with Cyanide Ion

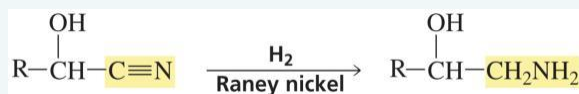


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Reactions of Cyanohydrins



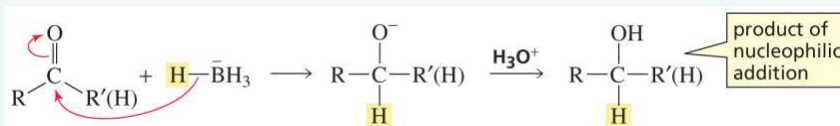
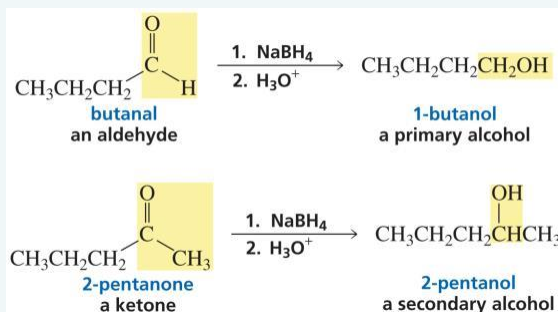
Hydrolysis to form an α -hydroxycarboxylic acid.



Reduction to a primary amine with an OH group on the β -carbon.

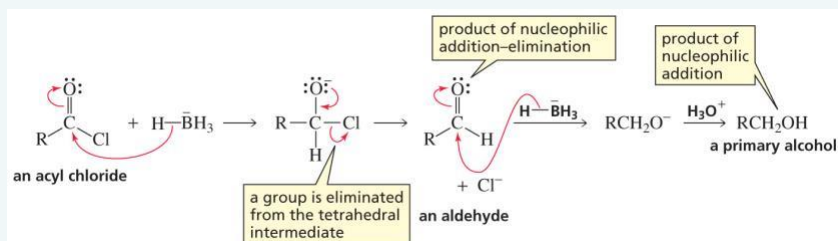
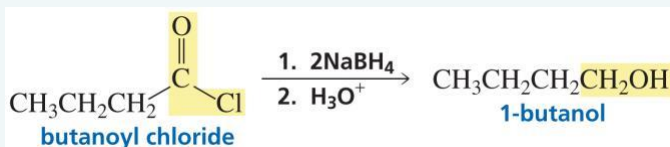
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Reactions of Aldehydes and Ketones with Hydride Ion



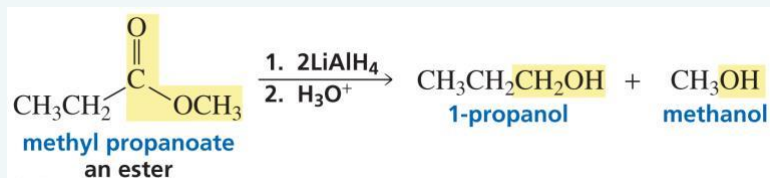
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Reactions of Acyl Chlorides with Hydride Ion



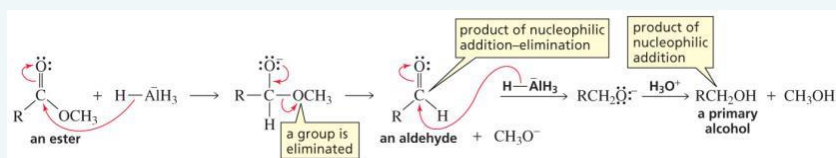
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Reaction of an Ester with Hydride Ion



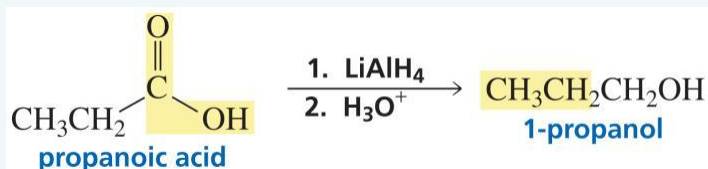
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Mechanism for the reaction of an Ester with Hydride Ion



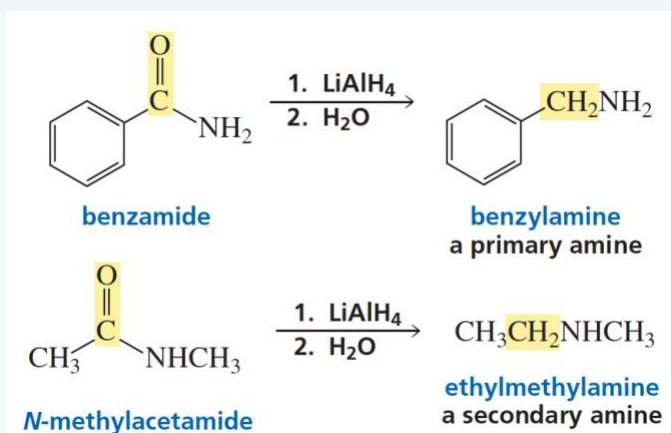
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Reactions of Carboxylic Acids with Hydride Ion



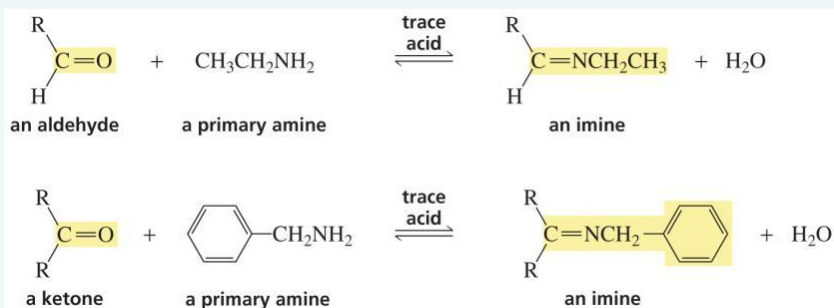
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The Reactions of Amides with Hydride Ion



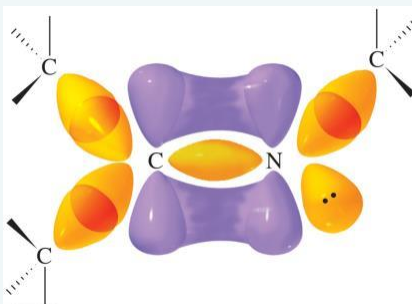
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Aldehydes and Ketones form Imines with Primary Amines



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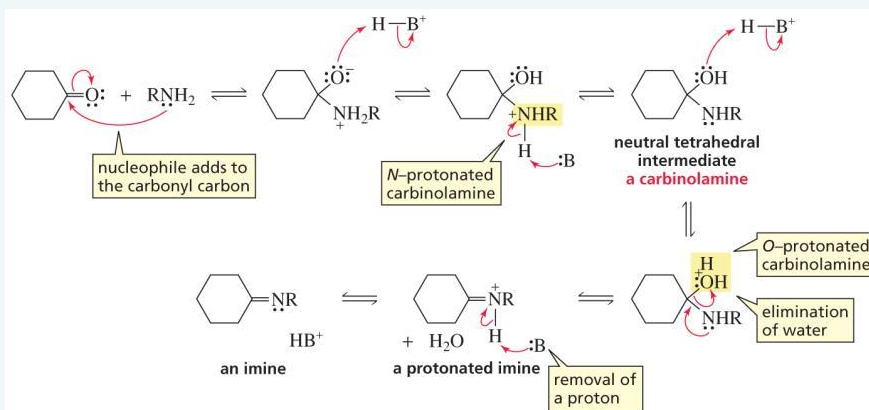
Bonding in an Imine



- The π bond is formed by side-to-side overlap of a p orbital of carbon with a p orbital of nitrogen.
- The π bond is perpendicular to the **orange orbitals**.

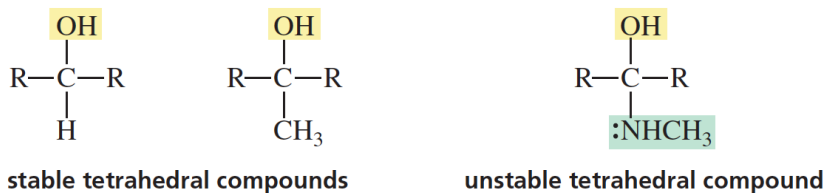
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Mechanism for Imine Formation



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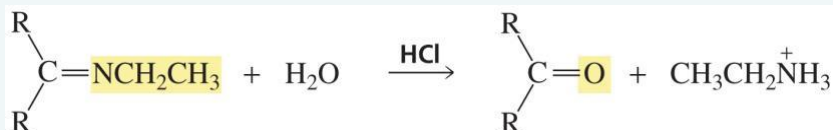
Intermediates in Imine Formation are unstable



- Imine formation is reversible because there are two protonated intermediates in the mechanism that can eliminate a group.
- Imine formation can be pushed to completion by removing water as it is formed.

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Imine Hydrolysis is Irreversible

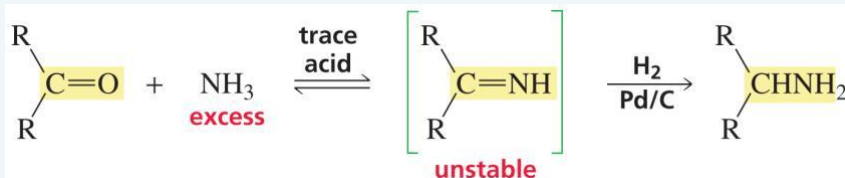


The **amine** is **protonated** in the acidic solution, so it is unable to react with the carbonyl compound.

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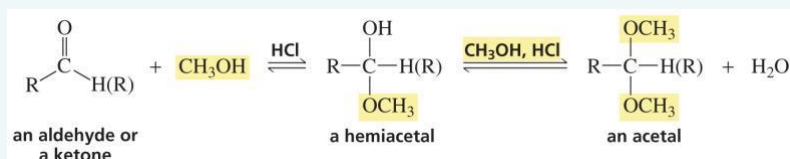
Reductive Amination

The unstable **imine** formed from ammonia is hydrogenated to an **amine**.



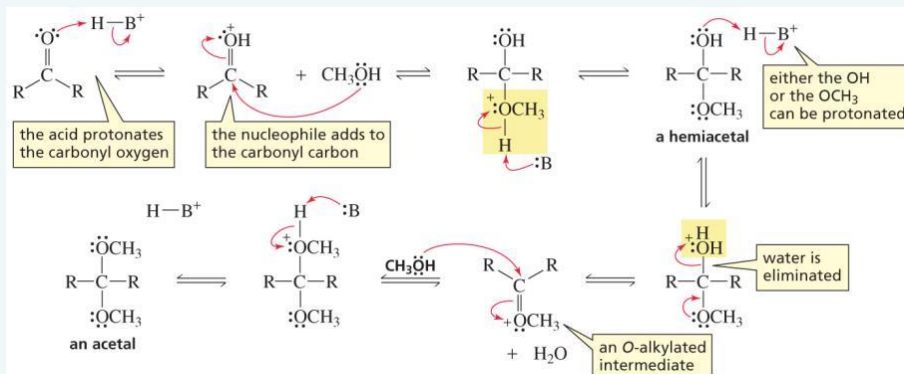
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The reaction of an Aldehyde or Ketone with an alcohol



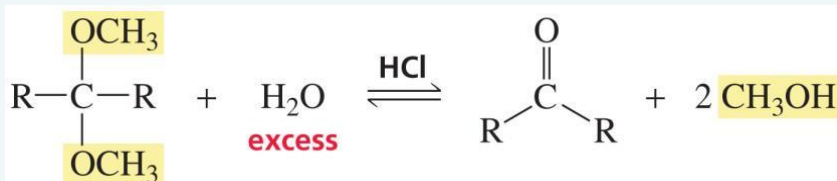
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Mechanism for the reaction of an Aldehyde or Ketone with an alcohol



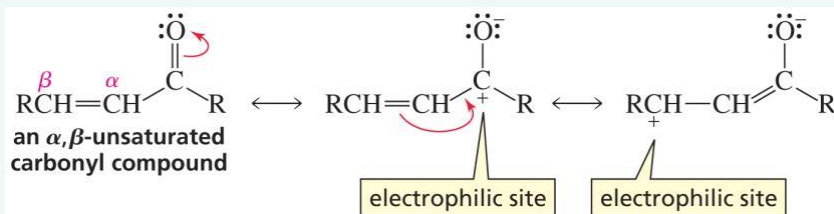
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Acid-Catalyzed Hydrolysis of an Acetal



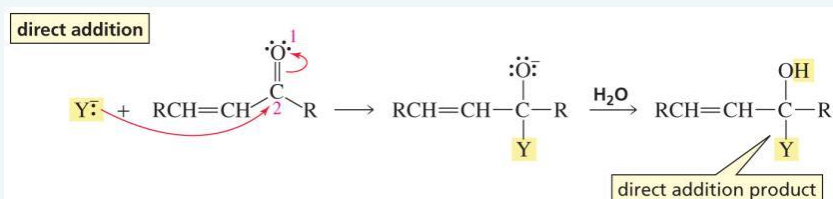
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α,β -Unsaturated Aldehydes and Ketones have two Electrophilic Sites



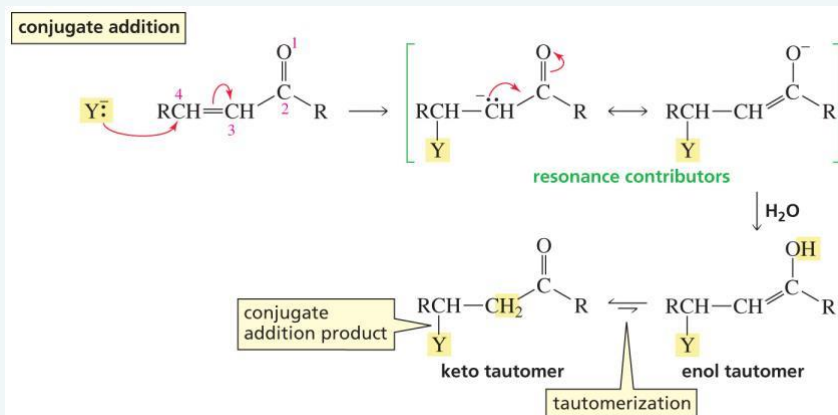
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Direct addition to α,β -Unsaturated Aldehydes and Ketones



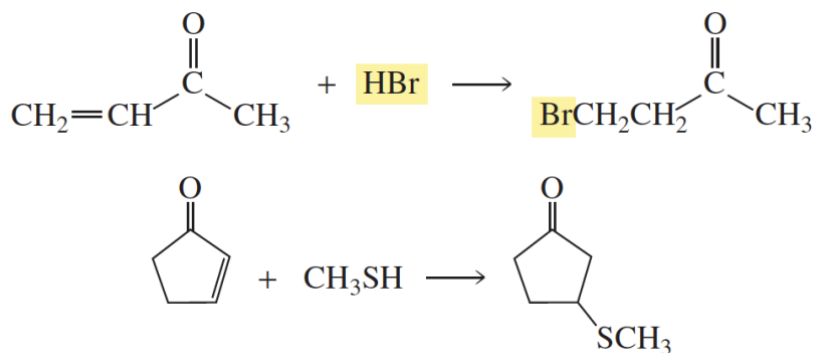
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Conjugate addition to α,β -Unsaturated Aldehydes and Ketones



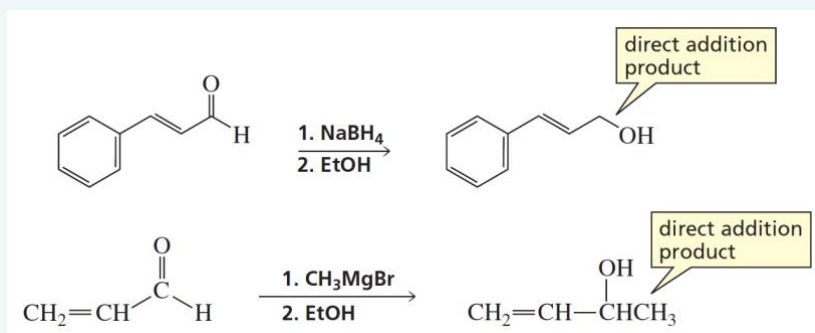
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Weak Bases form Conjugate Addition products



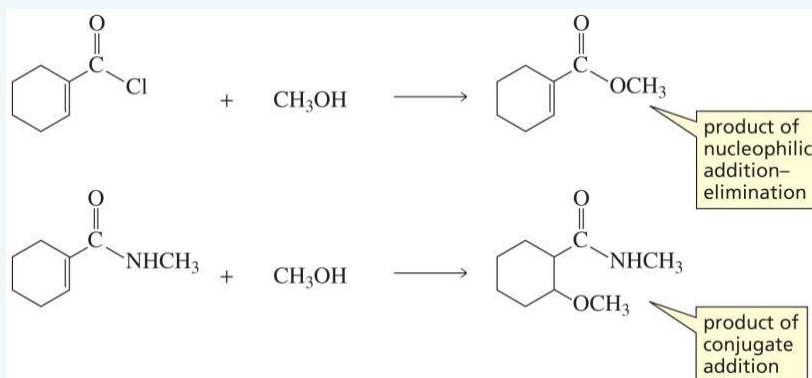
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Strong Bases form Direct Addition products



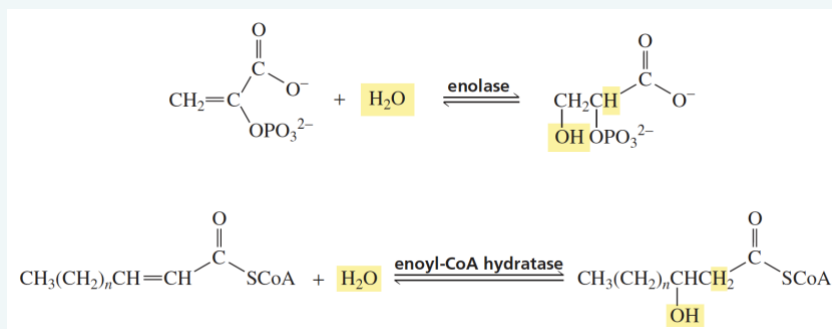
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Nucleophilic Addition to α,β -Unsaturated Carboxylic Acid Derivatives



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Conjugate Addition Reactions in Biological Systems



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