



ACADEMIC

ORGANIC CHEMISTRY

REACTIONS

FEATURES OF AN ORGANIC REACTION

Mechanism: Describes the overall reaction using a series of simple steps.

Stoichiometry: Calculate reactant and product masses using the balanced equation and molar masses.

Kinetics: The study of the reaction rate and mechanism.

Theoretical yield: Mass of product given by a complete reaction;

$$\% \text{yield} = 100\% \times (\text{product mass}) / (\text{theoretical yield})$$

Equilibrium: Reaction does not proceed to completion, instead, it reaches a balanced state of forward and reverse reactions.

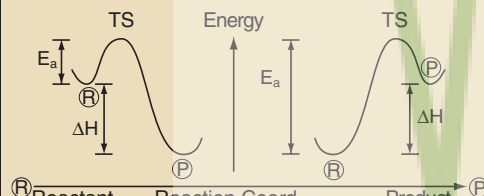
Major reaction types:

- Acid
- Base
- Oxidation Reduction
- Condensation
- Substitution (SN1, SN2)
- Ionic Reaction
- Elimination (E1, E2)
- Cyclization
- Hydrolysis
- Addition
- Radical reaction

Important named reactions:

- **Diels-Alder:** form cyclic alkene
- **Friedel-Crafts:** add acyl or alkyl group
- **Grignard:** add alkyl or aryl group
- **Wolf-Kischner, Clemmenson:** reduce ketone to alkane
- **Wittig:** convert aldehyde/ketone to alkene

KINETICS AND REACTION MECHANISM



Transition state (TS): Maximum on the reaction-coordinate curve: the least stable intermediate.

Activation Energy (E_a): Energy of the TS relative to the reactant. The change in enthalpy (ΔH) is < 0 for **exothermic**; > 0 for **endothermic**.

Hammond-Leffler postulate: The TS is more like the reactant or product that is closer in energy; *endothermic* TS is like the product, *exothermic* TS is like the reactant.

Kinetic vs. thermodynamic control: ΔG and ΔH describe Thermodynamic Stability.

- If ΔG is large and negative (**exergonic**), the product formation is likely controlled by "thermodynamics."

Large K_{eq} corresponds to a large amount of product, relative to reactant.

- A large E_a may give rise to "kinetic" control; the energy of the TS controls the reaction, instead of the product-reactant thermodynamics.

Solvent effects: A solvent may stabilize an intermediate, decreasing E_a and increasing the rate of the reaction. Charged-complexes are stabilized by polar solvents.

ORGANIC ACID AND BASE

Acid:

- Electron-pair acceptor (Lewis acid)
- Proton donor (Bronsted-Lowry acid); example: carboxylic acid

Base:

- Electron-pair donor (Lewis base)
- Proton acceptor (Bronsted-Lowry base); example: amine

Factors enhancing acid strength (HA):

- Weaker H-A bond
- Greater electronegativity of "A"
- Inductive effect of substituent on "A" (electron withdrawal enhances transfer).
- More "s" character in hybrid orbital (s-orbital is lower in energy than p-orbital)
- Resonance stabilized conjugate base (A^-)

Factors enhancing base strength:

- Reverse of acid-strength guidelines
- A base is a nucleophile; Electronic effects which shift electron density to the atom with the lone-pair increases base-strength.

ALKANE

Properties:

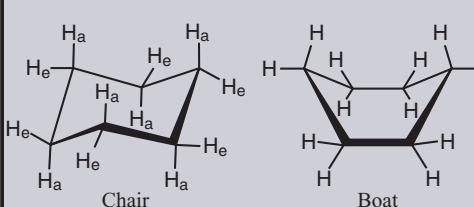
- Hydrocarbon
- Weak intermolecular forces
- Non-cyclic: general formula C_nH_{2n+2}
- Tetrahedral C-C-C (109°)

Nomenclature:

- Add "-ane" to prefix
- Locate substituent by position #
- Haloalkane: substitute halide for -H

Cycloalkane: (C_nH_{2n})

- Bicyclic - two fused or bridged rings
- n = 3: **cyclopropane:** (highly strained)
- n = 4: **cyclobutane:** (some flexibility)
- n = 5: **cyclopentane:** (slight puckering)
- n = 6: **cyclohexane:** *chair* - stable conformer; *boat* - less stable; *Axial* position: "perpendicular" to ring; *Equatorial* position: in ring "plane" (see H_a and H_e in chair diagram below)
- Cis - two substituents in *up* position
- Trans - *one up and one down*



Synthesis:

- Hydrogenate alkene or alkyne (H_2 , Pt catalyst)
- Free-radical reaction of alkene
- Reduce haloalkane (Zn, H^+)
- Friedel-Crafts alkylation

Reaction:

- Combustion: alkane + $O_2 \Rightarrow CO_2 + H_2O$
- Halogenation to haloalkane (Cl_2/Br_2 , light or heat)

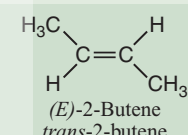
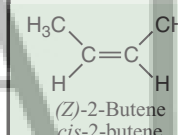
ALKENE >C=C<

Properties: Similar to alkane; non-polar, flammable

Nomenclature:

- Add -ene to prefix; Use # to denote C=C position
- **Isolated** C-C=C-C-C; **cumulative** -C=C=C-
- Polyunsaturated fatty acid: 2 or more C=C
- Allene: adjacent C=C-C
- Vinyl group: $H_2C=CH-$
- Methylene group: $H_2C=$
- Allyl group: $H_2C=CH-CH_2-$
- Vinyl halide: halide replaces -H on $>C=C<$
- Conjugated: alternate C-C and C=C (resonance)
- Alkadiene, 2 conjugated C=C; example: butadiene; s-cis and s-trans (rotate about C-C bond)
- Alkatriene, 3 conjugated C=C
- Annulene: conjugated monocyclic compound; example: [6] annulene = benzene
- Aromatic cyclic ions: cyclopentadiene anion, cycloheptatriene cation (6 electrons)

Isomers: no free rotation of C=C



- E/Z: prioritize groups by atomic weight (Z - higher priority groups on the same side)
- For noncyclic: cis is less stable (steric hindrance). For cyclic, cis more stable.
- **Hofmann Rule:** Form the least-substituted alkene
- **Markovnikov of Addition:** H adds to C with most -H's
- **Zaitsev Elimination:** Form alkene with more substitution

Synthesis:

- Dehydrate alcohol (H^+ , heat) (elimination)
- Dehydrohalogenate haloalkane (base, heat)
- Dehalogenate vic dihalide (Zn , acetic acid)
- Hydrogenate alkyne:
 - syn, Z/cis-isomer ($H_2/P-2$ catalyst)
 - anti, E/trans-isomer ($Li, NH_3, -78^\circ C$)
- Wittig, aldehyde/ketone + phosphorous ylide

Reaction:

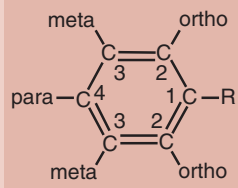
- Combustion (O_2)
- Hydrate to 2/3° alcohol (H^+, H_2O); 1° from ethene; can rearrange (Markovnikov)
- Hydrate to alcohol; hydroborate/oxidize ($THF/B_2H_6, H_2O_2/OH^-$) (syn, anti-Markovnikov)
- Oxymercuroate-demercurate to alcohol
- Hydrohalogenate (HX) (Markovnikov)
- Halogenate (Br_2/Cl_2), vic dihaloalkane (X_2, CCl_4 ; anti)
- Halohydrin (X_2, H_2O ; anti-addition)
- Hydroxylate to form a 1,2-diol ($KMnO_4$, cold OH^- ; syn addition)
- Oxidize to carboxylic acid ($KMnO_4$, hot OH^-)
- Ozonolyze to ketone ($O_3; Zn, H_2O$)
- Hydrogenate to alkane (Pt, H_2 ; syn-addition)
- Free radical polymerization
- Alkadiene Reaction
 - allylic halogenation (Cl_2 , heat)
 - Diels-Alder: cycloalkene from diene + alkene/alkyne

BENZENE/ARENES

Properties: insoluble in water, miscible with non-polar organic solvents.

Nomenclature:

- Aromatic** (or arene): Denote substituent using group name and ring position; ortho (1,2), meta (1,3), para (1,4); examples: benzene C_6H_6 ; phenol, Ar-OH (carboxic acid, hydroxybenzene, benzenol); aniline Ar-NH₂; toluene, Ar-Me (methyl benzene); xylene, dimethyl benzene
- Fused rings:** naphthalene, $C_{10}H_8$ (2 edge-sharing rings)
- Aryl or Phenyl group:** Ar- (remove H from benzene)
- Aryl halide:** halogen replaces an H atom; Ar-X
- Alkenyl benzene:** Ar-C=C<
- Benzyl:** Ar-CH₂-



Synthesis: Dehydrogenate cyclohexane (sulfur+ heat)

General Reaction:

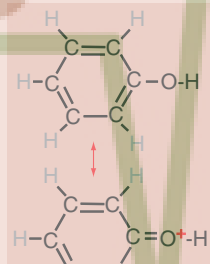
- Combustion (similar to alkane)
- Birch reduction \Rightarrow 1,4 cyclohexadiene (Na , NH_3 , $EtOH$)
- Hydrogenate to cyclohexane (H_2 , Pt)

Electrophilic substitution:

- Alkylation: Ar-R (*Friedels-Crafts*, RCl , $AlCl_3$)
- Nitration: Ar-NO₂ (HNO_3 , H_2SO_4)
- Halogenation: Ar-Br (Br_2 , $FeBr_3$)
 - Ar-Cl (Cl_2 , $FeCl_3$)
 - Ar-I (I_2 , HNO_3)
- Acylation: Ar-CR=O ($RCOCl$, $AlCl_3$)
- Sulfonation: Ar-SO₃H (SO_3H , H_2SO_4)

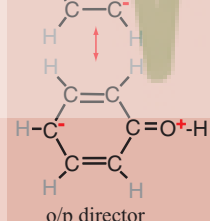
Reactivity of substituted benzene:

- A substituent alters the ring electronic structure.
- Activating group:** More reactive than benzene; add electrons to the ring, stabilize the arenium cation
- Deactivating group:** Less reactive; pull electrons from the ring, destabilize the arenium cation

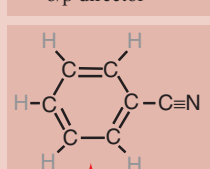


- Ortho/para-director:**
 - substituent tends to activate the ring (except for -X); electron density donated to ring creates “-” center on o/p sites, o/p isomers are preferred
 - examples: -NR₂, -OH, -R, -OR, -X (halogen)

- Meta-director:**
 - substituent tends to deactivate the ring; electron density withdrawn from the ring creating “+” center on o/p site, m-preferred reaction site.
 - examples: -NO₂, -CN, -COOH, -SO₃H, -COOR, -CHO, -CRO

**Reactivity of di-substituted benzene:**

- Directing effects may be cooperative; e.g. “o/p” plus “m” at 1,4 positions
- Otherwise: consider steric effects; activating group tends to dominate deactivating group.

**Reaction of alkyl substituted benzene:**

- Toluene to benzoic acid: ($KMnO_4$, OH^- , heat, H^+)
- Chlorinate -Me of Toluene (Cl_2)

ALKYNE -C≡C-**Properties:**

- Hydrocarbon, at least 1 C≡C triple bond
- Properties similar to alkane or alkene
- Linear $R''-C\equiv C-R'$

Nomenclature:

- Add -yne to prefix
- Number denotes position of triple bond; example: ethyne (acetylene) C_2H_2

Synthesis:

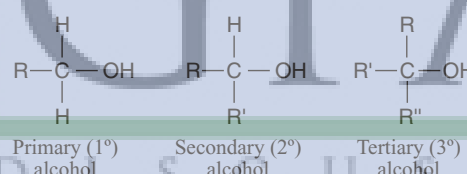
- $CaC_2 + H_2O \Rightarrow Ca(OH)_2 + C_2H_2$
- Dehydrohalogenate vic-haloalkene ($NaNH_2$, liq NH_3)
- Alkylate terminal alkyne ($NaNH_2$, liq NH_3 ; $R-X$)

Reaction:

- Addition: hydrogenate to alkane (H_2 , Pt or Ni)
 - syn to cis/Z alkene (H_2/Ni_2B P-2 catalyst)
 - anti to trans/E alkene (Li , liq NH_3)
- haloalkene to gem-dihalide (HX) (Markovnikov)
- halogenate to haloalkene or haloalkane (X_2)
- Ozonolyze to carboxylic acid (O_3 , H_2O)
- Oxidize to carboxylic acid ($KMnO_4$, OH^- ; H^+)

ALCOHOL R-OH**Properties:**

- Low molecular weight are water-soluble
- H-bonding, polar
- RO-H acidic proton
- Resonance stabilized ArO^- or RO^-

**Nomenclature:**

- Prefix + “anol”; example: methanol $Me-OH$ (methyl alcohol)
- Cyanohydrin:** -OH and -CN
- Halohydrin: -OH and halogen
- Diol or glycol (two -OH); **gem-diol:** 1,1 diol; **vic-diol:** 1,2 diol

Synthesis:

- Hydrate alkene (H_2O , H^+)
- Hydroborate/oxidize alkene (THF: BH_3 ; H_2O_2 , OH^-)
- Hydrogenate aldehyde (H_2/Ni or Pt catalyst)
- Hydrolyze 1^o alkyl halide (water, OH^-)
- Reduce aldehyde, ester, ketone or carboxylic acid
- Ethanol: Ferment sugar or starch
- Methanol: $CO + H_2$, catalyst; Pyrolyze cellulose
- Hydrolyze ester (water, acid)
- Dehydrate ether (H_2SO_4 , low heat)
- Grignard ($RMgX$): formaldehyde \Rightarrow 1^o alcohol; aldehyde \Rightarrow 2^o alcohol; ketone \Rightarrow 3^o alcohol
- Synthesis of Glycol from Ketone/aldehyde: (HIO_4 or $Pb(OAc)_4$; H_2SO_4 + heat)
- oxidize alkene: ($KMnO_4$: cis) (H_2O_2 , formic acid: trans)
- hydrolyze epoxide (H_2O , H_2SO_4)

Reaction:

- Oxidize 1^o to aldehyde (CuO , heat) or 2^o to ketone ($KMnO_4$, H^+)
- Oxidize 1^o to carboxylic acid ($KMnO_4$, H^+)
- Dehydrate to alkene; Zaitsev's rule; rate 3^o>2^o>1^o (hot H_2SO_4 , or Al_2O_3)
- Dehydrate to ether (H_2SO_4 , lower temperature)
- Oxidize to ketone (2^o alcohol) (H_2CrO_2)
- Form haloalkane (HX ; substitution)

AROMATIC ALCOHOL Ar-OH**Properties:**

- The most common is Phenol, Ar-OH
- Acidic hydrogen, $ArO-H$; $pK_a=9.9$
- Ring substituent alters acidity
- Benzendiol, $HO-Ar-OH$; para, hydroquinone; ortho, catechol; meta, resorcinol

Reaction of Phenol:

- Electrophilic substitution: o-p director
- Hydrogenate to cyclohexane (H_2 , catalyst)
- Form ester (acid anhydride or acid chloride)

Synthesis of Phenol:

- Electrophilic aromatic substitution
- Williamson reaction, phenyl ether ($NaOH$, RCH_2X)
- Arenediazonium salt intermediate: $Ar-NH_2 + HONO \Rightarrow Ar-N_2^+ + Cu_2O, H_2O \Rightarrow Ar-OH$
- Benzene + propene \Rightarrow cumene; oxidation/acid \Rightarrow phenol + acetone
- Aryl halide (Ar-X) + $NaOH$, heat and acid
- Ar-OR + HI/HBr , heat

HALOALKANE/ALKENE/ARENES R-X**Nomenclature:**

- Halogen (X = fluorine, chlorine, bromine or iodine) replaces -H on hydrocarbon group
- Denote halogen in the name; example: Chloromethane: $Cl-Me$; chlorobenzene $Ar-Cl$

Synthesis: alcohol (ROH) + HX**Reaction:**

- Dehydrohalogenate to alkene (often rearranges)
- Hydrolyze 1^o alkyl halide to alcohol ($RX + OH^-$)

HALOHYDRIN X-R-R'-OH**Synthesis:** Alkene + X_2 , H_2O **Reaction:**

- Halohydrin + ROH \Rightarrow β hydroxy ether
- Halohydrin + $RNH_2 \Rightarrow$ β hydroxy amine
- Halohydrin + $RSH \Rightarrow$ β hydroxy sulfide

ETHER R''-O-R'**Properties:**

- Polar, hydrogen bonding
- Oxygen lone-pair is a nucleophile
- Flammable liquid

Nomenclature:

- $R''-O-R'$, “R' R'' ether” or “alkoxy alkane”;
- Example: diethyl ether, common solvent: $Et-O-Et$
- Alkoxy group:** $-OR$ ($O-Me$, methoxy; $O-Et$, ethoxy)
- Oxa-: substitute an -O- for a -CH₂-
- Cyclic ether: tetrahydrofuran (THF)
- Epoxide or oxirane:** 3-member ring
- Dioxane:** cyclic double ether
- Peroxide:** $R-O-O-R'$; $-O-O-$ single-bond

Synthesis:

- Williamson synthesis ($R'I + NaOR$)
- Dehydrate 1^o alcohol (H_2SO_4 , heat)
- Epoxidation: alkene + peroxyacid
- Halohydrin + ROH \Rightarrow hydroxy ether

Reaction:

- Hydrolyze to alcohol (H^+ or OH^-)
- Autoxidize to peroxide (oxygen in air); **EXPLOSIVE HAZARD!**

Epoxide reaction:

- Hydrolyze 1,2 glycol (acid, H^+)
- Hydrolyze to 1,2 glycol (base, OH^- or OR^-)
- Grignard + epoxide \Rightarrow 1^o alcohol

ALDEHYDE & KETONE >C=O

Properties:

- Polar $>\text{C}^+=\text{O}^-$; low molecular weight are water-soluble
- Main chemical difference: ketone is harder to oxidize than aldehyde.

Aldehyde nomenclature (RCHO):

- Prefix +“anal”;
- Example: HCHO, methanal (formaldehyde); MeCHO, ethanal (acetaldehyde); Ar-CHO, benzaldehyde

Ketone nomenclature (RR'CO):

- Prefix +“anone,” also “R,R’ ketone”;
- Example: 2-propanone (acetone or dimethyl ketone);
- Diketone: 2 $>\text{C}=\text{O}$ groups
- Acyl group: $\text{RC}=\text{O}$ or $\text{Ar}-\text{C}=\text{O}$
- Ketene: $\text{C}=\text{C}=\text{O}$
- Ketal: $\text{RR}'\text{C}(\text{OR})(\text{OR})$;
- Acetal: $\text{RHC}(\text{OR})(\text{OR})$
- Hemiacetal: $\text{RHC}(\text{OH})(\text{OR})$
- Diketone: $\text{R}'-\text{CO}-\text{CH}_2-\text{CO}-\text{R}$

Keto-enol tautomerism:



- Nucleophile attacks $>\text{C}=\text{C}<$ of enol-form
- Acidic α -H, $-\text{CH}^*-\text{CHO}$ can form resonance stabilized carbanion (especially for diketone).
- Racemization via keto-enol:
chiral ketone \Rightarrow achiral enol \Rightarrow achiral ketone

Synthesis:

- Oxidize alcohol: aldehyde from 1° (Cu, heat); ketone from 2° (H_2CrO_4)
- Grignard:** nitrile (RCN) + $\text{R}'\text{MgX} \Rightarrow \text{RCR}'\text{O}$
- Reduce $\text{RCO}_2\text{R}'$ ($\text{i-Bu}_2\text{AlH}$)
- Reduce RCN ($\text{i-Bu}_2\text{AlH}$)
- Ozonolyze alkene (O_3 , H_2O_2)
- Friedel-Crafts acylation:** $\text{Ar-H} + \text{RCOCl} (\text{AlCl}_3)$

General Reaction:

- Wittig:** form alkene (phosphorous ylide)
- Form Oxime ($>\text{C}=\text{N}-\text{OH}$) (hydroxyl amine)
- Reduce to alcohol (Metal hydride, LiAlH_4)
- Wolff-Kishner:** $>\text{C}=\text{O}$ to $>\text{CH}_2$ (N_2H_4 , base, heat)
- Clemmensen reduction:** $>\text{C}=\text{O}$ to $>\text{CH}_2$ ($\text{Zn}(\text{Hg}), \text{HCl}$)
- Hydrogenate to ROH
(H_2 , metal; NaBH_4 , H^+ ; LiAlH_4 , H^+)
- Oxidize to RCOOH (peroxyacid)
- Form cyanohydrin (HCN)
- Form imine ($>\text{C}=\text{N}-\text{R}$) (1° amine)
- Aldol condensation,
 $>\text{CH}=\text{O} + \text{COOH} \Rightarrow >\text{C}=\text{C}-\text{CH}=\text{O}$
- Nucleophilic attack: $\text{RCHO} + \text{H-Nu} \Rightarrow \text{R-C(OH)-N}$
- Hemiacetal/ketal formation:
 $\text{ROH} + \text{R}'_2\text{C}=\text{O} \Rightarrow \text{R}'_2\text{C}(\text{OH})(\text{OR})$
- Formation of acetal ($\text{R}'\text{OH}, \text{HCl}$)
- Reductive amination: aldehyde or ketone \Rightarrow amine (amine or ammonia, H_2 , Rh)

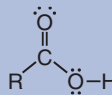
Specific Reaction:

- Acetaldehyde to gem-diol
($\text{H}_2\text{O}, \text{H}^+$ or OH^- catalyst)
- Synthesis of acetaldehyde ($\text{C}_2\text{H}_2, \text{Hg}^{2+}, \text{H}^+, \text{H}_2\text{O}$)
- Oxidize aldehyde to RCOOH : $\text{Ag}_2\text{O}, \text{OH}^-$ or $\text{Ag}(\text{NH}_3)_2^+$; **Tollen's** reagent, ketone is not oxidized
- Haloform, methyl ketone (X_2, OH^-)
- Halogenate -H of ketone (X_2, H^+ or OH^-)

CARBOXYLIC ACID

Properties:

- Organic acid, resonance stabilizes dissociation
- Soluble in water; H-bonding, acid strength given by pKa



Nomenclature:

- Prefix +“oic acid”;
- Examples: HCOOH , **methanoic acid** (formic acid)
 $\text{Me}-\text{COOH}$, **ethanoic acid** (acetic acid),
 $\text{Ar}-\text{COOH}$, **benzoic acid** (benzenecarboxylic acid)
oxalic acid (dicarboxylic acid, $\text{HOOC}-\text{COOH}$)
malonic acid ($\text{HOOC}-\text{CH}_2-\text{COOH}$)
- Fatty acid**, “R” long hydrocarbon (aliphatic) chain

Derivatives:

- Ester
- Amide
- Acid anhydride: RCO-O-CO-R
- Peroxyacid: $\text{R}-\text{CO}_3\text{H}$

Synthesis:

- Oxidize 1° alcohol ($\text{K}_2\text{Cr}_2\text{O}_7, \text{OH}^-$)
- Oxidize aldehyde ($\text{Ag}_2\text{O}, \text{H}^+$)
- Oxidize alkene ($\text{KMnO}_4, \text{OH}^-, \text{heat}, \text{H}^+$)
- Ozonolyze alkene ($\text{O}_3, \text{H}_2\text{O}_2$)
- Hydrolyze nitrile or acyl chloride ($\text{H}^+, \text{H}_2\text{O}$)
- Acid anhydride + water
- Grignard and carbonation ($\text{RMgX} + \text{CO}_2, \text{H}^+$)
- Benzoic acid: oxidize 1°/2° alkylbenzene ($\text{KMnO}_4, \text{OH}^-, \text{heat}, \text{H}^+$)
- From methyl ketone ($\text{Ar}-\text{CO}-\text{CH}_3$) ($\text{X}_2, \text{OH}^-, \text{H}^+$)

Reaction:

- Form acyl chloride ($\text{SOCl}_3, \text{PCl}_3$ or PCl_5)
- Reduce to alcohol (LiAlH_4)
- Neutralize with a base, form a salt
- Esterification: ($\text{R}'\text{OH}, \text{H}^+$)
- Reduce to ketone ($\text{Ba}(\text{OH})_2, \text{heat}$)
- Decarboxylate keto acid to ketone (heat)
- α halo acid: ($\text{X}_2, \text{P}, \text{H}_2\text{O}$); HVZ (Hell-Volhard-Zelinski)
- α hydroxy acid from halo acid (OH^-, H^+)
- α amino acid from halo acid (NH_3 or amine)

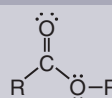
Carbonic acid and derivatives:

- Carbonic acid: H_2CO_3 or $\text{HO}-\text{CO}-\text{OH}$
- Carbonyl dichloride (phosgene), $\text{Cl}-\text{CO}-\text{Cl}$; toxic gas
- Phosgene + $\text{EtOH} \Rightarrow$ diethylcarbonate, $\text{EtO}-\text{CO}-\text{OEt}$
- Phosgene + $\text{NH}_3 \Rightarrow \text{H}_2\text{N}-\text{CO}-\text{NH}_2$ (urea)
- Phosgene + $\text{ROH} \Rightarrow \text{RO}-\text{CO}-\text{Cl}$ (alkyl chloroformate)
- $\text{RO}-\text{CO}-\text{Cl} + \text{RNH}_2 \Rightarrow \text{RO}-\text{CO}-\text{NHR}$ (urethane, carbamates)

ESTER

Properties:

- Derive from carboxylic acid; polar, weak H-bonding; pleasant or fruity odor



Nomenclature:

- Denote “alcohol” component with “-yl” suffix, acid with “-oate” or “-ate” suffix.
- Examples: $\text{Me}-\text{CO}-\text{O-Eth}$, ethyl acetate (ethanol+acetic acid);
- Lactone: cyclic ester

Synthesis:

- Esterification: $\text{ROH}+\text{R}'\text{COOH} \Rightarrow \text{R}'\text{COOR}$ (acid)
- Acid chloride (RCOCl) + $\text{R}'\text{OH}$
- $\text{R}-\text{CN} + \text{R}'\text{OH} (\text{H}^+)$
- Acid anhydride + alcohol \Rightarrow ester + carboxylic acid
- Aromatic ester: phenol + carboxylic anhydride
- β -keto ester: Claisen condensation from ethyl acetate (NaOEt, HCl)
- Transesterification:
 $\text{R}'\text{COOR} + \text{R}''\text{OH} \Rightarrow \text{R}'\text{COOR}'' + \text{ROH} (\text{H}^+, \text{heat})$

ESTER continued

Reaction:

- Acid-catalyzed hydrolysis
- Saponification: base-catalyzed hydrolysis
- Three fatty acids + ethylene glycol \Rightarrow triglyceride
- Grignard to 3° alcohol ($\text{R}'\text{MgX} + \text{R}-\text{COOR}'$)
- Reduce to 1° alcohol (H_2, Ni)
- Form amide ($\text{RCOOR} + 1^\circ/2^\circ$ amine)
- Pyrolyze to alkene and carboxylic acid
- Lactone:** Cyclic ester
 - Intramolecular esterification of δ -hydroxy acid (H^+)
 - Hydrolyze δ/γ lactone to δ/γ hydroxy acid (OH^-, H^+)

AMINE RR'R"N

Properties:

- Substituted ammonia; polar, water soluble; $>\text{N}-\text{H}$ forms H-bonds
- Organic base:** strength denoted by pKb
- Structure:** distorted pyramid (AX_3E)

Nomenclature:

- “R1 R2 R3 amine”
- Example: $\text{Me}-\text{NH}_2$, methyl amine;
 $\text{Ar}-\text{NH}_2$, phenylamine (aniline, amino benzene)

Types of amines:

Primary (1°) amine	Secondary (2°) amine	Tertiary (3°) amine
$\text{R}-\overset{\text{..}}{\underset{\text{H}}{\text{N}}}-\text{H}$	$\text{R}-\overset{\text{..}}{\underset{\text{R}'}{\text{N}}}-\text{H}$	$\text{R}-\overset{\text{..}}{\underset{\text{R}'}{\text{N}}}-\text{R}'$

- Quaternary ammonium salt (4°)
 $\text{NR}'\text{R}''\text{R}'''$ cation (no lone-pair)

Synthesis:

- 1°: aminate haloalkane: $\text{RCH}_2\text{X} + \text{NH}_3$
 - reduce nitrile, $\text{RCN} (\text{LiAlH}_2)$ or (H_2, Ni)
 - reduce nitroalkane, $\text{RNO}_2 (\text{LiAlH}_4)$
 - reduce oxime (Na, EtOH)
 - from aldehyde/ketone (NH_3, H^+)
- 2°: haloalkane + 1° amine
- 3°: haloalkane + 2° amine
- reduce amide ($\text{LiAlH}_4, \text{H}_2\text{O}$)
- aldehyde/ketone + $\text{R}'\text{NH}_2 (\text{H}^+)$
- Aromatic Amine: $\text{Ar}-\text{NO}_2 \Rightarrow \text{Ar}-\text{NH}_2$
(H_2 , catalyst; $\text{Fe}, \text{HCl}, \text{OH}^-$)

Reaction of amine:

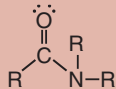
- React as a base: amine + $\text{H}^+ \Rightarrow \text{R}_3\text{NH}^+$
- Nucleophilic N lone-pair
- Amine + sulfonyl chloride \Rightarrow sulfonamide
- amide formation: $1^\circ + \text{R}'\text{COCl} \Rightarrow \text{R}'\text{CO-NHR}$
- $1^\circ + \text{CH}_3\text{COOH} \Rightarrow \text{R}-\text{NO}_2$
- amide formation: $2^\circ + \text{RCOCl}$
- Cope Elimination:** oxidize 3° amine to tertiary ammonium oxide ($\text{R}_3\text{N}^+-\text{O}^-$), heat produces $\text{RHC}=\text{CH}_2$
- Ar- NH_2 : o-p director, electrophilic aromatic substitution
- Ar- NH_2 : nucleophilic aromatic substitution:
 - Step 1: $\text{Ar}-\text{NH}_2 + \text{H}_2\text{N}-\text{NO}_2 \Rightarrow \text{Ar}-\text{N}_2^+$ (diazonium salt, unstable)
 - Step 2: Depends on substitution:
 - $+ \text{Cu}_2\text{O}, \text{Cu}^{2+}, \text{H}_2\text{O} \Rightarrow \text{Ar}-\text{OH}$
 - $+ \text{CuCl} \Rightarrow \text{Ar}-\text{Cl}$
 - $+ \text{CuCN} \Rightarrow \text{Ar}-\text{CN}$
 - $+ \text{H}_3\text{PO}_2 \Rightarrow \text{Ar}-\text{H}$

Hofmann elimination:

- Quaternary ammonium hydroxide \Rightarrow alkene (heat)

AMIDE**Nomenclature:**

- Example: Me-CO-NH₂, acetamide
- Cyclic amide (lactam): N of amide forms ring with β , γ or δ carbon;
 - β forms 4 membered ring; γ forms a 5 membered ring, δ form a 6 membered ring.
- Observed in amino acids

**Synthesis:**

- Nitrile hydrolysis (R-CN + H₂O, conc. H₂SO₄)
- Acyl chloride + 1°/2° amine or ammonia
- Pyrolysis of ammonium salt + RCOOH
- Ammonolysis of ester: 1° or 2° amine + ester
- Polyamide => polypeptide => protein

Reaction:

- Reduce to amine (LiAlH₄)
- Hydrolyze to acid (H₂O, H⁺ or OH⁻)
- Dehydrate to nitrile, RCN (P₄O₁₀, heat)
- Hofmann Reaction: Form 1° amine (NaOBr)
- Grignard (R-MgX) to ketone, R-CO-R~
- Form aldehyde and 2° amine (LiAlH₂ (OEt)₂)
- Nucleophilic substitution; Form R-CO-Nu + amine

AMINO ACID**Properties:**

- Basic (-NH₂) and acidic (-COOH) functionality
- Chiral isomers
- Zwitterion: self-ionization of amino acid to produce COO⁻ and -NH₃⁺
- Isoelectric point, pH which produces equal + and - charges

Nomenclature:

- Common name based on "R" group;
- examples: glycine (-H), alanine (-CH₃)

Synthesis:

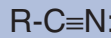
- Gabriel synthesis: RCH₂COOH + Br₂, PCl₃, NH₃

Reaction:

- Lactam formation (cyclic amide)
- Polypeptide formation (peptide bond); dehydration: R-NH₂ and HO-R' moieties
- Protein, amino acid polymer

OTHER NITROGEN-COMPOUNDS**Nitrile:**

example: H₃C-CN; methane nitrile

**Synthesis:**

- Haloalkane + NaCN
- Aldehyde/ketone => cyanohydrin (CN⁻, H⁺)
- Dehydrate amide (P₄O₁₀, heat)

Reaction:

- Hydrolyze to carboxylic acid (acid, heat)
- Hydrolyze to carboxylate (base, heat)
- Reduce to 1° amine (Raney Ni; LiAlH₄)
- Form aldehyde (DIBAL-H (i-Bu₂AlH, H₂O)
- Form ketone (Grignard reagent or R-Li, H⁺)

Imine: >C=N-R

Synthesis: Aldehyde/ketone + 1° amine (H⁺)

Reaction: Intermediate in amination of aldehyde/ketone

NITROGEN continued**Imide:** R-CO-NH-CO-R'**Synthesis:**

- Dehydration, amide + carboxylic acid

Oxime: >C=NOH**Synthesis:**

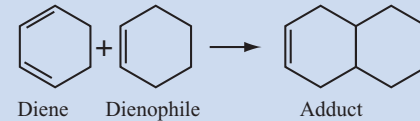
- aldehyde/ketone + hydroxylamine

Reaction:

- oxime to 1° amine (Na, EtOH)

CYCLIZATION: SYNTHESIS OF A CYCLIC COMPOUND**Synthesis:**

- Diels-Alder: diene + dienophile + heat => adduct



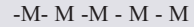
Diene Dienophile Adduct

• Freund-Gustavson: 3-membered ring from 1,3 dihalide (EtOH, Zn, heat)

• [2,2] cycloaddition of alkenes giving cyclobutane adduct (two alkenes, photochemical reaction)

Reaction of cyclic compound:

- Retro-Diels-Alder: thermally decompose cycloalkene
- Reduce aromatic to symmetric 1,4 cycloalkene (Li or Na, EtOH, Liq NH₃) (Birch)
- Small ring is strained, may decompose to linear chain
- Epoxide ring opening reaction

ORGANIC POLYMER

Monomers (M) bond to form a high molecular weight compound.

Factors which influence properties: chain length, branching vs. linear, nature of the monomer, density, interchain bonds, hydrophobic and hydrophilic interactions.

Examples:

- PE (polyethylene)
- PS (polystyrene)
- HDPE (high density polyethylene)
- LDPE (low density polyethylene)
- PET (polyethylene terephthalate)

Synthesis:

- Free-radical synthesis: ethylene => PE; styrene => PS (radical initiation)
- Condensation:
 - HO-R-OH+HO-R'-OH => HO-R-R'-OH + H₂O
 - Example: ethylene glycol and terephthalic acid => PET

Reaction:

- Hydrolysis of polymer (reverse of condensation)
- Cross-link adjacent polymer chains or segments

METAL REACTION**Organometallic:**

- Carbon atom bonded to a metal atom
- Types of bonding:
 - ionic bond, Na,K; R⁻M⁺
 - partial covalent, Mg, Li; R electrophilic character
 - covalent, Pb, Sn, Hg; R-M

Grignard reagent:

- Strong base gives R electrophilic character:
 - Li + R-Br => R-Li
 - RX + Mg => RMgX
 - ArX + Mg => ArMgX

Organoborane:

- Boron hydride, B_nH_m
- example: diborane, B₂H₆

Synthesis:

- Hydroboration: Alkene + Boron hydride syn addition

Reaction:

- Organoborane => alcohol (H₂O₂/OH⁻)
- R-B<- => R-H (acetic acid; addition of H)

Organolithium: R-Li**Synthesis:**

- Li + haloalkane (R-X or Ar-X) (cold, Et₂O)

Organomagnesium: RMgX or ArMgX

- Grignard: RX + Mg (Et₂O); R behaves as R-

Organocupper: R-Cu

- Add R- to C=C of unsaturated carbonyl

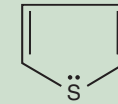
Organolead/mercury:

- Stable compound, VOLATILE AND TOXIC
- Tetraethyl lead (anti-knock agent in gasoline)

SULFUR CHEMISTRY**Sulphur Compounds**

• Thiol:	R-SH
• Sulfide or Thioether:	R-S-S-R'
• Disulfide:	R-S-S-R'
• Thiol ester:	R-CO-SR'
• Sulfoxide:	R-SO ₂ -R'
• Sulfone:	R-SO ₂ -R'
• Thiophenol:	Ar-SH
• Thiotetone:	R-CS-R'
• Sulfonic acid:	R-SO ₃ H
• Sulfenic Acid:	R-SO ₂ H
• Hydrogen sulfate:	R-OSO ₃ H

Thiophene, Heterocyclic sulphur compound

**Synthesis:**

- Thiol: From alkyl bromide/iodide (KOH, H₂S)
- Thiol: RCH₂X + NaSH => RCH₂SH (EtOH, heat)
- Thiol ester: Acyl chloride + thiol
- Alkyl hydrogen sulfate (Alkene + cold conc. H₂SO₄)
- Thiol: Alkene + H₂S (H₂SO₄, heat) (Markovnikoff addition)
- Thiol: Alkene+H₂S (peroxide or UV) (Anti-Markovnikoff addition)

Reaction:

- Form sulfide from thiol (NaOH, R'CH₂X)
- Form disulfide from thiol (I₂ or H₂O₂)
- Oxidize thiol to form sulfonic acid, RSO₃H, (HNO₃)
- Desulfurization of thiol to alkane (H₂, Ni)
- Sulfonate benzene (SO₃, conc. H₂SO₄)

ISBN-13: 978-1-42320285-1

ISBN-10: 142320285-6

5 0 4 9 5

CREDITS

Author: Mark Jackson, PhD.

PRICE
U.S.\$4.95

Layout: Andre Brisson

CAN.\$7.50

Note: Due to the condensed nature of this chart, use as a quick reference guide, not as a replacement for assigned course work. The reaction reagents are not for illustrative purposes only; this should not serve as guide for lab experiment procedures.

All rights reserved. No part of this publication may be reproduced or transmitted in any form, or by any means, electronic or mechanical, including photocopy, recording, or any information storage and retrieval system, without written permission from the publisher.

©2001 BarCharts, Inc. 1106

Customer Hotline # 1.800.230.9522

hundreds of titles at
quickstudy.com



PRINTED WITH
SOYINK