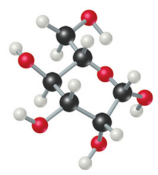
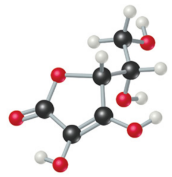
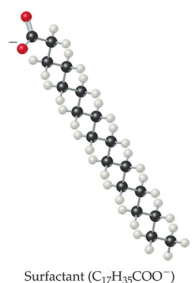
Chapter 25 Organic and Biological Chemistry



Glucose ($C_6H_{12}O_6$)



Ascorbic acid (HC₆H₇O₆)

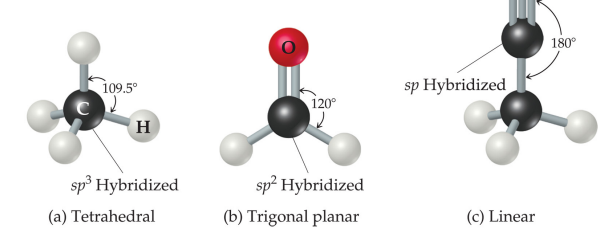


Organic Chemistry

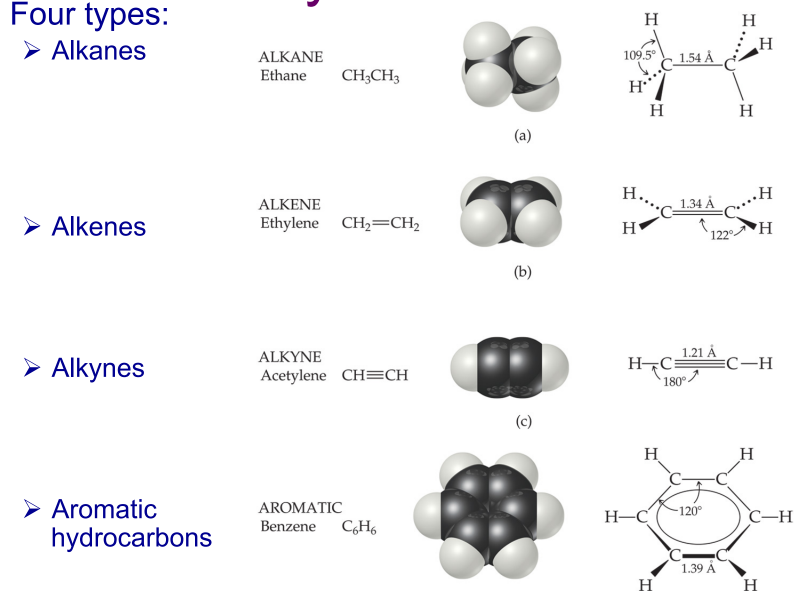
- The chemistry of carbon compounds.
- What's special about carbon?
 - tetravalent (sp³ hybridization)
 - wide choice in oxidation states
 - ≻ CO₂ C, +4
 - ≻ CH₄ C, -4
 - bonds well to O,N,halides,itself,etc.
 - Covalent bonds are very strong

Structure of Carbon Compounds

- There are three hybridization states and geometries found in organic compounds:
 - ≻ sp³ Tetrahedral
 - > sp² Trigonal planar
 - ≻ sp Linear



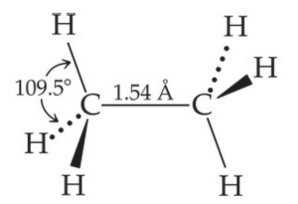
Hydrocarbons



Alkanes

ALKANE Ethane CH₃CH₃

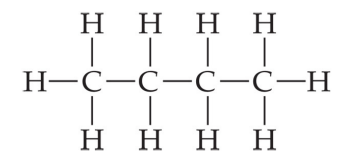


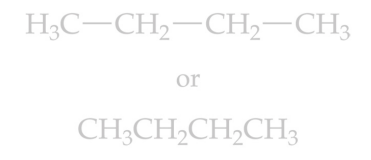


- Only single bonds.
- Saturated hydrocarbons.
 - \succ "Saturated" with hydrogens.

Formulas

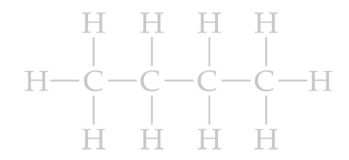
- Lewis structures of alkanes look like this.
- Also called structural formulas.
- Often not convenient, though...

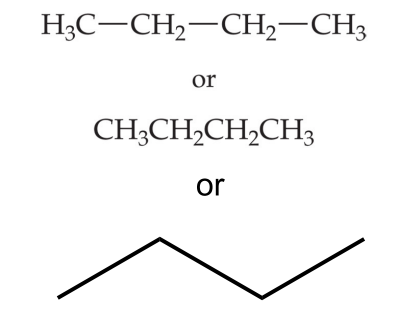




Formulas

...so more often condensed formulas are used.





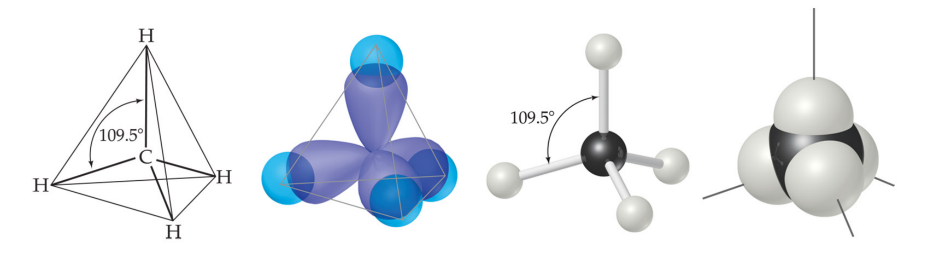
Note: always 4 bonds to Carbon.

Properties of Alkanes

Molecular Formula	Condensed Structural Formula	Name	Boiling Point (°C)
CH_4	CH_4	Methane	-161
C_2H_6	CH ₃ CH ₃	Ethane	-89
C_3H_8	$CH_3CH_2CH_3$	Propane	-44
$C_{4}H_{10}$	$CH_3CH_2CH_2CH_3$	Butane	-0.5
$C_{5}H_{12}$	$CH_3CH_2CH_2CH_3$	Pentane	36
$C_{6}H_{14}$	$CH_3CH_2CH_2CH_2CH_3$	Hexane	68
$C_{7}H_{16}$	CH ₃ CH ₂ CH ₂ CH ₂ CH ₂ CH ₂ CH ₃	Heptane	98
C_8H_{18}	CH ₃ CH ₂ CH ₂ CH ₂ CH ₂ CH ₂ CH ₂ CH ₃	Octane	125
C_9H_{20}	CH ₃ CH ₂ CH ₃	Nonane	151
$C_{10}H_{22}$	CH ₃ CH ₂	Decane	174

- Only van der Waals force: London force.
- Boiling point increases with length of chain.

Structure of Alkanes

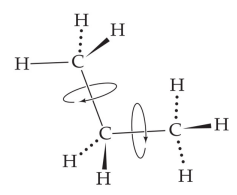


- Carbons in alkanes *sp*³ hybrids.
- Tetrahedral geometry.
- 109.5° bond angles.

Structure of Alkanes



- Only *o*-bonds in alkanes
- Free rotation about C—C bonds.



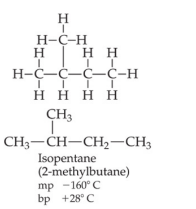


 $\begin{array}{ccccccc} H & H & H & H \\ H - C - C & - C - C - C - H \\ H & H & H & H \\ C H_3 C H_2 C H_2 C H_2 C H_3 \\ & Butane \\ mp & -135^{\circ} C \\ bp & -0.5^{\circ} C \end{array}$











H-Ċ-H

H-Ċ-H

Ĥ

 $CH_3 - C - CH_3$ (2,2-dimethylpropane)

mp −20° C

bp +9°C

Η

Η

Ċ–Н

Η

Η

H-C

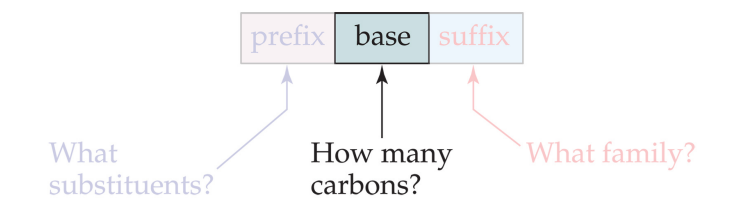
CH₃

CH₃

Have same molecular formulas, but atoms are bonded in different order.

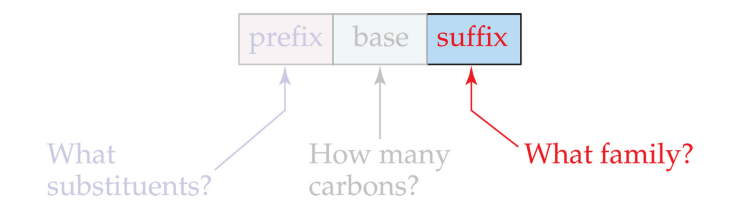
Organic Nomenclature

- Three parts to a compound name:
 - Base: Tells how many carbons are in the longest continuous chain.



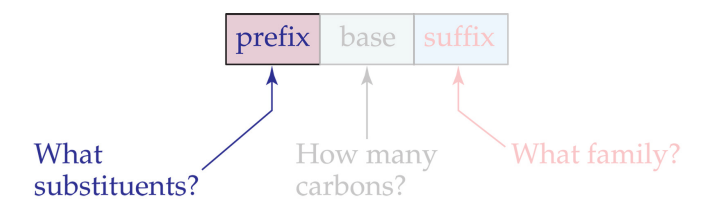
Organic Nomenclature

- Three parts to a compound name:
 - Base: Tells how many carbons are in the longest continuous chain.
 - > Suffix: Tells what type of compound it is.

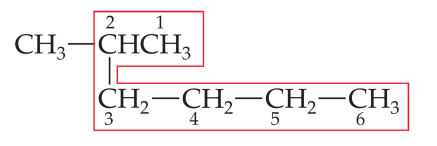


Organic Nomenclature

- Three parts to a compound name:
 - Base: Tells how many carbons are in the longest continuous chain.
 - > Suffix: Tells what type of compound it is.
 - Prefix: Tells what groups are attached to chain.



To Name a Compound...

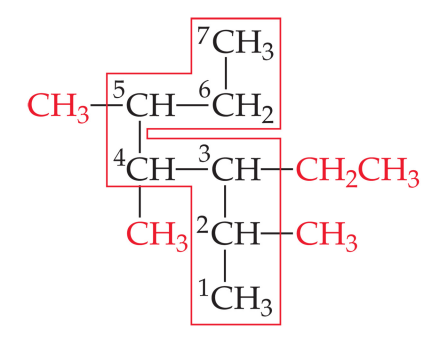


2-Methylhexane

Group	Name	
$CH_3 - CH_3CH_2 - CH_3CH_2CH_2 - CH_3CH_2CH_2 - CH_3CH_2CH_2CH_2 - CH_3CH_2CH_2CH_2 - CH_2CH_2 - CH_2CH_2 - CH_3CH_2CH_2 - CH_3CH_2CH_2CH_2 - CH_3CH_2CH_2 - CH_3CH_2CH_2CH_2 - CH_3CH_2CH_2CH_2CH_2 - CH_3CH_2CH_2CH_2CH_2CH_2CH_2CH_2CH_2CH_2CH_2$	Methyl Ethyl Propyl Butyl	
CH ₃ HC— CH ₃	Isopropyl	
$CH_{3} - CH_{3} - CH_{3} - CH_{3}$	<i>tert-</i> Butyl	

- 1. Find the longest chain in the molecule.
- 2. Number the chain from the end nearest the first substituent encountered.
- 3. List the substituents as a prefix along with the number(s) of the carbon(s) to which they are attached.

To Name a Compound...

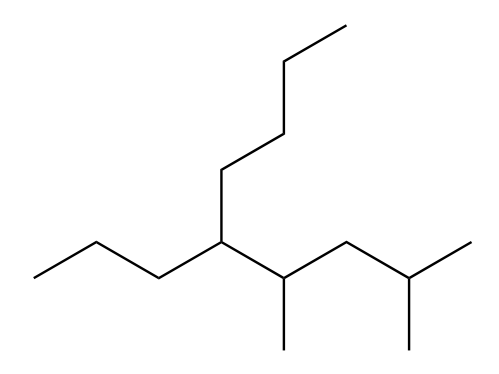


3-Ethyl-2,4,5-trimethylheptane

list substiutents alphabetically.

More than one longest chain: pick the one with the most substituents.

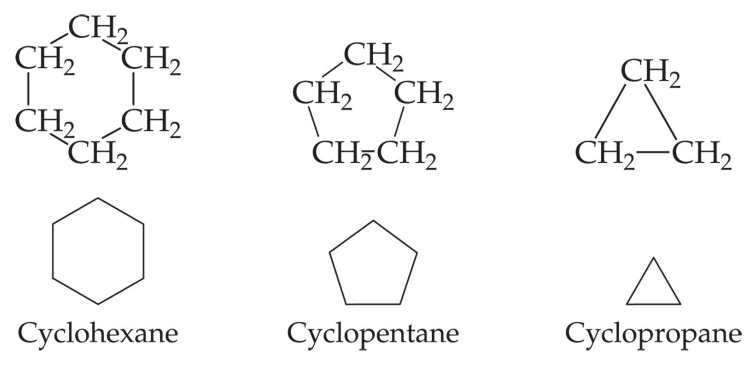
The most substituents:



2,4-dimethyl-5-propylnonane

Cycloalkanes

- Carbon can also form ringed structures.
- Five- and six-membered rings are most stable.
 - Can take on conformation in which angles are very close to tetrahedral angle.
 - > Smaller rings are quite strained.



Reactions of Alkanes

- Rather unreactive due to presence of only C —C and C—H σ -bonds.
- Therefore, great nonpolar solvents.
- General rule of organic chemistry;
 - reactivity comes from the functional groups, ie. the part of the molecule that is not a straight alkane.
 - different functional groups give rise to different kinds of activity.

- EXAM 4
- •
- 2 VB
- 2 color wheel
- 6 xtal field
- 2 isomers
- 1 magnetism
- 1 oxid state
- 1 organic
- 1 organic naming
- 1 intermolecular forces
- 1 MO
- •
- ۲

Alkenes



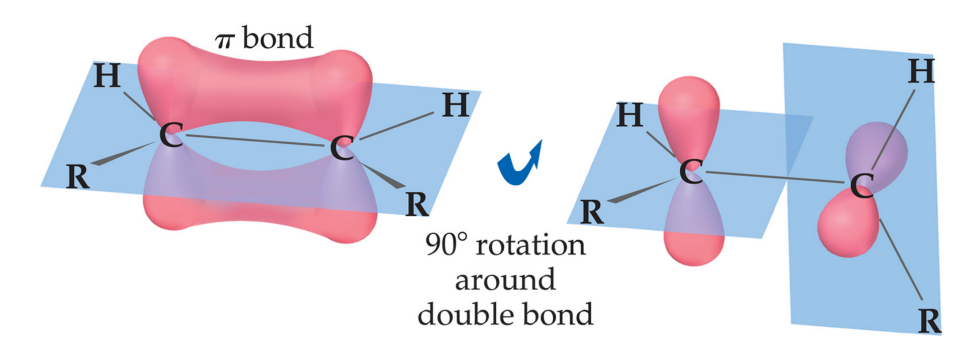
- Contain at least one carbon-carbon double bond.
- Unsaturated.

> Have fewer than maximum number of hydrogens.

Structure of Alkenes

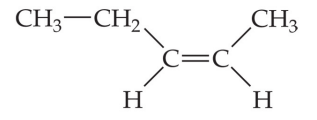
• Unlike alkanes, alkenes cannot rotate freely about the double bond.

> Side-to-side overlap makes this impossible without breaking π -bond.

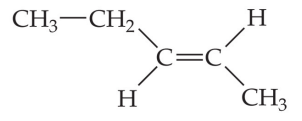


Structure of Alkenes

This creates geometric isomers, which differ from each other in the spatial arrangement of groups about the double bond.

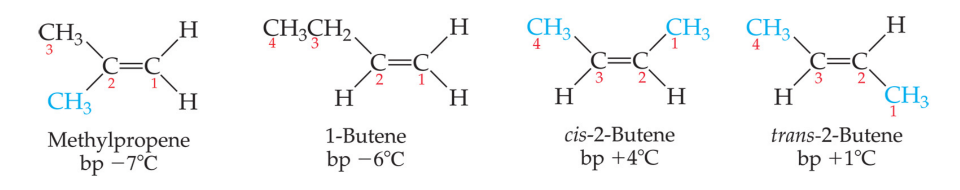


cis-2-Pentene



trans-2-Pentene

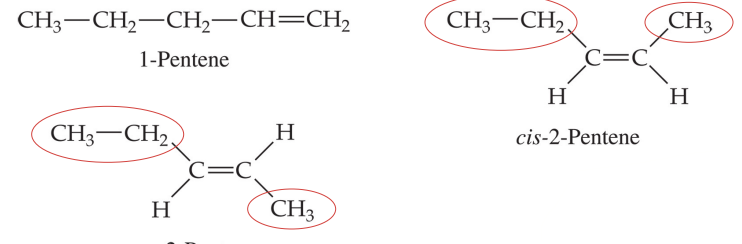
Properties of Alkenes



Structure also affects physical properties of alkenes.

Nomenclature of Alkenes

- Chain numbered so double bond gets smallest possible number.
- cis- alkenes have carbons in chain on same side of molecule.
- *trans* alkenes have carbons in chain on opposite side of molecule.



trans-2-Pentene

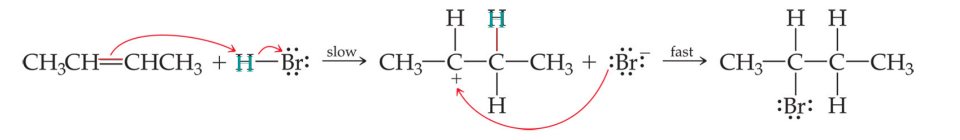
Reactions of Alkenes

$H_2C = CH_2 + \frac{Br_2}{H_2} \longrightarrow H_2C - CH_2$ $| | | \\Br Br$

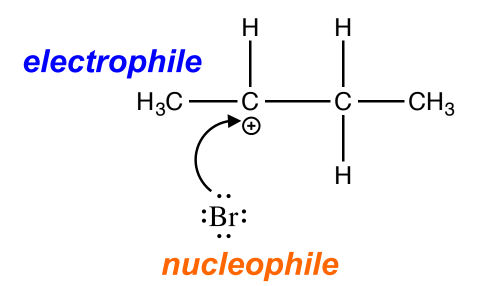
- Addition Reactions
 - Two atoms (e.g., bromine) add across the double bond.
 - > One π -bond and one σ -bond are replaced by two σ -bonds; therefore, ΔH is negative.

"Arrow pushing

• The organic chemist's language of reaction mechanism



- The basics of arrow pushing:
 - Arrow goes from where electrons come from to where they are going.
 - Double headed arrow indicates 2 electrons
 - Single headed arrow indicates 1 electron
- Alkene addition two-step mechanism:
 - First step is slow, rate-determining step.
 - Second step is fast.

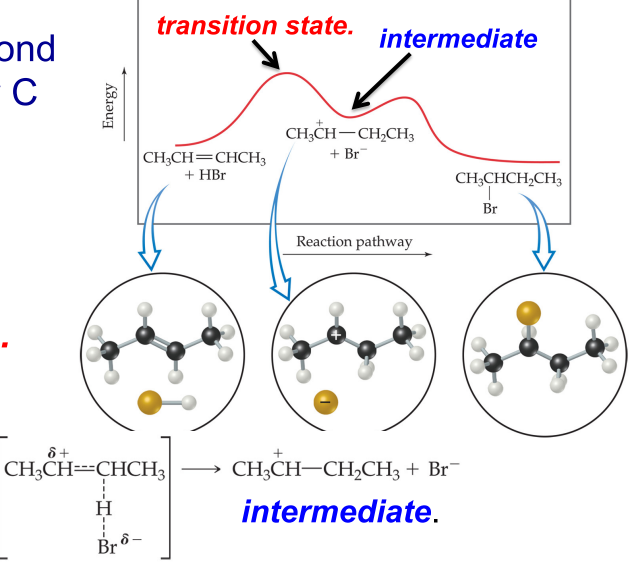


- The basics of arrow pushing:
 - Arrow goes from where electrons come from to where they are going.
 - Double headed arrow indicates 2 electrons
 - Single headed arrow indicates 1 electron
 - Arrow goes from *nucleophile* to electrophile

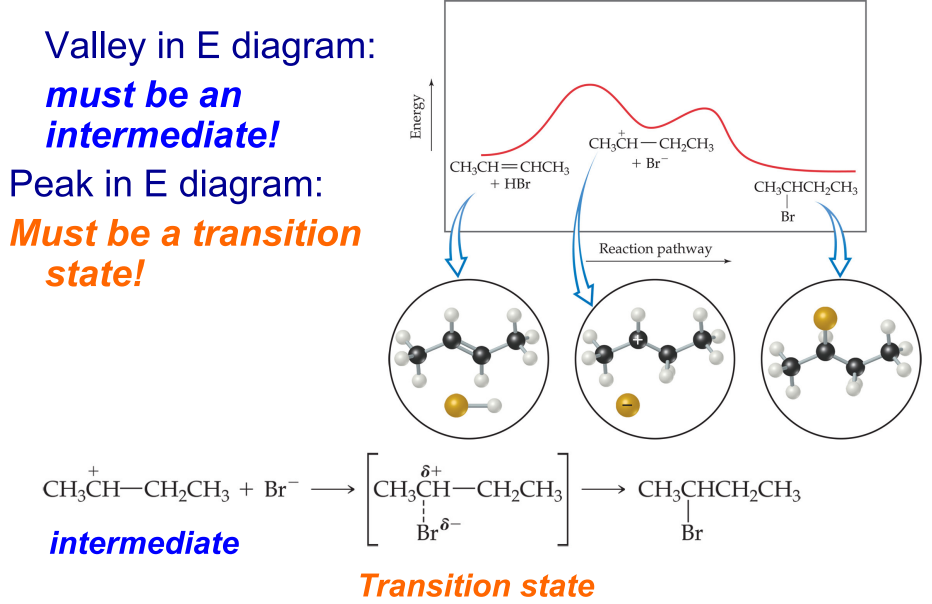
In first step, *π*-bond breaks and new C —H bond and cation form. An *intermediate*.

The top of the hill: a *transition state.*

 $CH_3CH = CHCH_3 + HBr \longrightarrow$



transition state.







- Contain at least one carbon–carbon triple bond.
- Carbons in triple bond *sp*-hybridized and have linear geometry.
- Also unsaturated.

Nomenclature of Alkynes

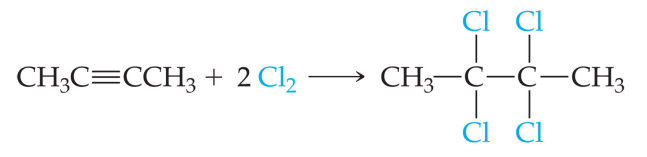
$\begin{array}{c} CH_3 - C \equiv C - CH - CH_3 \\ | \\ CH_3 \end{array}$

4-methyl-2-pentyne

- Analogous to naming of alkenes.
- Suffix is -yne rather than -ene.

Reactions of Alkynes

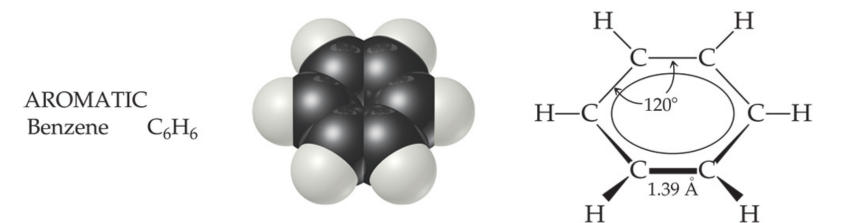
- Undergo many of the same reactions alkenes do.
- As with alkenes, impetus for reaction is replacement of π -bonds with σ -bonds.



2-Butyne

2,2,3,3-Tetrachlorobutane

Aromatic Hydrocarbons

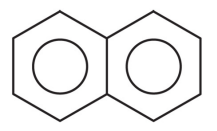


- Cyclic hydrocarbons.
- *p*-Orbital on each atom.
 Molecule is planar.
- Odd number of electron pairs in π -system.
- 4n+2 pi electrons in a cycle.

Aromatic Nomenclature

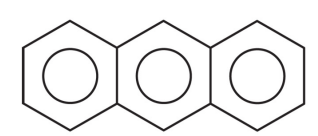
Many aromatic hydrocarbons are known by their common names.

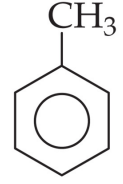




Benzene

Naphthalene



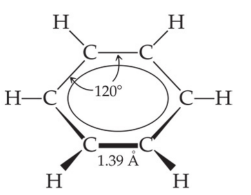


Anthracene

Toluene Methylbenzene

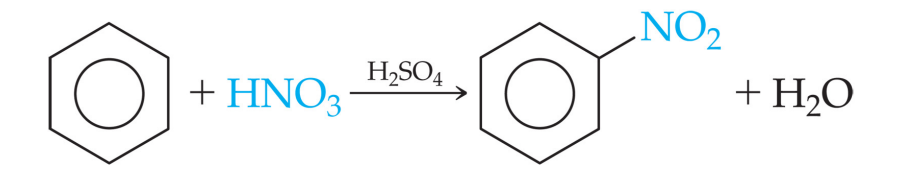
Reactions of Aromatic Compounds





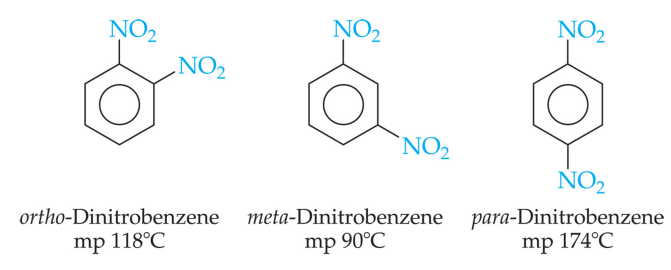
- Unlike in alkenes and alkynes, πelectrons do not sit between two atoms.
- Electrons are delocalized; this stabilizes aromatic compounds.

Reactions of Aromatic Compounds



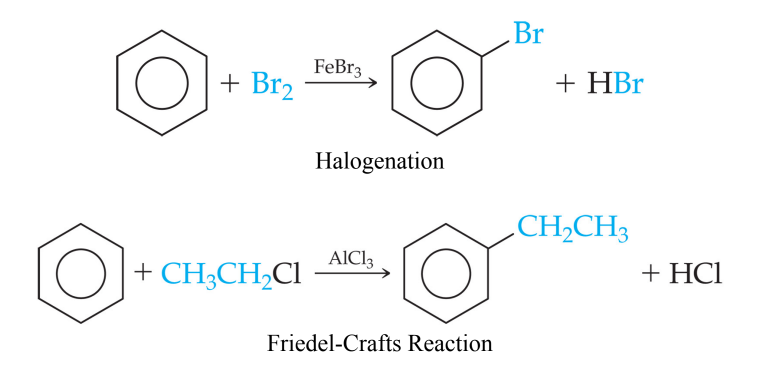
- Due to stabilization, aromatic compounds do not undergo addition reactions; they undergo substitution.
- Hydrogen is replaced by substituent.

Structure of Aromatic Compounds

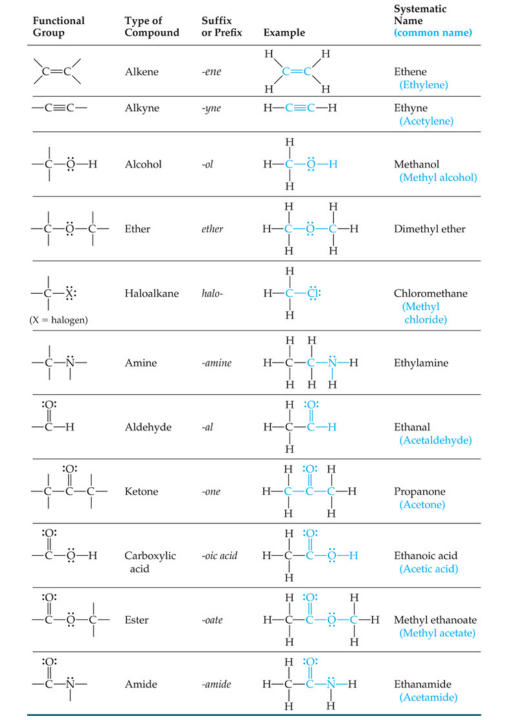


- Two substituents on a benzene ring could have three possible relationships
 - ➤ ortho-: On adjacent carbons.
 - ➤ meta-: One carbon between them.
 - ➢ para-: On opposite sides of ring.

Reactions of Aromatic Compounds



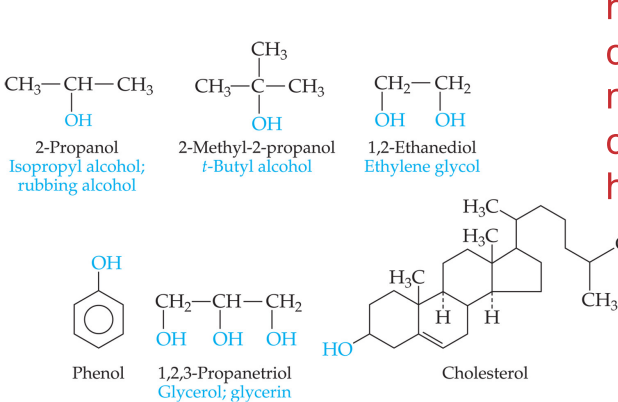
Reactions of aromatic compounds often require a catalyst.



Functional Groups Term used to refer to parts of organic molecules where reactions tend to occur.

Alcohols

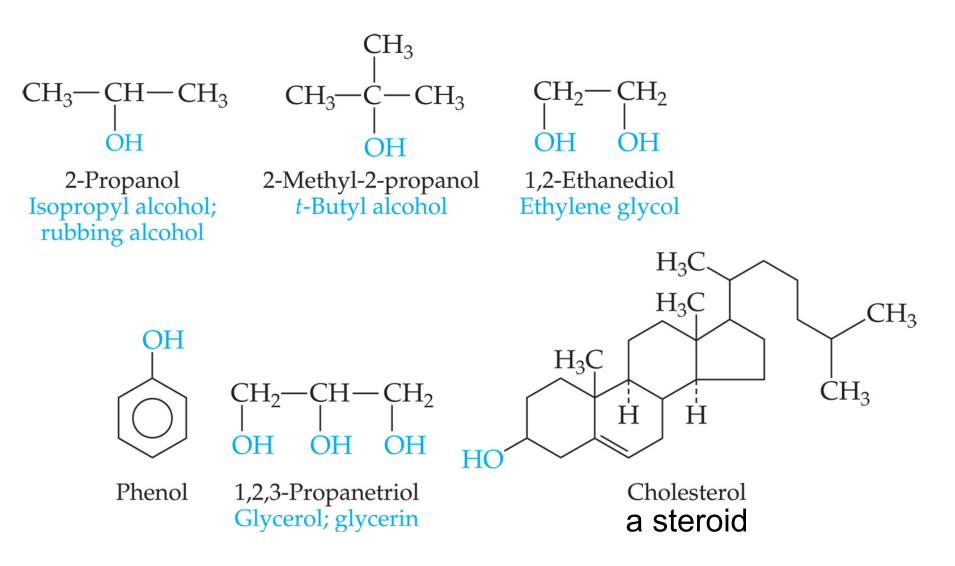
Contain one or more hydroxyl groups, —OH



 Named from parent hydrocarbon; suffix changed to -ol and number designates carbon to which hydroxyl is attached.

 CH_3

Alcohols

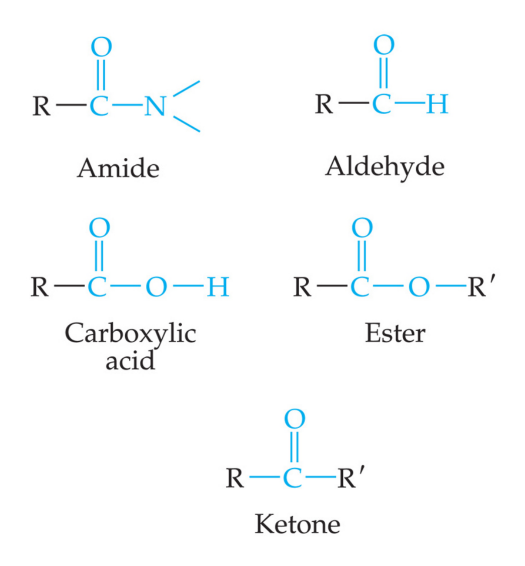


Ethers

$CH_2 = CH - O - CH = CH_2$ (an anesthetic)

- Tend to be quite unreactive.
- Therefore, they are good polar solvents.

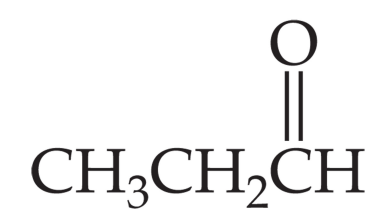
Carbonyl Compounds



- Contain C—O double bond.
- Include many classes of compounds.

Aldehydes

At least one hydrogen attached to carbonyl carbon.

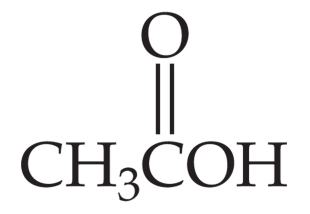


Ketones

 $\begin{array}{ccc} & \text{Two carbons} \\ & \text{bonded to} \\ & \text{carbonyl carbon.} \\ & \text{CH}_3\text{CH}_2\text{CCH}_2\text{CH}_3 \end{array}$

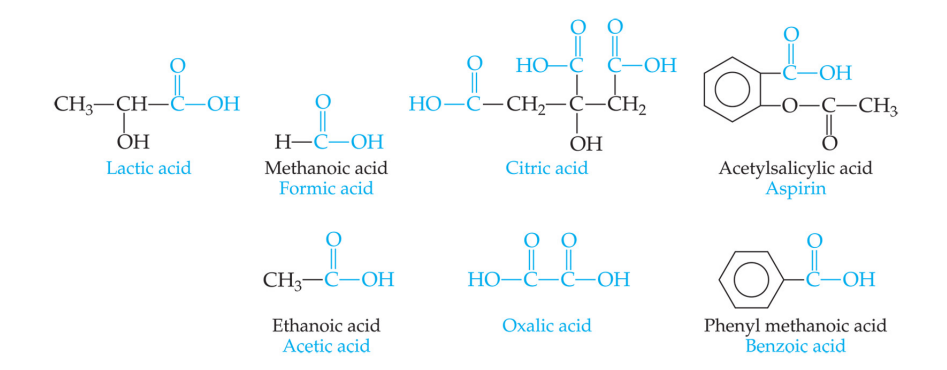
Carboxylic Acids

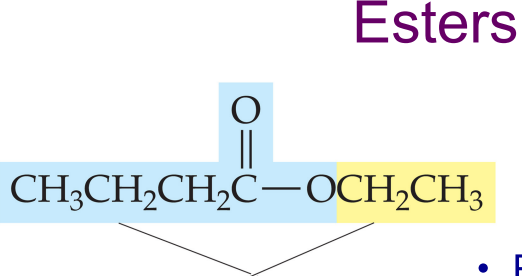
- Have hydroxyl group bonded to carbonyl group.
- Tart tasting.
- Carboxylic acids are weak acids .



CH₃COOH Acetic acid

Carboxylic Acids





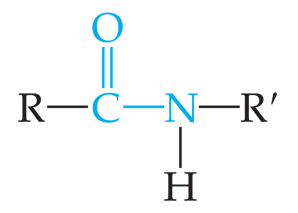
'butyrate

'Ethyl'

- Products of reaction between carboxylic acids and alcohols.
- Found in many fruits and perfumes.

Amides

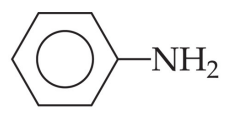
Formed by reaction of carboxylic acids with amines.



Amines

- Organic bases.
- Generally have strong, unpleasant odors.

 $(CH_3)_3N$



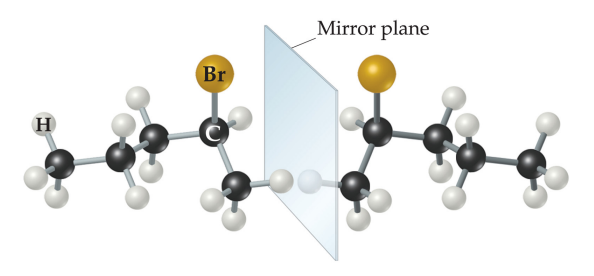
Ethylamine

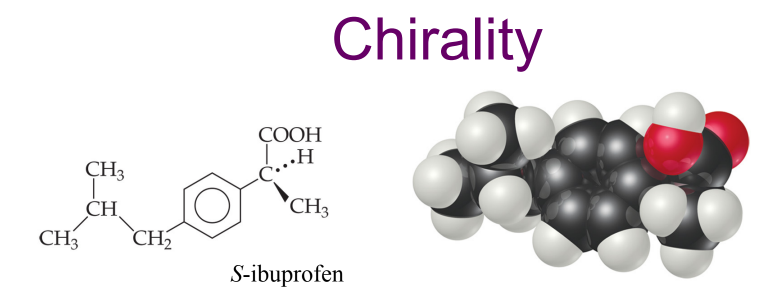
Trimethylamine

Phenylamine Aniline

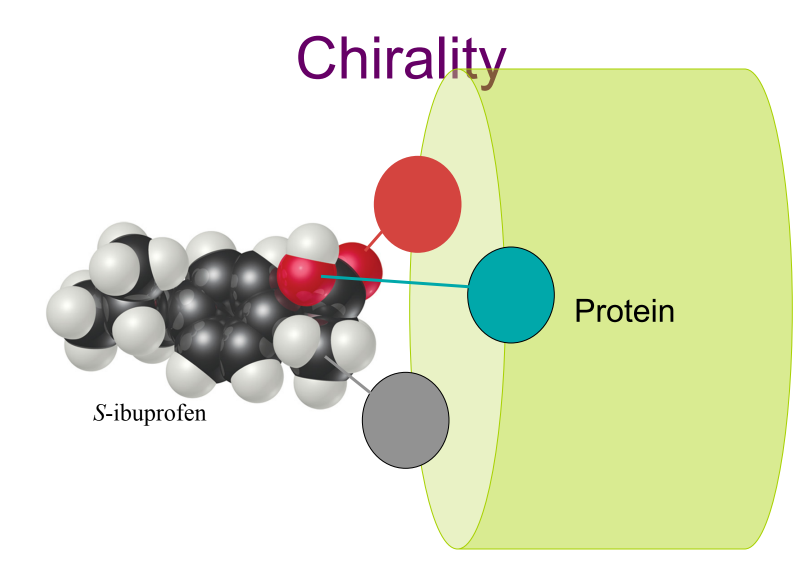
Chirality

- Carbons with four different groups attached to them are handed, or chiral.
- Optical isomers or stereoisomers
- If one stereoisomer is "right-handed," its enantiomer is "left-handed."





- Many pharmaceuticals are chiral.
- Often only one enantiomer is clinically active.
- Why?



Because they interact with a chiral protein binding site.

Exam 4 Topics

- 1. Valence bond theory
- 2. Molecular orbital theory
- 3. Chapter 24, coordination chemistry
- 4. Chapter 25, Organic (a little)

Valence bond theory:

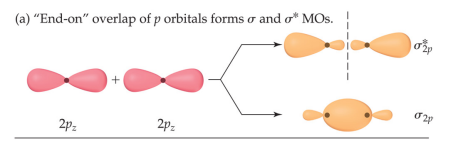
1.Hybridization (mostly covered in last exam)

2.Double bonds due to overlap of atomic p orbitals (pi bonds) 3.Concept of delocalization what orbitals are overlaping in a delocalized system? Exam 4, MO theory and coordination compounds Chapter 9, end and Chapter 24.

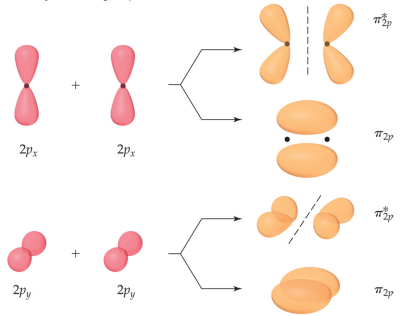
MO theory: Rules:

- 1. The number of MO's equals the # of Atomic orbitals
- 2. The overlap of two atomic orbitals gives two molecular orbitals, 1 bonding, one antibonding
- 3. Atomic orbitals combine with other atomic orbitals of *similar energy.*
- 4. Degree of overlap matters. More overlap means bonding orbital goes *lower* in E, antibonding orbital goes *higher* in E.
- 5. Each MO gets two electrons
- 6. Orbitals of the *same energy* get filled 1 electron at a time until they are filled.

Difference between pi and sigma orbitals



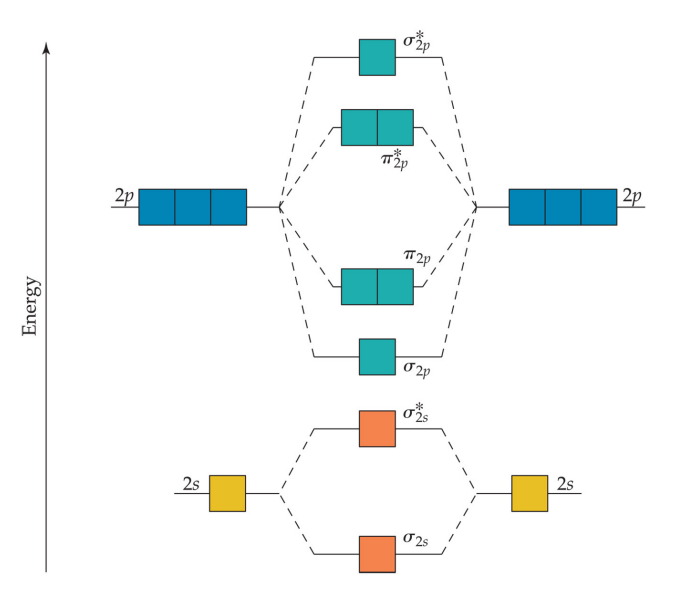
(b) "Sideways" overlap of *p* orbitals forms two sets of π and π^* MOs.



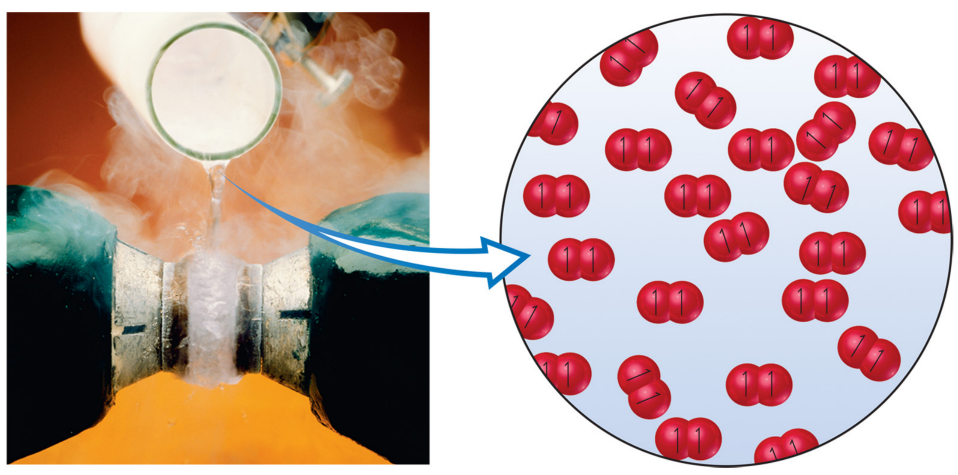
End on

Side to side.

A typical MO diagram, like the one below. For 2p and 2s atomic orbital mixing.



Oxygen O₂ is Paramagnetic, why?



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Show me why.

	Large 2 <i>s</i> -2 <i>p</i> interaction				Small 2 <i>s</i> -2 <i>p</i> interaction			
		B ₂	C ₂	N ₂		O ₂	F ₂	Ne ₂
	σ_{2p}^{*}				σ_{2p}^{*}			11
	π^*_{2p}				π_{2p}^{*}	1 1	11 11	11 11
	σ_{2p}			11	π_{2p}	11 11	11 11	11 11
	π_{2p}	1 1	11 11	11 11	σ_{2p}	11	11	11
	σ^*_{2s}	11	11	11	σ^*_{2s}	11	11	11
	σ_{2s}	11	11	11	σ_{2s}	11	11	11
Bond order Bond enthalpy (kJ/mol) Bond length (Å) Magnetic behavior		1 290 1.59 Paramagnetic	2 620 1.31 Diamagnetic	3 941 1.10 Diamagnetic		2 495 1.21 Paramagnetic	1 155 1.43 Diamagnetic	0

Exam 4 Chapter 24.

Concentrate on the homeworks and the quiz! Terms:

- 1. Coordination sphere
- 2. Ligand
- 3. Coordination compound
- 4. Metal complex
- 5. Complex ion
- 6. Coordination
- 7. Coordination number

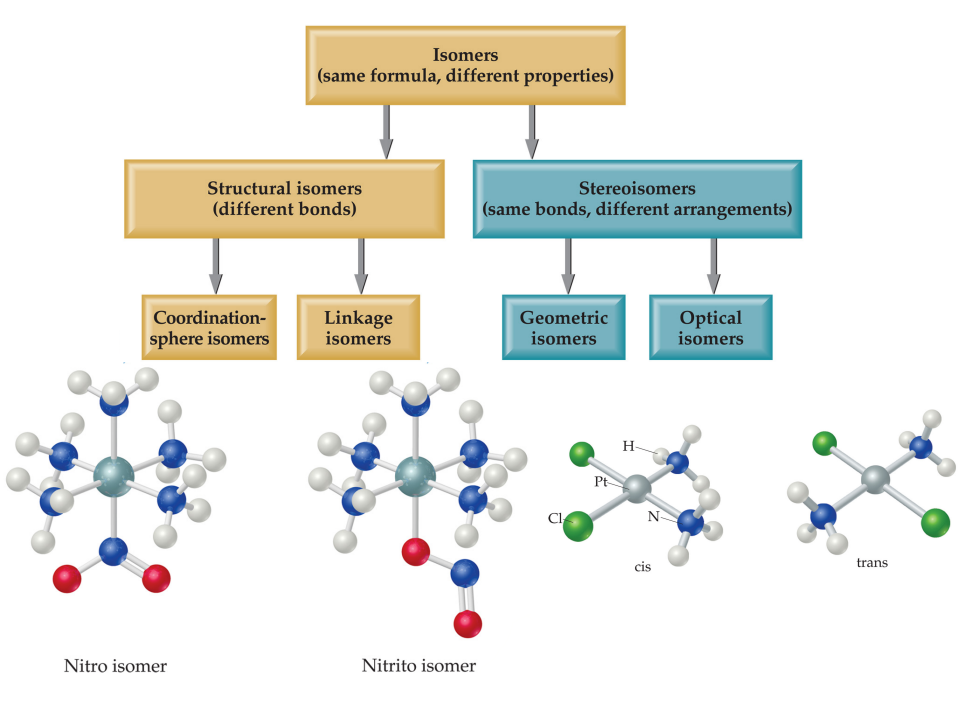
Same ligands different properties? Figuring oxidation number on metal Polydentate ligands (what are they)? Only ethylene diamine will be used (en) NH_2 - CH_2 - CH_2NH_2

Isomers.

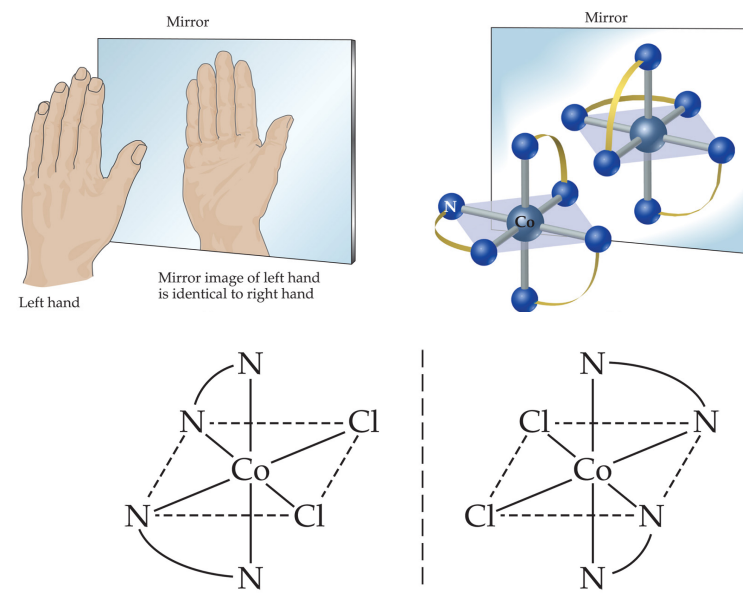
structural isomers (formula same, bonds differ) geometric isomers (formula AND bonds same, structure differs)

Stereoisomers:

Chirality, handedness,



Stereoisomers

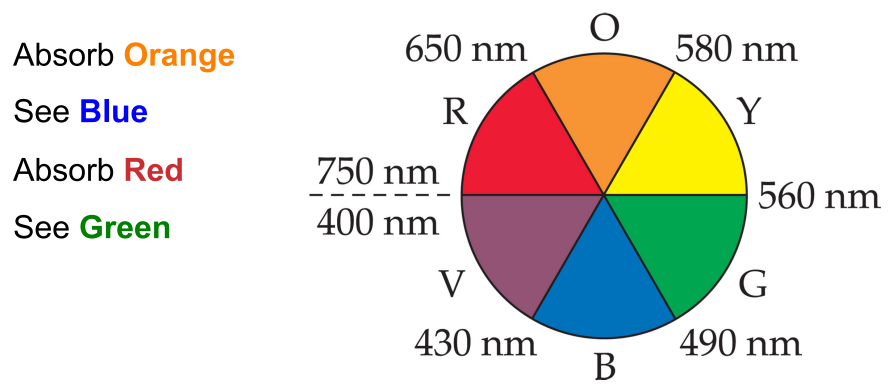


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Explaining the properties of metal complexes

Magnetism and color

How does seeing color work?



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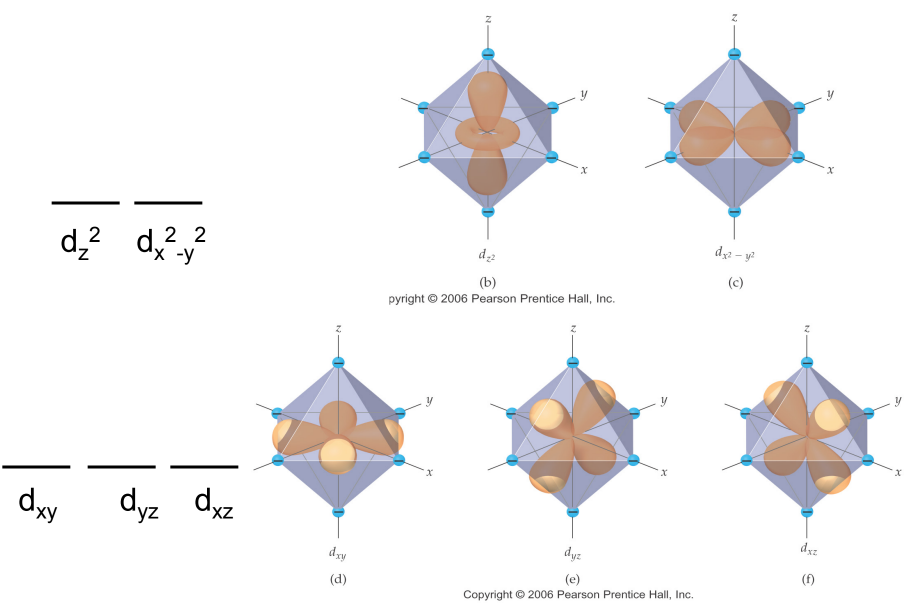
Different ligands on same metal give different colors



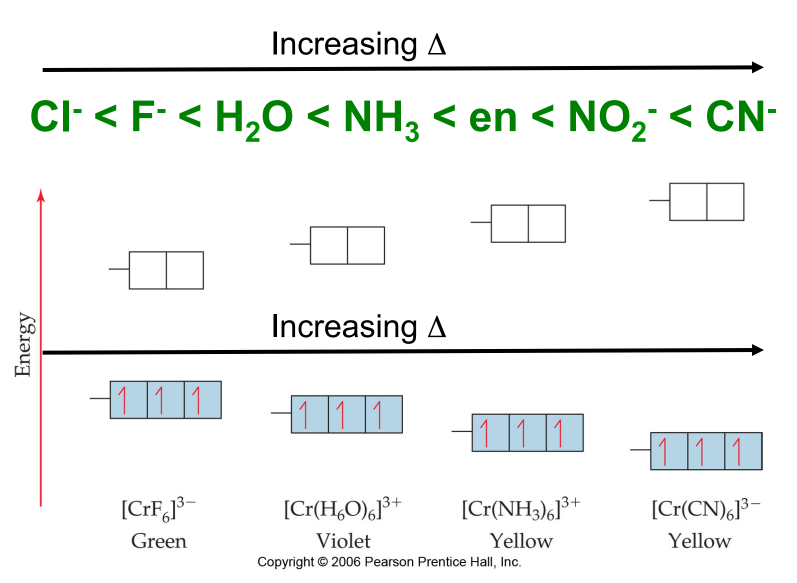
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Addition of NH₃ ligand to Cu(H₂O)₄ changes its color

Splitting of d orbitals in an octahedral ligand field

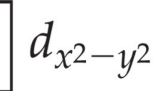




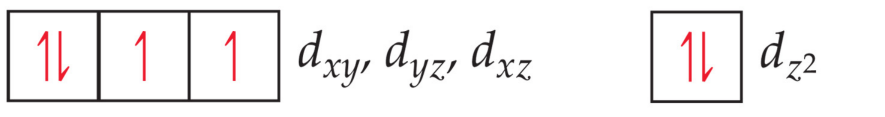


Know low spin versus high spin

There is also splitting from tetrahedral And square planar. Know they are different, don't remember exactly what square planer looks like.







$$1 d_{x^2-y^2}, d_{z^2}$$

Tetrahedral

Square planar

 $| 1| | 1| | d_{xz}, d_{yz}$

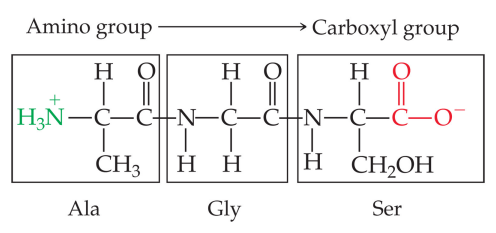
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Exam breakdown:

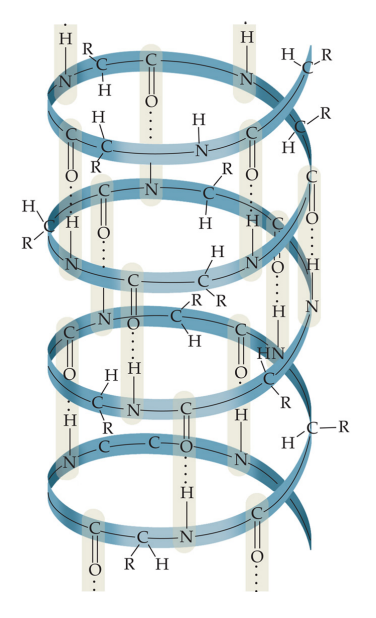
- Ch. 9: 3 Valence bond/hybridization M.O. diagrams Intermolecular forces 1 Ch 24 (23 in newer books): Color/absorption 3 Spec series 2 3 Isomers Magnetism (dia/para) 2 Ox#/coord# 3 High spin/low spin
- Ch. 25 (24 in newer books) Naming org. compounds Aromaticity Arrow pushing

Amino Acids and Proteins

- Proteins are polymers of α -amino acids.
- A condensation reaction between the amine end of one amino acid and the acid end of another produces a peptide bond.

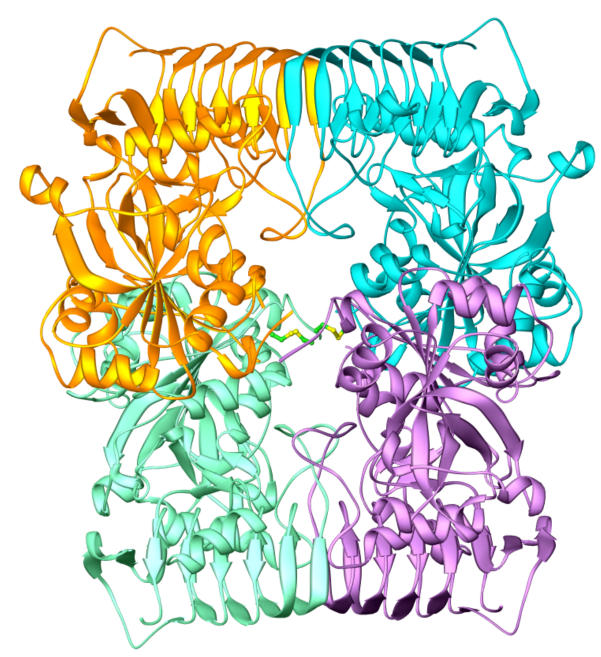


Amino Acids and Proteins



- Hydrogen bonding in peptide chains causes coils and helices in the chain.
- Kinking and folding of the coiled chain gives proteins a characteristic shape.

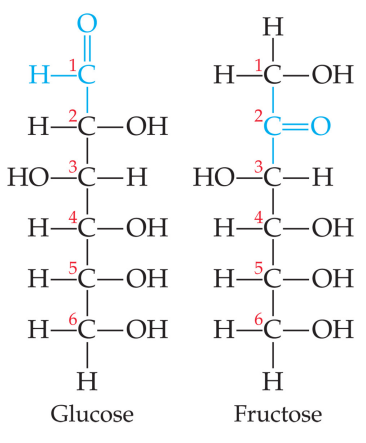
Amino Acids and Proteins



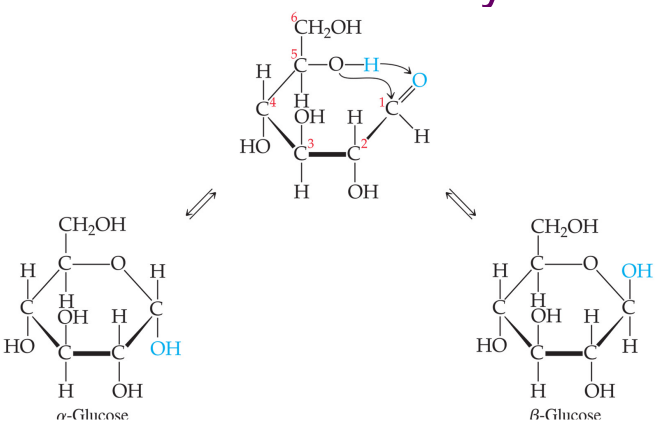
pyrophosphorylase makes starch in plants The complete molecule is a tetramter. It's mass is 240,000 amu.

Carbohydrates

Simple sugars are polyhydroxy aldehydes or ketones.



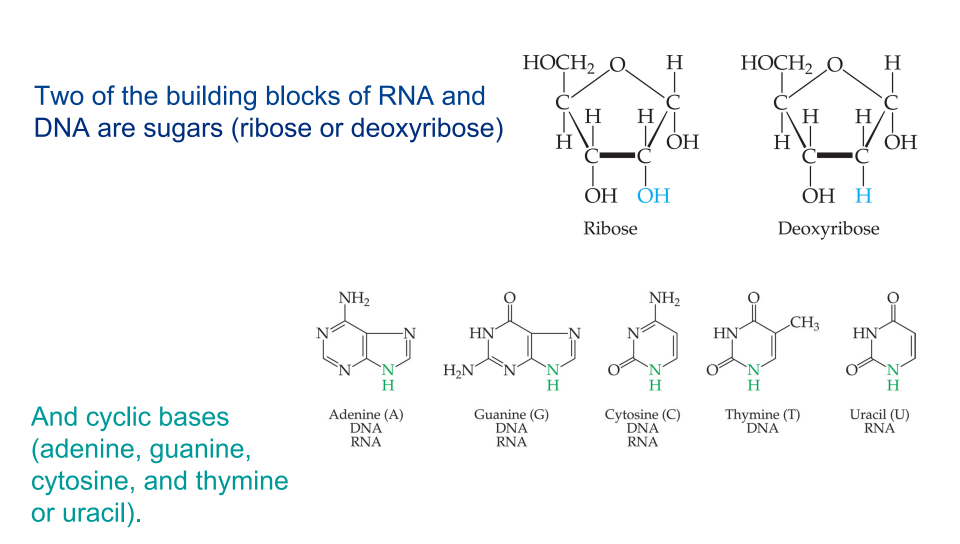
Carbohydrates



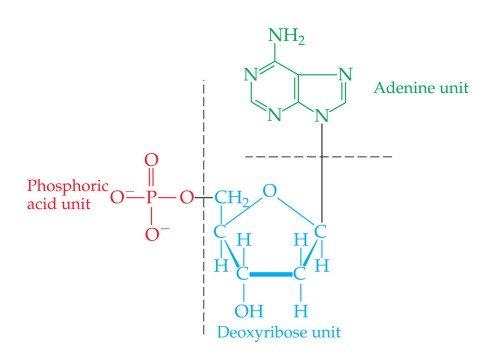
 In solution they form cyclic structures.

•These can form chains of sugars that form structural molecules such as starch and cellulose.

Nucleic Acids



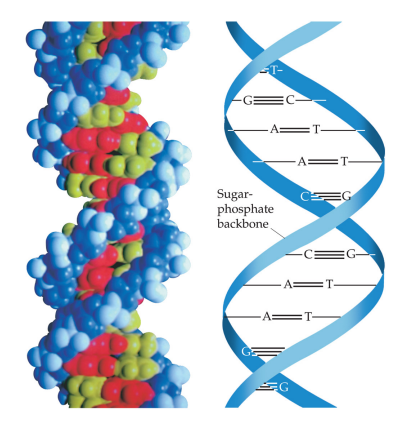
Nucleic Acids



These combine with a phosphate to form a nucleotide.

Nucleic Acids

Nucleotides combine to form the familiar double-helix form of the nucleic acids.



The best preparation:

- 1. The four exams and 6 quizzes.
 - "I' II bet, since I' m, after all, a little lazy, that I' II use some of these questions"

Topics:

Chapter 1. Dimensional analysis (at least 1 problem) significant figures

Chapter 2. History of atomic structure (2 problems) dalton cathode ray tubes rutherford gold foil experiment Miliken's oil drop experiment atomic numbers, mass, isotopes average at. weights The periodic table groups, periods, etc. molecular and empirical formulas ions, ionic compounds naming inorganic compounds naming binary molecular compounds (nitrogen triodide)

Chapter 2. compounds naming binary molecular compounds (nitrogen triodide) atomic structure, protons, neutrons, electrons

Chapter 3. stoichiometry calculations dealing with chemical reactions limiting reagent calculate empirical formula

Chapter 4, aqueous reactions, solution stoichiometry

strong and weak electrolytes the strong acids and the strong bases Know your anions and cations. precipitation reactions acid/base reactions Redox, oxidations numbers, redox reactions. the activity series solution stoichiometry titrations

Chapter 5, thermochemistry Kinetic and potential energy what is work versus heat? ΔH and ΔE what is a state function pV work calorimetry Hess' s law enthalpies of formation

Chapter 6, electronic structure of atoms wave and particle nature of light black body radiation, quantization of energy photons The photoelectric effect Line spectra & the bohr model Quantum mechanics atomic numbers atomic orbitals electron spin electron configuration The periodic table explained

Chapter 7, Periodic properties of the elements effective nuclear charge sizes of atoms sizes of ions ionization energies electron affinities group trends for: alkali metals alkine earch metals oxygen group, halogens, noble gases

Chapter 8, chemical bonding ionic bonding metal bonding covalent bonding bond polarity electronegativity lewis structures multiple bonds resonance structures octet rule and exceptions bond enthalpy and bond length

Chapter 9, VSEPR, valence bond and molecular orbital theory self explanatory.

Chapter 25, organic and biochemistry hydrocarbons functional groups naming organic compounds isomers what's an amino acid? What's a protein? What's a sugar? polysaccharide?

The exam, the lowdown:

36 questions from the previous 4 exams.
Exam 1: 9-10 questions
Exam 2: 8-9 questions
Exam 3: 9-10 questions
Exam 4: 8-9 questions