

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.

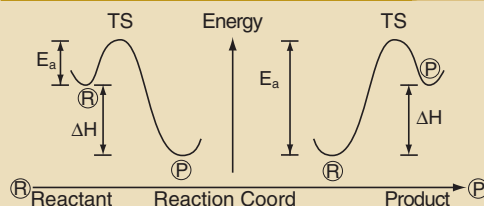
Major reaction types:

- 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-Kirschner, Clemmensen:** 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.

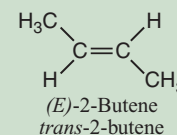
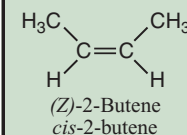
ALKENE $>C=C<$

Properties: Similar to alkane; non-polar, flammable

Nomenclature:

- Add -ene to prefix; Use # to denote C=C position
- **Isolated** C-C=C-C-C=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)
- Alkatriline, 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 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)
- Oxymercuration-demercuration 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^-)
- Ozonolyse 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

ALKANE



Properties:

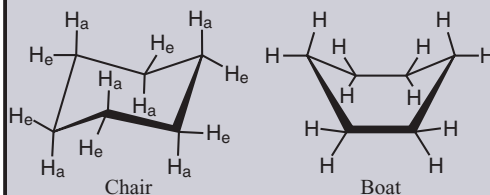
- Hydrocarbon
- Weak intermolecular forces
- Non-cyclic: general formula C_nH_{2n+2}
- Tetrahedral C-C-C (109°)

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:

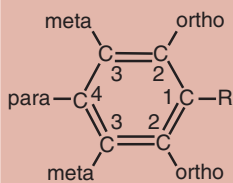
- Combustion: alkane + $O_2 \Rightarrow CO_2 + H_2O$
- Halogenation to haloalkane (Cl_2/Br_2 , light or heat)

BENZENE/ARENE

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 (carbolic 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:

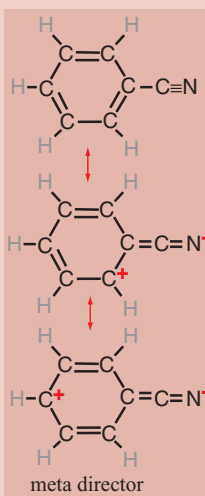
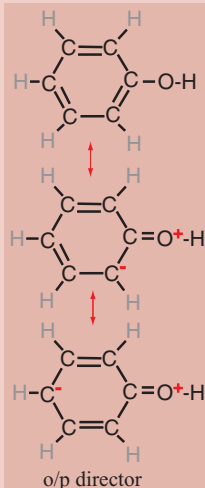
- Combustion (similar to alkane)
- Birch reduction $\Rightarrow$ 1,4 cyclohexadiene (Na, NH₃, EtOH)
- Hydrogenate to cyclohexane (H_2 , Pt)

Electrophilic substitution:

- Alkylation: Ar-R (*Friedels-Craft*, RCl, AlCl₃)
- Nitration: Ar-NO₂ (HNO₃, H₂SO₄)
- Halogenation: Ar-Br (Br₂, FeBr₃)
- Ar-Cl (Cl₂, FeCl₃) Ar-I (I₂, HNO₃)
- Acylation: Ar-CR=O (RCOCl, AlCl₃)
- Sulfonation: Ar-SO₃H (SO₃, H₂SO₄)

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: (KMnO₄, OH⁻, heat, H⁺)
- Chlorinate -Me of Toluene (Cl₂)

ALKYNE -C≡C-**Properties:**

- Hydrocarbon, at least 1 C $\equiv$ 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) C₂H₂

Synthesis:

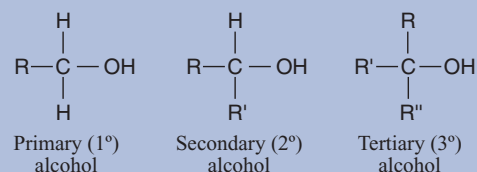
- CaC₂ + H₂O $\Rightarrow$ Ca(OH)₂ + C₂H₂
- Dehydrohalogenate vic-haloalkene (NaNH₂, liq NH₃)
- Alkylate terminal alkyne (NaNH₂, liq NH₃; R-X)

Reaction:

- Addition: hydrogenate to alkane (H₂, Pt or Ni)
 - syn to cis/Z alkene (H₂/Ni₂B P-2 catalyst)
 - anti to trans/E alkene (Li, Liq NH₃)
- haloalkene to gem-dihalide (HX) (Markovnikov)
- halogenate to haloalkene or haloalkane (X₂)
- Ozonolyze to carboxylic acid (O₃, H₂O)
- Oxidize to carboxylic acid (KMnO₄, 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 (H₂O, H⁺)
- Hydroborate/oxidize alkene (THF:BH₃; H₂O₂, OH⁻)
- Hydrogenate aldehyde (H₂/Ni or Pt catalyst)
- Hydrolyze 1o alkyl halide (water, OH⁻)
- Reduce aldehyde, ester, ketone or carboxylic acid
- Ethanol: Ferment sugar or starch
- Methanol: CO + H₂, catalyst; Pyrolyze cellulose
- Hydrolyze ester (water, acid)
- Dehydrate ether (H₂SO₄, low heat)
- Grignard (RMgX): formaldehyde $\Rightarrow$ 1° alcohol; aldehyde $\Rightarrow$ 2° alcohol; ketone $\Rightarrow$ 3° alcohol
- Synthesis of Glycol from Ketone/aldehyde: (HIO₄ or Pb(OAc)₄; H₂SO₄ + heat)
- oxidize alkene: (KMnO₄: cis) (H₂O₂, formic acid: trans)
- hydrolyze epoxide (H₂O, H₂SO₄)

Reaction:

- Oxidize 1° to aldehyde (CuO, heat) or 2° to ketone (KMnO₄, H⁺)
- Oxidize 1° to carboxylic acid (KMnO₄, H⁺)
- Dehydrate to alkene; Zaitsev's rule; rate 3° > 2° > 1° (hot H₂SO₄, or Al₂O₃)
- Dehydrate to ether (H₂SO₄, lower temperature)
- Oxidize to ketone (2° alcohol) (H₂CrO₂)
- 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
- Benzendiols, HO-Ar-OH; para, hydroquinone; ortho, catechol; meta, resorcinol

Reaction of Phenol:

- Electrophilic substitution: o-p director
- Hydrogenate to cyclohexane (H₂, catalyst)
- Form ester (acid anhydride or acid chloride)

Synthesis of Phenol:

- Electrophilic aromatic substitution
- Williamson reaction, phenyl ether (NaOH, RCH₂X)
- Arenediazonium salt intermediate: Ar-NH₂ + HONO $\Rightarrow$ Ar-N₂⁺ + Cu₂O, H₂O $\Rightarrow$ 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/ARENE 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° alkyl halide to alcohol (RX + OH⁻)

HALOHYDRIN X-R-R'-OH

Synthesis: Alkene + X₂, H₂O

Reaction:

- Halohydrin + ROH $\Rightarrow$ β hydroxy ether
- Halohydrin + RNH₂ $\Rightarrow$ 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° alcohol (H₂SO₄, 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° alcohol

ALDEHYDE & KETONE >C=O**Properties:**

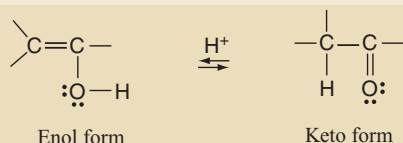
- Polar $>C^+=O$; low molecular weight are water-soluble
- Main chemical difference: ketone is harder to oxidize than aldehyde.

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 $>C=O$ groups
- Acyl group: $RC=O$ or $Ar-C=O$
- Ketene: $C=C=O$
- Ketal: $RR'C(OR)(OR)$;
- Acetal: $RHC(OR)(OR)$
- Hemiacetal: $RHC(OH)(OR)$
- Diketone: $R'-CO-CH_2-CO-R$

Keto-enol tautomerism:

- Nucleophile attacks $>C=C<$ of enol-form
- Acidic α -H, $-CH^*-CHO$ 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) + $R'MgX \Rightarrow RCR'O$
- Reduce RCO_2R' (i-Bu₂AlH)
- Reduce RCN (i-Bu₂AlH)
- Ozonolyze alkene (O_3 , H_2O_2)
- Friedel-Craft acylation: $Ar-H + RCOCl (AlCl_3)$

General Reaction:

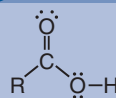
- Wittig, form alkene (phosphorous ylide)
- Form Oxime ($>C=N-OH$) (hydroxyl amine)
- Reduce to alcohol (Metal hydride, $LiAlH_4$)
- Wolff-Kishner: $>C=O$ to $>CH_2$ (N_2H_4 , base, heat)
- Clemmenson reduction, $>C=O$ to $>CH_2$ ($Zn(Hg)$, HCl)
- Hydrogenate to ROH (H_2 , metal; $NaBH_4$, H^+ ; $LiAlH_4$, H^+)
- Oxidize to RCOOH (peroxyacid)
- Form cyanohydrin (HCN)
- Form imine ($>C=N-R$) (1° amine)
- Aldol condensation, $>CH=O + COOH \Rightarrow >C=C-CH=O$
- Nucleophilic attack: $RCHO + H-Nu \Rightarrow R-C(OH)-Nu$
- Hemiacetal/ketal formation: $ROH + R'_2C=O \Rightarrow R'_2C(OH)(OR)$
- Formation of acetal ($R'OH$, HCl)
- Reductive amination: aldehyde or ketone $\Rightarrow$ amine (amine or ammonia, H_2 , Rh)

Specific Reaction:

- Acetaldehyde to gem-diol (H_2O , H^+ or OH^- catalyst)
- Synthesis of acetaldehyde (C_2H_2 , Hg^{2+} , H^+ , H_2O)
- Oxidize aldehyde to RCOOH: Ag_2O, OH^- or $Ag(NH_3)_2^+$; 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), Me-COOH, **ethanoic acid** (acetic acid), Ar-COOH, **benzoic acid** (benzenecarboxylic acid)
- **oxalic acid** (dicarboxylic acid, HOOC-COOH)
- **malonic acid** (HOOC-CH₂-COOH)
- **Fatty acid**, "R" long hydrocarbon (aliphatic) chain

Derivatives:

- Ester
- Amide
- Acid anhydride: $RCO-O-CO-R$
- Peroxyacid: $R-CO_3H$
- Acyl chloride
- Amino acid

Synthesis:

- Oxidize 1° alcohol ($K_2Cr_2O_7$, OH^-)
- Oxidize aldehyde (Ag_2O , H^+)
- Oxidize alkene ($KMnO_4$, OH^- , heat, H^+)
- Ozonolyze alkene (O_3 , H_2O_2)
- Hydrolyze nitrile or acyl chloride (H^+ , H_2O)
- Acid anhydride + water
- Grignard and carbonylation ($RMgX + CO_2$, H^+)
- Benzoic acid: oxidize $1^\circ/2^\circ$ alkylbenzene ($KMnO_4, OH^-$, heat, H^+)
- From methyl ketone ($Ar-CO-CH_3$) (X_2 , OH^- , H^+)

Reaction:

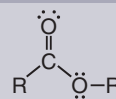
- Form acyl chloride ($SOCl_2$, PCl_3 or PCl_5)
- Reduce to alcohol ($LiAlH_4$)
- Neutralize with a base, form a salt
- Esterification: ($R'OH$, H^+)
- Reduce to ketone ($Ba(OH)_2$, heat)
- Decarboxylate keto acid to ketone (heat)
- α halo acid: (X_2 , P, H_2O): HVZ (Hell-Volhard-Zelinski)
- α hydroxy 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 $HO-CO-OH$
- Carbonyl dichloride (phosgene), $Cl-CO-Cl$; toxic gas
- Phosgene + EtOH $\Rightarrow$ diethylcarbonate, $EtO-CO-OEt$
- Phosgene + $NH_3 \Rightarrow H_2N-CO-NH_2$ (urea)
- Phosgene + ROH $\Rightarrow RO-CO-Cl$ (alkyl chloroformate)
- $RO-CO-Cl + RNH_2 \Rightarrow RO-CO-NHR$ (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: Me-CO-O-Eth, ethyl acetate (ethanol+acetic acid);
- Lactone: cyclic ester

Synthesis:

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

ESTER continued**Reaction:**

- Acid-catalyzed hydrolysis
- Saponification: base-catalyzed hydrolysis
- Three fatty acids + ethylene glycol $\Rightarrow$ triglyceride
- Grignard to 3° alcohol ($R''MgX + R-COOR'$)
- Reduce to 1° alcohol (H_2 , Ni)
- Form amide ($RCOOR + 1^\circ/2^\circ$ amine)
- Pyrolyze to alkene and carboxylic acid

Lactone: Cyclic ester

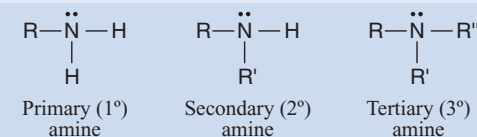
- Intramolecular esterification of δ -hydroxy acid (H^+)
- Hydrolyze δ/γ lactone to δ/γ hydroxy acid (OH^- , H^+)

AMINE**RR'R''N****Properties:**

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

Nomenclature:

- "R1 R2 R3 amine"
- Example: Me-NH₂, methyl amine; Ar-NH₂, phenylamine (aniline, amino benzene)

Types of amines:

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

Synthesis:

- **1°:** amine haloalkane: $RCH_2X + NH_3$
 - reduce nitrile, RCN ($LiAlH_2$) or (H_2 , Ni)
 - reduce nitroalkane, RNO_2 ($LiAlH_4$)
 - reduce oxime (Na, EtOH)
 - from aldehyde/ketone (NH_3 , H^+)
- **2°:** haloalkane + 1° amine
- aldehyde/ketone + $R'NH_2$ (H^+)
- **3°:** haloalkane + 2° amine
- reduce amide ($LiAlH_4$, H_2O)
- aldehyde/ketone + $R'R''NH$ (H^+)
- Aromatic Amine: $Ar-NO_2 \Rightarrow Ar-NH_2$ (H_2 , catalyst; Fe, HCl, OH^-)

Reaction of amine:

- React as a base: amine + $H^+ \Rightarrow R_3NH^+$
- Nucleophilic N lone-pair
- Amine + sulfonyl chloride $\Rightarrow$ sulfonamide
- amide formation: $1^\circ + R'COCl \Rightarrow R'CO-NHR$
- $1^\circ + CH_3COOOH \Rightarrow R-NO_2$
- amide formation: $2^\circ + RCOCl$
- **Cope Elimination:** oxidize 3° amine to tertiary ammonium oxide ($R_3N^+-O^-$), heat produces $RHC=CH_2$
- Ar-NH₂: o-p director, electrophilic aromatic substitution
- Ar-NH₂: nucleophilic aromatic substitution:
 - Step 1: $Ar-NH_2 + cold\ nitrous\ acid \Rightarrow Ar-N_2^+$ (diazonium salt, unstable)
 - Step 2: Depends on substitution:
 - + Cu_2O , Cu^{2+} , $H_2O \Rightarrow Ar-OH$
 - + $CuCl \Rightarrow Ar-Cl$
 - + $CuCN \Rightarrow Ar-CN$
 - + $H_3PO_2 \Rightarrow Ar-H$

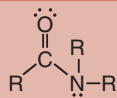
Hofmann elimination:

- 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



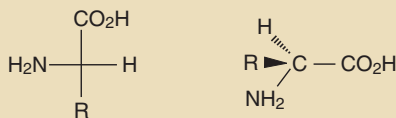
Synthesis:

- Nitrile hydrolysis ($R-CN + H_2O$, conc. H_2SO_4)
- Acyl chloride + 1°/2° amine or ammonia
- Pyrolysis of ammonium salt + $RCOOH$
- Ammonolysis of ester: 1° or 2° amine + ester
- Polyamide $\Rightarrow$ polypeptide $\Rightarrow$ protein

Reaction:

- Reduce to amine ($LiAlH_4$)
- Hydrolyze to acid (H_2O , H^+ or OH^-)
- Dehydrate to nitrile, RCN (P_4O_{10} , heat)
- *Hofmann* Reaction: Form 1° amine ($NaOBr$)
- *Grignard* ($R^{\sim}MgX$) to ketone, $R-CO-R^{\sim}$
- Form aldehyde and 2° amine ($LiAlH_2(OEt)_2$)
- Nucleophilic substitution; Form $R-CO-Nu$ + amine

AMINO ACID



Properties:

- Basic ($-NH_2$) and acidic ($-COOH$) functionality
- Chiral isomers
- **Zwitterion:** self-ionization of amino acid to produce COO^- and $-NH_3^+$
- Isoelectric point, pH which produces equal + and - charges

Nomenclature:

- Common name based on "R" group; examples: glycine ($-H$), alanine ($-CH_3$)

Synthesis:

- Gabriel synthesis: $RCH_2COOH + Br_2, PCl_3, NH_3$

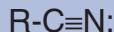
Reaction:

- Lactam formation (cyclic amide)
- Polypeptide formation (peptide bond); dehydration: $R-NH_2$ and $HO-R^{\sim}$ moieties
- Protein, amino acid polymer

OTHER NITROGEN-COMPOUNDS

Nitrile:

example: H_3C-CN ; methane nitrile



Synthesis:

- Haloalkane + $NaCN$
- Aldehyde/ketone $\Rightarrow$ cyanohydrin (CN^- , H^+)
- Dehydrate amide (P_4O_{10} , heat)

Reaction:

- Hydrolyze to carboxylic acid (acid, heat)
- Hydrolyze to carboxylate (base, heat)
- Reduce to 1° amine (Raney Ni; $LiAlH_4$)
- Form aldehyde (DIBAL-H ($i-Bu_2AlH$), H_2O)
- Form ketone (*Grignard* reagent or $R-Li$, H^+)

Imine:



Synthesis:

Aldehyde/ketone + 1° amine (H^+)

Reaction:

Intermediate in amination of aldehyde/ketone

NITROGEN continued

Imide:



Synthesis:

- Dehydration, amide + carboxylic acid

Oxime:



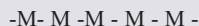
Synthesis:

- aldehyde/ketone + hydroxylamine

Reaction:

- oxime to 1° amine (Na , $EtOH$)

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 $\Rightarrow$ PE; styrene $\Rightarrow$ PS (radical initiation)
- Condensation:
 - $HO-R-OH + HO-R^{\sim}-OH \Rightarrow HO-R-R^{\sim}-OH + H_2O$
 - Example: ethylene glycol and terephthalic acid $\Rightarrow$ PET

Reaction:

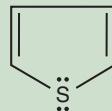
- Hydrolysis of polymer (reverse of condensation)
- Cross-link adjacent polymer chains or segments

SULFUR CHEMISTRY

Sulphur Compounds

- Thiol: $R-SH$
- Sulfide or Thioether: $R-S-R^{\sim}$
- Disulfide: $R-S-S-R^{\sim}$
- Thiol ester: $R-CO-SR^{\sim}$
- Sulfoxide: $R-S(=O)-R^{\sim}$
- Sulfone: $R-SO_2-R^{\sim}$
- Thiophenol: $Ar-SH$
- Thioketone: $R-CS-R^{\sim}$
- Sulfonic acid: $R-SO_3H$
- Sulfinic Acid: $R-SO_2H$
- Hydrogen sulfate: $R-OSO_3H$

Thiophene, Heterocyclic sulphur compound



Synthesis:

- Thiol: From alkyl bromide/iodide (KOH , H_2S)
- Thiol: $RCH_2X + NaSH \Rightarrow RCH_2SH$ ($EtOH$, heat)
- Thiol ester: Acyl chloride + thiol
- Alkyl hydrogen sulfate (Alkene + cold conc. H_2SO_4)
- Thiol: Alkene + H_2S (H_2SO_4 , heat) (Markovnikov addition)
- Thiol: Alkene + H_2S (peroxide or UV) (Anti-Markovnikov addition)

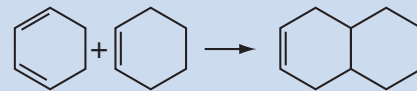
Reaction:

- Form sulfide from thiol ($NaOH$, $R^{\sim}CH_2X$)
- Form disulfide from thiol (I_2 or H_2O_2)
- Oxidize thiol to form sulfonic acid, RSO_3H , (HNO_3)
- Desulfurization of thiol to alkane (H_2 , Ni)
- Sulfonate benzene (SO_3 , conc. H_2SO_4)

CYCLIZATION: SYNTHESIS OF A CYCLIC COMPOUND

Synthesis:

- Diels-Alder: diene + dienophile + heat $\Rightarrow$ 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 , $EtOH$, $Liq NH_3$) (Birch)
- Small ring is strained, may decompose to linear chain
- Epoxide ring opening reaction

METAL REACTION

Organometallic:

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

Grignard reagent:

- Strong base gives R electrophilic character:
 - $Li + R-Br \Rightarrow R-Li$
 - $RX + Mg \Rightarrow RMgX$
 - $ArX + Mg \Rightarrow ArMgX$

Organoborane:

- Boron hydride, B_nH_m
example: diborane, B_2H_6
- **Synthesis:**
 - Hydroboration: Alkene + Boron hydride syn addition
- **Reaction:**
 - Organoborane $\Rightarrow$ alcohol (H_2O_2/OH^-)
 - $R-B < \Rightarrow R-H$ (acetic acid; addition of H)

Organolithium: $R-Li$

- **Synthesis:**
 - $Li + haloalkane (R-X \text{ or } Ar-X)$ (cold, Et_2O)

Organomagnesium: $RMgX$ or $ArMgX$

- *Grignard*: $RX + Mg$ (Et_2O); R behaves as $R^{\sim}$

Organocopper: $R-Cu$

- Add R - to $C=C$ of unsaturated carbonyl

Organolead/mercury:

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

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CREDITS

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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 noted for illustrative purposes only; this should not serve as guide for lab experiment procedures.

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