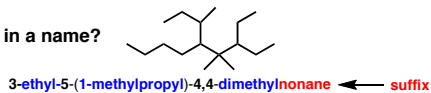


Introduction to Alkane Nomenclature

<http://masterorganicchemistry.com>

What's in a name?



the NUMBERS are called "locators"
items in BLUE are called "substituents".
the name in RED at the end is called the suffix.

The purpose of this sheet is to demonstrate the rules by which alkanes are named.

ORDER OF BUSINESS

A. Determine the priority of functional groups (not covered here since we're dealing with alkanes only)

B. Find the longest linear chain of your molecule, or the largest ring (whichever is greatest). This is the **Chain length rule** which defines both the "main chain" and also the suffix.

-**tiebreaker:** where more than one "path" along the molecule leads to the longest chain, the **main chain is the one that contains the most substituents.**

C. Identify the substituents along your main chain. Substituents are classified according to length of carbon chain and the suffix "yl" is attached.

D. Number your chain from one of the ends. The **LOWEST LOCATOR RULE** determines which end is chosen as carbon #1: "Number the chain such as to provide the lowest possible locators for the chain."

-**tiebreaker** for lowest-locator rule: **alphabetization**

E. Multiple instances of substituents are given the prefixes di, tri, tetra, etc.

-**note:** must have locator for all substituents. Example: 1,1-dimethyl is correct. 1-dimethyl is incorrect.

F. Branched substituents are numbered and named separately from the main chain, and put in brackets.

G. The FINAL name is assembled such as to arrange the substituents in alphabetical order.

- "di", "tri", "tetra" are ignored for alphabetization purposes.

- prefixes like "n", "tert", "i", and "sec" are ignored for alphabetization purposes.

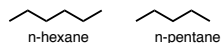
THE EXCEPTION is "isopropyl" and "isobutyl". For some reason these count as "i"

- not covered here, but this is also where one puts in descriptors like "cis", "trans", (R), (S) (E), (Z) and so on.

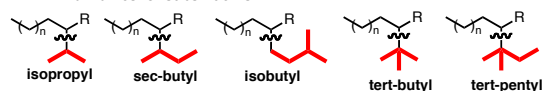
Names for Hydrocarbon Chains and Rings

1 methane		3 cyclopropane
2 ethane		4 cyclobutane
3 propane		5 cyclopentane
4 butane		6 cyclohexane
5 pentane		7 and higher follow the same pattern.
6 hexane		
7 heptane		
8 octane		
9 nonane		
10 decane		
11 undecane		
12 dodecane		
20 eicosane		

Sometimes you will see "n" in front to indicate that it is a straight-chain alkane



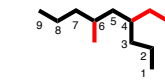
Trivial names for substituents



A. Determining the Priority of Functional Groups.

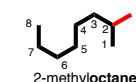
Too big a subject to cover on one sheet! This paper will focus on alkanes. Determining functional group priority will be the subject of a subsequent sheet.

B. Applying the Chain Length Rule



Longest chain is 9 carbons - suffix will be **nonane**

Watch out! Longest chain might not be drawn as a "straight chain"



Longest chain is 8 carbons - suffix will be **octane**

Chains vs. rings

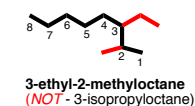
Rings take priority over chains, assuming there are only alkyl groups in the chain. ***see note below



propylcyclobutane butylcyclopropane

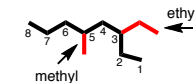
Tiebreaker: Alphabetization

Where more than one "longest chain" exists, the more substituted chain is chosen as the "longest chain"

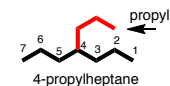


C. Identifying Substituents

Substituents are carbon fragments branching off the main chain. They are named according to the number of carbons like the main chain would be, except the "ane" is dropped and replaced with "yl"



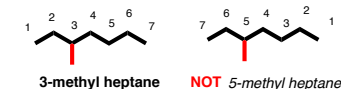
3-ethyl-5-methyloctane



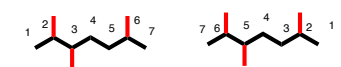
*** there seems to be some degree of confusion on this point. Both "1-butylcyclopropane" and "1-cyclopropylbutane" are commonly used. If someone can clarify, please write me.

D. Applying the Lowest Locator Rule

Number the chain from one end so as to provide the lowest locator possible for the first substituent.



This also applies for subsequent substituents, if either direction would give the same number.

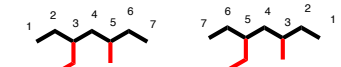


For rings with one substituent, the locator "1" can be dropped.

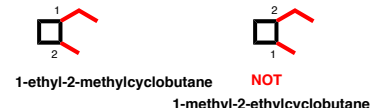


Tiebreaker: Alphabetization

If the same locators are obtained from either direction of the chain, the chain is numbered according to alphabetical order of the substituents.

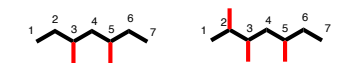


3-ethyl-5-methylheptane NOT 3-methyl-5-ethylheptane



E. Multiples of the Same Substituent

Multiples of the same substituent are given the prefixes "di", "tri", "tetra", etc. The lowest locator rule still applies.



3,5-dimethylheptane 2,3,5-trimethylheptane

Also applies to substituents on the same carbon:



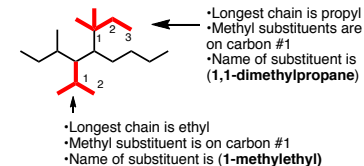
NOT 4-dimethylheptane
NOT 4,4-methylheptane

Extremely common mistakes!

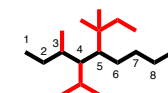
F. Dealing With Branched Substituents (the IUPAC Way)

Treat each branched substituent as its own naming problem.

Carbon #1 of the branched substituent will be where it meets the main chain.



We put the name in brackets to avoid confusing the numbers of the branched substituent with the numbers of the main chain.



In certain instances, you may see the trivial names isopropyl, isobutyl, tert-butyl, tert-pentyl used.

e.g. 4-isopropyl-3-methyl-5-tert-pentylnonane.

The official IUPAC nomenclature system will never let you down. It would probably be best to just go with that.

G. Putting the Name Together.

1. Put your substituents together in **alphabetical order**.

For alphabetization purposes:

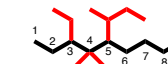
IGNORE "di", "tri", "tet", etc.
"sec-", "tert-", "n-", "cyclo-

DO NOT IGNORE "iso". For some strange reason "isopropyl" is alphabetized under "i" and not "p". Don't ask me why.

2. Affix the **locators**.

3. Make sure any **branched substituents** are in brackets.

4. Attach the **suffix** at the end.



3-ethyl-5-(1-methylpropyl)-4,4-dimethylnonane

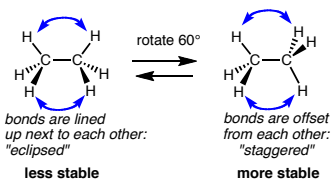
This sheet © 2010 James A. Ashenhurst
<http://masterorganicchemistry.com>

Errors/omissions/suggestions?
james@writechem.com
Version 1.1, Jan 2011

For more complete resources on nomenclature consult:
1) "Organic Chemistry Online" by William Reusch:
<http://www2.chemistry.msu.edu/80/faculty/reusch/VirtTxJml/intro1.htm>
2) IUPAC "Blue Book"
<http://www.acdlabs.com/iupac/nomenclature/>

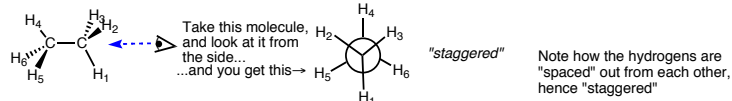
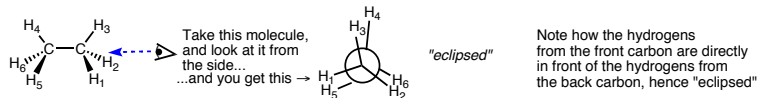
What are conformations?

•Sigma bonds (single bonds) are free to rotate.
 •Through the rotation of bonds, molecules can adopt different 3-dimensional shapes. These are called **conformations**. Two identical molecules with different 3-D shapes are called **conformational isomers**.
 •Not all conformations are equal in energy. Because electron clouds repel, conformations which maximize the distance between atoms are generally favored (i.e. lower in energy).

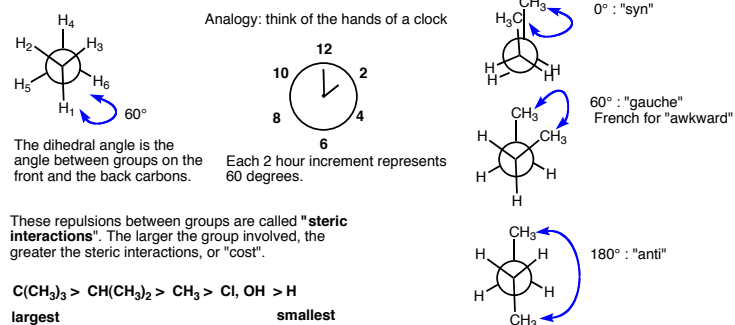


difference: 2.4 kcal/mol (10 kJ/mol)

Key Tool: The Newman projection



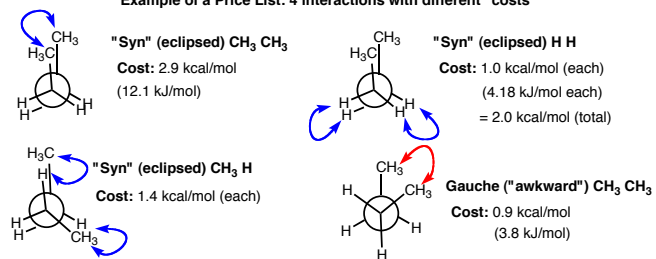
Key Tool: The "Dihedral Angle"



Interestingly, CH_3 is larger than Cl or OH , since it "sweeps out" a larger area

Repulsions between groups cause **strain**, which can be relieved through bond rotation; this is often called "**torsional strain**".

Example of a Price List: 4 interactions with different "costs"



Example: how to figure out the rotational energies for butane ($CH_3CH_2CH_2CH_3$) ?

Step 1: like a clock, pick one group as the "hour hand" and one as the "minute hand". Then rotate it through 360°, in 60° (2h) increments.

Pick the two methyl groups as your "hands".

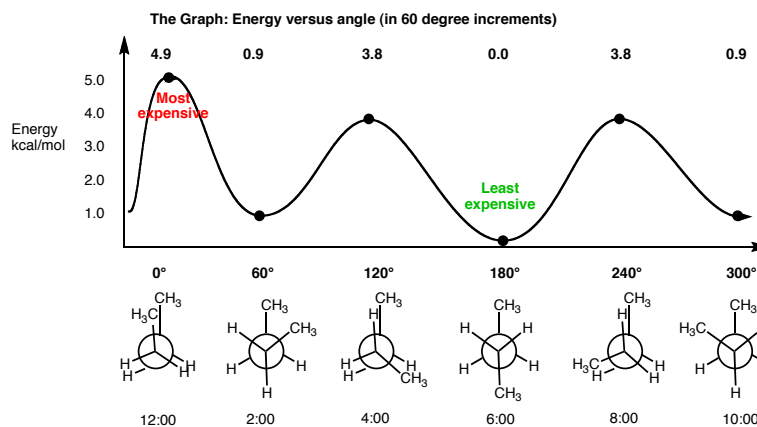
Step 2: Calculate the "costs" for each of the six conformations

Example: Adding up the "costs" of each conformation for butane

Dihedral angle	0°	60°	120°	180°	240°	300°
Eclipsed CH_3-CH_3	2.9	-	-	-	-	-
Eclipsed CH_3-H	-	-	2 x 1.4	-	2 x 1.4	-
Eclipsed $H-H$	2 x 1.0	-	1.0	-	1.0	-
Gauche CH_3-CH_3	-	0.9	-	-	-	0.9
Total Cost (kcal/mol)	4.9	0.9	3.8	0.0	3.8	0.9

Most "expensive" (at 0° and 300°)
 Least "expensive" (at 180°)

Step 3: Make a graph



Omissions, Mistakes, Suggestions?

james@masterorganicchemistry.com
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Introduction to Cycloalkanes

"Master Organic Chemistry"
 masterorganicchemistry.com
 Nov 2012. Version 1.0

Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

Cyclic molecules

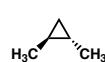
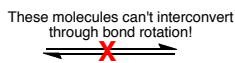
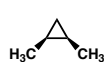
Molecules in organic chemistry don't just exist as chains - they can also exist as rings. Compare propane and cyclopropane:



note how there are two fewer hydrogens in the molecular formula of cyclopropane than for propane

Stereochemistry

When groups are on a ring, they are constrained - they can't rotate in the same way that linear molecules can. This can give rise to "cis" and "trans" isomers - stereoisomers.

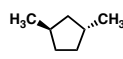
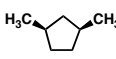


cis-1,2-dimethylcyclopropane

trans-1,2-dimethylcyclopropane

stereoisomers: same molecular formula, different boiling points/melting points

Other examples:



cis-1,3-dimethylcyclopentane

trans-1,3-dimethylcyclopentane

cis-1,2-dimethylcyclohexane

trans-1,2-dimethylcyclohexane

Ring Strain

In contrast to chains, where sp^3 hybridized carbons can adopt their ideal geometry of 109.5° , the geometry of carbons in some cycloalkanes can be far from ideal, leading to **ring strain**



Name: Cyclopropane

Cyclobutane

Cyclopentane

Cyclohexane

Interior angle: 60°

90°

108°

120°

Ring strain (kcal/mol)

27.6 kcal/mol

26.4 kcal/mol

6.5 kcal/mol

0

(kJ/mol)

115.5 kJ/mol

110.4 kJ/mol

27.2 kJ/mol

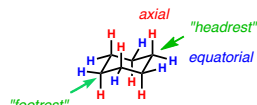
0

Ring strain is a combination of **angle strain** (where geometry of the carbon deviates from the ideal) and **torsional strain** (arising from eclipsing interactions)

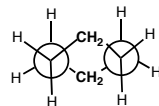
The Chair Conformation of Cyclohexane

The cyclohexane ring is not a flat hexagon. Why not? Two reasons: 1) the bond angles would be 120° (creating angle strain) and 2) the neighboring hydrogen atoms would be eclipsed with each other. Instead cyclohexane adopts a *chair* conformation where all bond angles are close to 109 degrees and all neighboring groups are staggered.

Groups that point "straight" up and down are said to be "axial" - in red



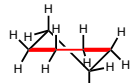
Newman projection of chair



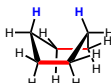
In the chair form, all substituents are staggered along every C-C bond



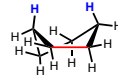
Planar
 angle strain (120°)
 eclipsing strain
 (two hydrogens omitted for clarity)



"Half-Chair"
 eclipsing strain
 steric crowding
 10.7 kcal/mol less stable than chair



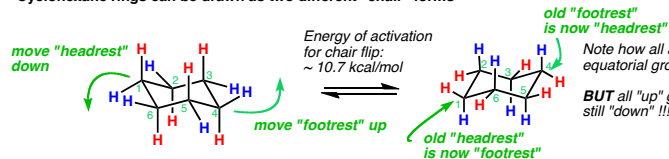
"Boat"
 eclipsing strain
 steric crowding
 6.9 kcal/mol less stable than chair



"Twist boat"
 eclipsing strain
 steric crowding
 5.3 kcal/mol less stable than chair

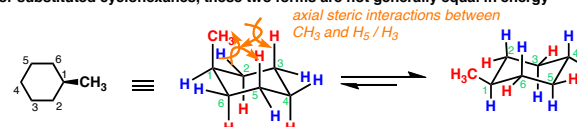
Other conformations of cyclohexane (higher in energy than the chair)

Cyclohexane rings can be drawn as two different "chair" forms



Note how all axial groups became equatorial and all equatorial groups became axial...
BUT all "up" groups are still "up", and all "down" groups are still "down" !!!

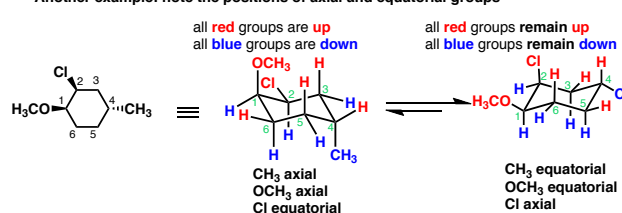
For substituted cyclohexanes, these two forms are not generally equal in energy



CH₃ axial (less stable by 1.8 kcal/mol)

CH₃ equatorial

Another example: note the positions of axial and equatorial groups



all red groups are up
 all blue groups are down

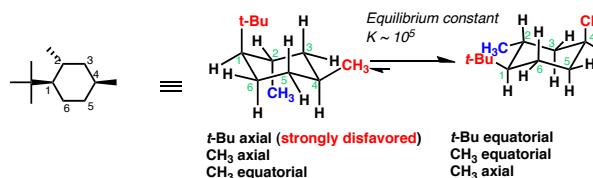
all red groups remain up
 all blue groups remain down

CH₃ axial
 OCH₃ axial
 Cl equatorial

CH₃ equatorial
 OCH₃ equatorial
 Cl axial

Important: the *t*-butyl group is conformationally "locked" in the equatorial position

The *t*-butyl group is so bulky that the axial position is extremely disfavored. Equilibrium favors the equatorial position to such an extent that one can consider the chair to be "locked" in the conformation with *t*-Bu equatorial

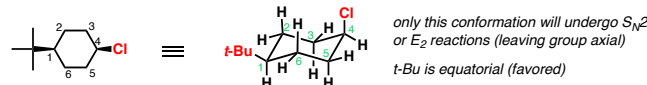


t-Bu axial (strongly disfavored)
 CH₃ axial
 CH₃ equatorial

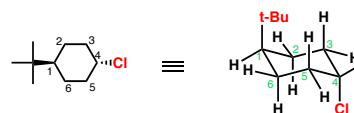
t-Bu equatorial
 CH₃ equatorial
 CH₃ axial

Application in substitution and elimination reactions (S_N2 and $E2$)

Understanding the proportions of different chair forms is important because S_N2 and $E2$ reactions will only occur when the leaving group is in the axial position.



only this conformation will undergo S_N2 or $E2$ reactions (leaving group axial)
t-Bu is equatorial (favored)



only this conformation will undergo S_N2 or $E2$ reactions (leaving group axial)
t-Bu is axial (disfavored)
Substitution and elimination reactions will be extremely slow for this molecule

The axial position is less favored for substituents since it can undergo unfavorable steric interactions with other axial substituents. These are avoided in the equatorial position. (Technically these are "gauche" interactions)

Each interaction between axial CH₃ and axial H "costs" ~ 0.9 kcal/mol, for a total of about 1.75 kcal/mol.

Using this value we can calculate the % of 1-methylcyclohexane that is present in the axial form at equilibrium:

$$\Delta G = -RT \ln K \quad K = e^{-\Delta G/RT}$$

At 1.75 kcal/mol this gives us $K = 0.052$, or approximately 95:5 ratio in favor of the equatorial CH₃

This value: 1.75 kcal/mol is sometimes referred to as the "A" value of CH₃

The size of the A value is related to the magnitude of the steric interactions between axial substituents

Group	A Value	Group	A Value
CH ₃	1.74	Cl	0.43
CH ₂ CH ₃	1.75	OH	0.87
<i>i</i> -Pr	2.15	OCH ₃	0.6
<i>t</i> -Bu	4.7	NH ₂	1.6

This table gives an idea of the preference of the substituent for the equatorial position. For example *t*-Bu is the bulkiest of these groups, whereas Cl is the least bulky. Using the Gibbs equation it's possible to calculate the position of the equilibrium for each substituent.

For example

A value	% equatorial (at 25°C)
0	50%
0.5	70%
1	84%
1.5	93%
2	97%
2.5	98.5%
3	99.4%
3.5	99.7%
4.0	99.9%
4.5	99.95%
5	99.98%

Omissions, Mistakes, Suggestions?

jes@masterorganicchemistry.com

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Summary of Functional Groups (1)

"Master Organic Chemistry"

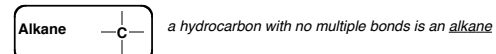
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August 2012 Version 1.2

Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

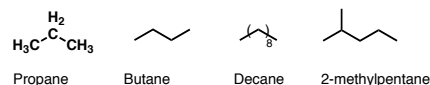
What Are Functional Groups?

Functional groups are collections of atoms that have a common pattern of chemical reactivity



Suffix: "-ane". As a substituent: "alkyl"

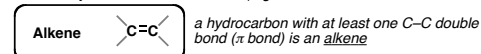
Examples:



Characteristics: nonpolar

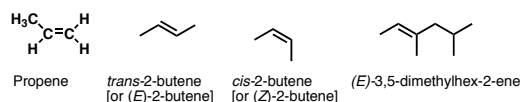
Geometry: tetrahedral (sp^3 hybridized)

Reactivity: **free radical reactions** (e.g. free radical chlorination or bromination)



Suffix: "-ene". As a substituent: "alkenyl"

Examples:

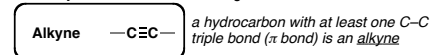


Characteristics: nonpolar. Molecule cannot rotate along double bond.

Geometry: trigonal planar (sp^2 hybridized)

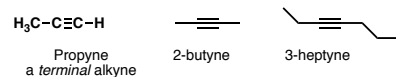
Reactivity: undergo **addition reactions**, as well as **oxidative cleavage**

Stability increases with increasing # of carbons attached



Suffix: "-yne". As a substituent: "alkynyl"

Examples:



Alkynes with a C-H bond are called "terminal" alkynes

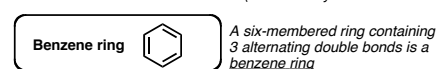
Geometry: linear (sp hybridized)

Characteristics: non polar

Reactivity: **addition reactions**

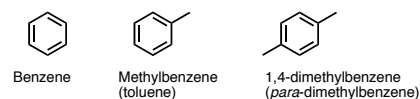
oxidative cleavage reactions

acid-base reactions (terminal alkynes are unusually acidic)



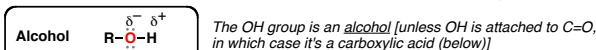
Suffix: "benzene". As a substituent: "phenyl"

Examples:



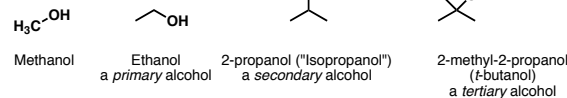
Reactivity: **substitution reactions** (e.g. electrophilic aromatic substitution or nucleophilic aromatic substitution)

Less reactive than normal alkenes due to aromatic stability



Suffix: "-ol". As a substituent: "hydroxy"

Examples:

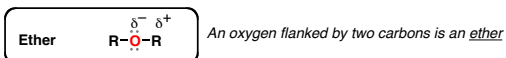


Characteristics: polar (O-H group participates in hydrogen bonding)

Reactivity: **acid-base reactions** (can act as acids or bases)

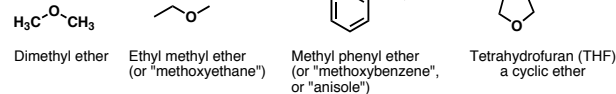
substitution reactions (can act as nucleophiles)

oxidation reactions (*primary and secondary alcohols (and methanol) can be oxidized to aldehydes, ketones, or carboxylic acids, depending on structure and reagent used*)



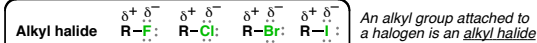
As a substituent: "alkoxy"

Examples:



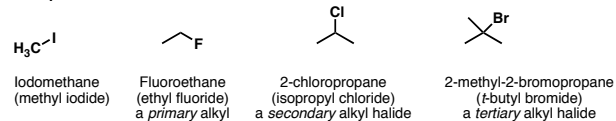
Characteristics: borderline between nonpolar and polar (due to dipole-dipole)

Reactivity: **acid-base reactions** (oxygen can act as a very weak base)



Suffix: "-ane". As a substituent: "haloalkyl"

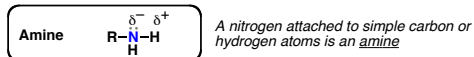
Examples:



Characteristics: generally considered non polar (but more polar than alkanes)

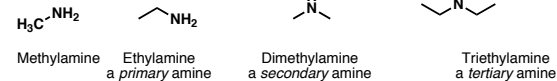
Reactivity: **substitution reactions** (Cl, Br, I can be good leaving groups)

elimination reactions (Cl, Br, I can be good leaving groups)



Suffix: "-ine". As a substituent: "amino"

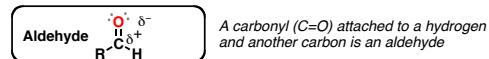
Examples:



Characteristics: polar (N-H group participates in hydrogen bonding, although not as much as a hydroxy group)

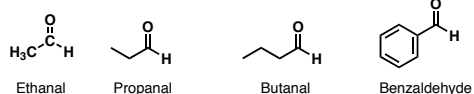
Reactivity: **acid-base reactions** (tend to act as bases)

substitution reactions (can act as nucleophiles)



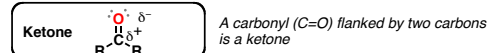
Suffix: "-al" (if attached to ring: carbaldehyde) As a substituent: "oxo"

Examples:



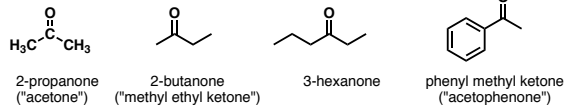
Characteristics: the C=O bond is somewhat polar

Reactivity: **addition reactions** (the carbonyl carbon reacts easily with nucleophiles)



Suffix: "-one". As a substituent: "oxo"

Examples:



Characteristics: the C=O bond is somewhat polar (less so than O-H however)

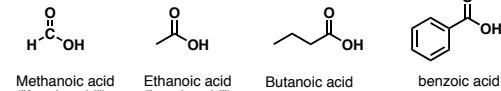
Reactivity: **addition reactions** (the carbonyl carbon reacts easily with nucleophiles)

acid-base reactions (carbons adjacent to the ketone can be deprotonated to give enolates)



Suffix: "-oic acid"

Examples:



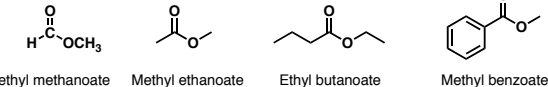
Reactivity: **acid-base reactions** (the O-H is acidic)

acyl substitution reactions (can replace OH with other groups under acidic conditions)



Suffix: "-oate"

Examples:



Reactivity: **acyl substitution reactions** (can replace OR with other functional groups under acidic conditions)

addition reactions (the carbonyl carbon reacts easily with nucleophiles)

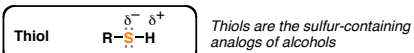
Omissions, Mistakes, Suggestions?

james@masterorganicchemistry.com

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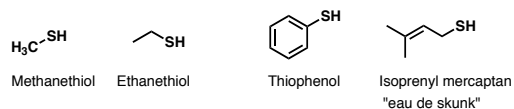
What Are Functional Groups?

Functional groups are collections of atoms that have a common pattern of chemical reactivity

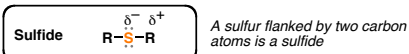


Suffix: "-thiol". As a substituent: "mercapto"

Examples:

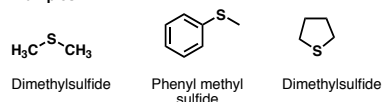


Reactivity: **acid-base reactions** (thiols can act as weak acids)
substitution reactions (thiols are good nucleophiles)

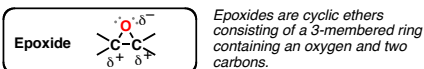


Suffix: "-sulfide".

Examples:



Reactivity: **reduction reactions** (will reduce the intermediate of ozonolysis reactions)

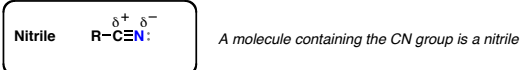


Suffix: "-oxide"

Examples:

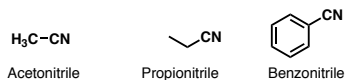


Reactivity: epoxides have considerable ring strain, and will undergo **ring opening** when treated with various types of nucleophiles (e.g. Grignard reagents, organolithium reagents)



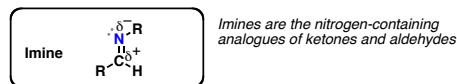
Suffix: "-nitrile"

Examples:



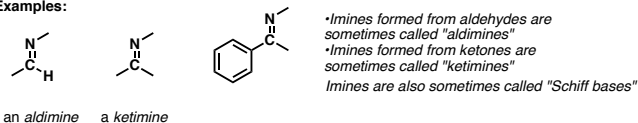
Reactivity: **reduction reactions** (nitriles can be reduced to amines with strong reductants such as LiAlH_4)
hydrolysis reactions (nitriles can be converted to carboxylic acids with aqueous acid)

Summary of Functional Groups (2)

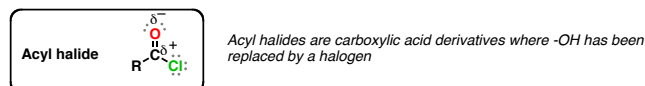


Suffix: "-imine".

Examples:

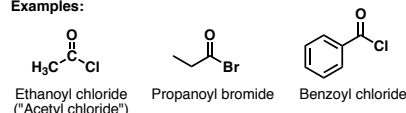


Reactivity: **reduction reactions** (imines can be reduced to amines)
hydrolysis reactions (imines can be hydrolyzed to give back aldehydes or ketones)

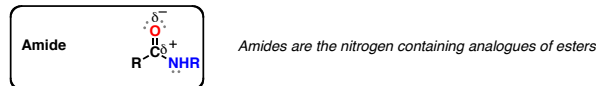


Suffix: "-oyl halide"

Examples:

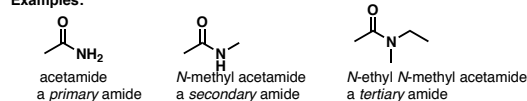


Reactivity: **acyl substitution reactions** (halides are good leaving groups and can be replaced by good nucleophiles)

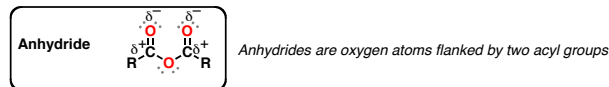


Suffix: "-amide"

Examples:

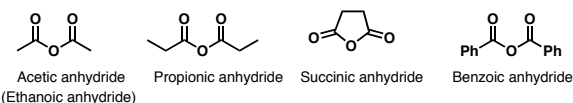


Reactivity: **reduction reactions** (amides can be reduced to amines with strong reductants such as LiAlH_4)
hydrolysis reactions (amides can be hydrolyzed to carboxylic acids with aqueous acid)

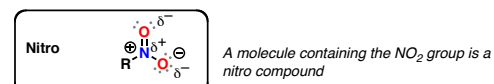


Suffix: "-anhydride"

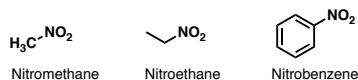
Examples:



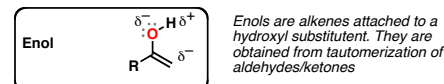
Reactivity: **acyl substitution reactions** (anhydrides can be cleaved with good nucleophiles)
reduction reactions (anhydrides can be reduced to aldehydes with various reducing agents)



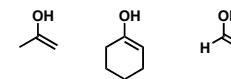
Examples:



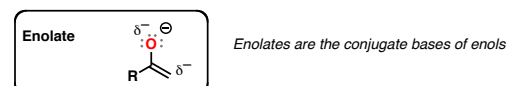
Reactivity: **reduction reactions** (nitro groups can be reduced to amines)
acid-base reactions (C-H bonds adjacent to the NO₂ group are acidic)



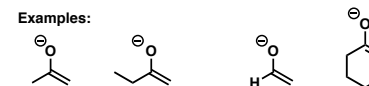
Examples:



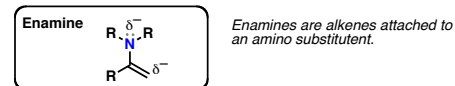
Reactivity: enols are good **nucleophiles**. They will perform **addition** reactions to aldehydes and ketones, among other reactions.
 Enols will **tautomerize** to aldehydes/ketones



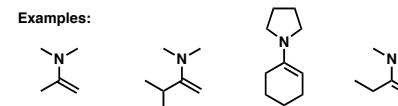
Examples:



Reactivity: enolates are excellent **nucleophiles** (better than enols) They will perform **addition** reactions to aldehydes and ketones, as well as **substitution** reactions with alkyl halides (among others)



Examples:



Reactivity: enamines are excellent **nucleophiles**. They will perform **addition** reactions to aldehydes and ketones, as well as **substitution** reactions with alkyl halides (among others)

Omissions, Mistakes, Suggestions?

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Introduction to Acids and Bases

"Master Organic Chemistry"
 masterorganicchemistry.com
 Nov 2012. Version 1.0

Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

1. Introduction

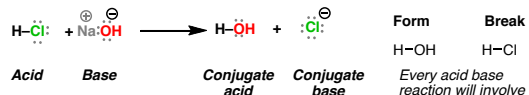
Two definitions of acids:

- Bronsted definition:** An acid donates a proton (H⁺)
- Lewis definition:** An acid accepts a lone pair of electrons

Two definitions of bases:

- Bronsted definition:** A base accepts a proton (H⁺)
- Lewis definition:** A base donates a lone pair of electrons

Simple example of an acid-base reaction



The four actors:

- Acid:** donates a proton (or - accepts a lone pair)
- Base:** accepts a proton (or - donates a lone pair)
- "Conjugate base":** is what remains of the acid after it donates H⁺
- "Conjugate acid"** is the species formed when the base accepts H⁺

2. How Do We Define and Measure Acidity?

Acidity is defined as the position of the equilibrium between an acid and its conjugate base. We obtain this number through **experimental measurement**.

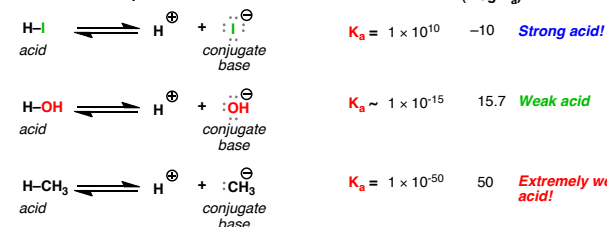
For a generic acid-base reaction:



These values can differ by an extremely large amount (10⁶⁰) so it is convenient to work with the logarithm of this number, defined as the pKa:

$$\text{p}K_a = -\log K_a$$

3 concrete examples:



3. What factors make an acid strong or weak?

Generally, the strength of an acid is related to the **stability** of the conjugate base.

The greater the stability of the conjugate base, the stronger the acid. The lower the stability of the conjugate base, the weaker the acid.

A different way to put it: **stability is inversely related to basicity**.

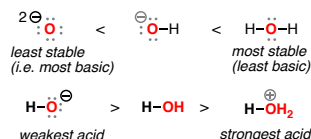
Stable conjugate base = a "weak" base
 Unstable conjugate base = a "strong" base

The stronger the acid, the weaker the conjugate base
The weaker the acid, the stronger the conjugate base.

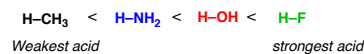
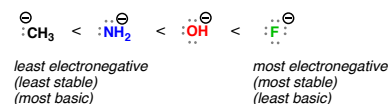
The best way to understand acidity/basicity is to understand the factors that stabilize negative charge.

There are seven important factors (placed here in roughly decreasing order of importance).

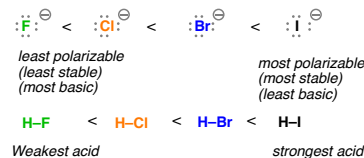
Factor #1: The less charge, the better



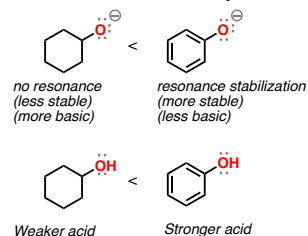
Factor #2: Going across the periodic table, greater electronegativity stabilizes negative charge.



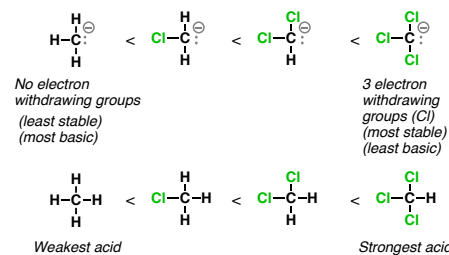
Factor #3: Going down the periodic table, greater polarizability stabilizes negative charge.



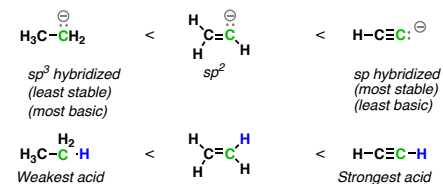
Factor #4: Negative charge is stabilized by resonance, which delocalizes electron density



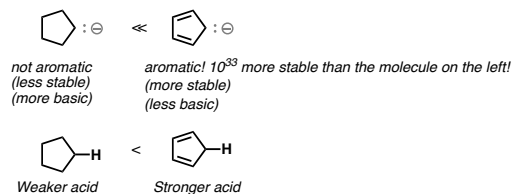
Factor #5: Negative charge is stabilized by adjacent electron withdrawing groups (inductive effects)



Factor #6: Negative charge is stabilized as the hybridization includes more s orbitals



Factor #7: A Special case - negative charge is especially stable if the electrons are part of an aromatic π-system



4. What makes an acid-base reaction favorable (or not)?

Key principle: In **favorable** acid-base reactions, a stronger acid adds to a stronger base to give a weaker acid and a weaker base

Key questions to ask: Is the "acid" stronger than the "conjugate acid"?
 Is the "base" stronger than the "conjugate base"?
If so, the acid-base reaction will be favorable.

How do you tell which acid is stronger or weaker?

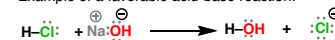
A **pKa table** collates **experimental measurements** of acidity, and thus incorporates all the factors mentioned above.

Strong acids have **low** pKa values. Weak acids have **high** pKa values.

A very abridged pKa table:

	Acid	pKa	Conjugate Base
↑ strongest acid weakest acid	H-Cl	-8	Cl ⁻
	H-OH ₂ ⁺	-1.7	H ₂ O
	H-O-C(=O)CH ₃	4.5	O=C(=O)CH ₃
	H-NH ₃ ⁺	10	:NH ₃
	H-OH	15.7	OH ⁻
	H-OCH ₂ CH ₃	16	OCH ₂ CH ₃ ⁻
	H-C≡C-H	25	C≡C-H ⁻
	H-NH ₂	38	NH ₂ ⁻
	H-CH ₃	~50	CH ₃ ⁻

Example of a favorable acid-base reaction:



pKa ~8

pKa ~15

Here, we're going from a stronger acid (HCl, pKa ~8) to a weaker acid (H₂O, pKa 15) so this acid-base reaction is favorable.

Omissions, Mistakes, Suggestions?

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Introduction to Resonance

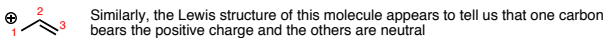
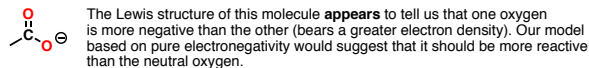
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masterorganicchemistry.com
 May 2012. Version 1.0

Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

1. What is Resonance?

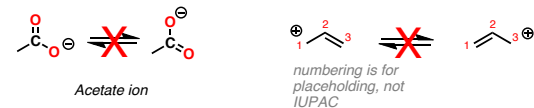
The true distribution of electrons in a molecule cannot always be accurately depicted through chemical formulae. Resonance is a tool to show this.

So what about applying this concept to these molecules?

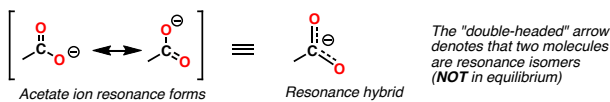


In reality, the charge in both of these molecules is distributed equally between the different ends

Important: this is **not** an equilibrium!



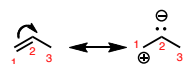
Instead, the "true" structure of the molecule is a **hybrid** of these two forms.



2. How to Use Curved Arrows

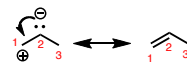
There are only **three** legal "moves" we can do with curved arrows to show resonance.

π Bond \rightarrow Lone pair



Tail is at π bond between C1 and C2
 Head is at C2
 Arrow shows movement of electrons from the C1-C2 π bond to become a lone pair on carbon 2.

Lone pair \rightarrow π Bond



Tail is on lone pair of C2
 Head is between C1 and C2.
 Arrow shows movement of electrons from the lone pair on C2 to become a π bond between C1 and C2

π Bond \rightarrow π Bond

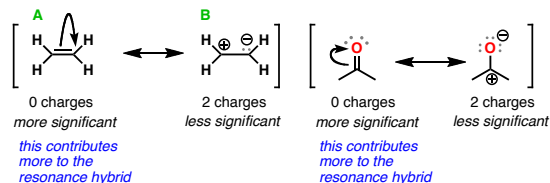


Tail is at π bond between C1 and C2
 Head is at π bond between C2 and C3
 Arrow shows movement of electrons from the C1-C2 π bond (which breaks) to the C2-C3 π bond (which forms).

3. When Resonance Forms are Not Equal: How to tell which are most stable?

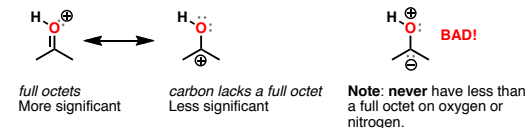
Rule #1 - The Fewer Charges, the Better

The more stable a resonance form is, the more it will contribute to the resonance hybrid.



Rule #2 - Full Octets Are Preferred

A resonance form in which all atoms have a full octet is more stable than one where at least one atom lacks a full octet.



Rule #3 - Place a negative charge on the most electronegative atom

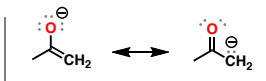
"Amide anions"



negative charge is on least basic atom (O)

negative charge is on most basic atom (N)

"Enolates"



negative charge is on least basic atom (O)

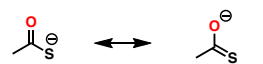
negative charge is on most basic atom (C)

If given a choice, the best place to put a negative charge on a resonance form is on the **least basic atom**

Use acidity trends to determine the "best" resonance form in these cases

- Electronegativity
- Polarizability
- Induction
- Orbitals (hybridization)

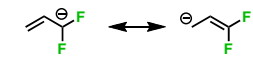
Polarizability



negative charge is on least basic atom (S)

negative charge is on most basic atom (O)

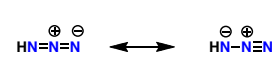
Electron withdrawing groups



negative charge is on least basic atom

negative charge is on most basic atom

Orbitals



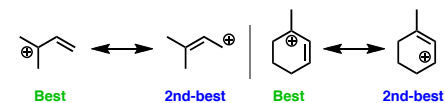
negative charge is on least basic atom (sp² hybridized nitrogen)

negative charge is on most basic atom (sp³ hybridized nitrogen)

Rule #4 Stabilizing positive charge

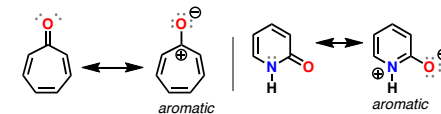
If the resonance form has a positive charge, make sure all octets are full (see rule #2).

If all octets are **not** full, place positive charge on the atom best able to support a carbocation (i.e. the **most substituted carbon**)



Rule #5 - Aromaticity (Org 2 Topic)

A resonance form that is aromatic will have a disproportionate contribution to the resonance hybrid

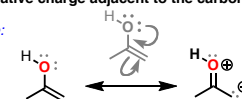


4. Applications of Resonance

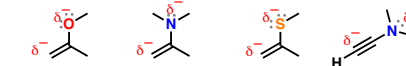
1) π -donation

Alkenes attached to an atom with a lone pair such as O, N, S, etc. (often called "heteroatoms") have an **important resonance form with a negative charge adjacent to the carbon-atom bond**.

Example:

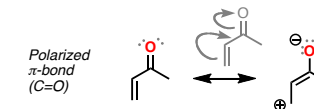


This effect makes these alkenes more nucleophilic (electron-rich).

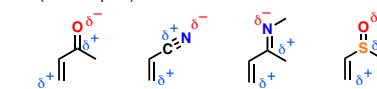


2) π -accepting

When double bonds are connected to a **polarized π bond**, the molecule will have a significant resonance form where there is positive charge on the adjacent carbon.



This effect makes these alkenes more electrophilic (electron-poor)



Omissions, Mistakes, Suggestions?







james@masterorganicchemistry.com

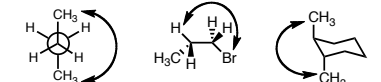
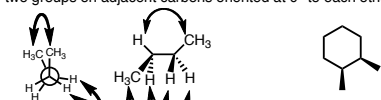
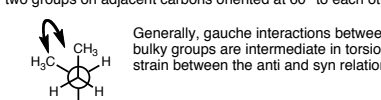
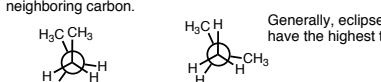
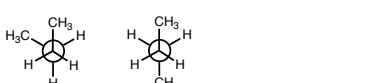
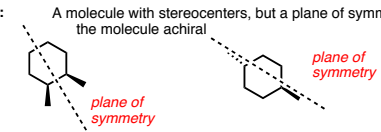
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









Introduction to Stereochemistry

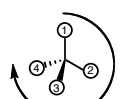
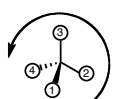
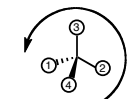
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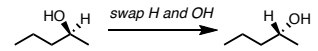
Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

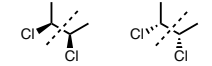
Symbols	Example
 "wedge" denotes a group pointing "out of the page"	
 "dash" denotes a group pointing "into the page"	
 "squiggle" undefined (mixture of stereoisomers) Rarely used.	

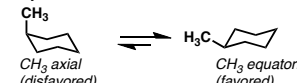
Terms	
anti	two groups on adjacent carbons oriented at 180° to each other 
syn	two groups on adjacent carbons oriented at 0° to each other 
gauche	two groups on adjacent carbons oriented at 60° to each other  Generally, gauche interactions between bulky groups are intermediate in torsional strain between the anti and syn relations
eclipsed	when all 3 substituents overlap completely with all 3 substituents on a neighboring carbon.  Generally, eclipsed conformations have the highest torsional strain
staggered	refers to the orientation when all 3 substituents on a carbon are arranged at a 60 deg angle to all 3 substituents on another carbon. 
isomers	two molecules with the same molecular formula, but different in their structures
configuration	the 3-D arrangement of bonds around a carbon.
racemic mixture	a 50:50 mixture of two enantiomers
Meso compound:	A molecule with stereocenters, but a plane of symmetry that makes the molecule achiral  plane of symmetry
Cis	on the same side of a double bond or ring
Trans	on opposite sides of a double bond or ring.
Chiral molecule	a molecule with an enantiomer
Stereocenter	Has 4 different constituents
Torsional Strain	Strain that arises from the proximity of bonds (and the electrons in them) - generally eclipsing

Types of isomers					Isomers: same empirical formula, different structural formula.	
Type	Connectivity?	Configuration?	3-D Shape?	Physical properties	Example	
Conformational isomers	Same	Same	Different	Identical, as long as they can interconvert through bond rotation		
Constitutional isomers	Different			Different boiling points, melting points, and other physical properties		
Stereoisomers	Same	Different				
Enantiomers	→	Stereoisomers that are non-superimposable mirror images		Identical. Differ in optical rotation		
Diastereomers	→	Stereoisomers that are NOT non-superimposable mirror images		Different boiling points, melting points, etc.		
Note: another name for stereoisomers is "configurational isomers": they have the same connectivity, but differ in the configurations of the carbons.						

The R,S convention	
Rank according to atomic weight Put #4 ranked substituent in back.	What if #4 is in the front? One approach is to trace 1,2 and 3 as you normally would. Then flip!
 1,2,3 goes CW: R	 1,2,3 goes CCW: S
 1,2, and 3 go clockwise. 4 ranked substituent is in front. Therefore, flip: this is R!	

The single swap rule:
Single swap rule: switching any two groups on a stereocenter will flip (R) to (S) and vice versa.
 swap H and OH
(S)-2-pentanol → (R)-2-pentanol
works for any other two groups as well (e.g. OH and CH ₃ , H and CH ₃ , OH and CH ₂ CH ₂ CH ₃ , etc.)

Recognizing Enantiomers/Diastereomers Using Only Their Names
Diastereomers: same name, but R/S designations are not exactly opposite
Example: (2S, 3R)-2-bromo-3-chlorobutane and (2S, 3S)-2-bromo-3-chlorobutane (2S, 3S, 4S, 5R)-2-bromo-3-chloro-4-methyl-5-propyldecane and (2S, 3R, 4R, 5S)-2-bromo-3-chloro-4-methyl-5-propyldecane also, diastereomers will have the same name, but differ in E/Z (or cis/trans). e.g. (Z)-2-butene and (E)-2-butene. cis-1,2-dimethylcyclohexane and trans-1,2-dimethylcyclohexane
Enantiomers: same name, but have all stereocenters have opposite R/S designations
Example: (R)-2-butanol, (S)-2-butanol (2S, 3R)-2-bromo-3-chlorobutane and (2R, 3S)-2-bromo-3-chlorobutane (2S, 3S, 4S, 5R)-2-bromo-3-chloro-4-methyl-5-propyldecane and (2R, 3R, 4R, 5S)-2-bromo-3-chloro-4-methyl-5-propyldecane
Important exception: If the molecule has a mirror plane, then it is meso , and the two "enantiomers" are in fact the same molecule
Example: (2R,3S)-2,3-dichlorobutane and (2S,3R)-2,3-dichlorobutane are the same  mirror plane

Cyclohexane chair conformations
in the cyclohexane chair conformation: all C-C bonds staggered  CH ₃ axial (disfavored) ↔ CH ₃ equatorial (favored)
Chair flips: all axial groups become equatorial, and all equatorial groups become axial. BUT all groups that are "up" stay "up" and all groups that are "down" stay "down". Bulky groups prefer the equatorial position. The bulkier the group, the greater the energy difference will be. Bulkiness: tertiary C (3°) > secondary C (2°) > primary C (1°), methyl (CH ₃) >> H The energy difference between two different chair forms (in ΔG) is related to the equilibrium constant by K = e ^{-(ΔG/RT)}

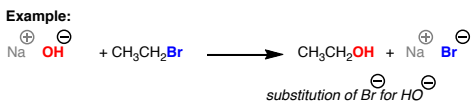
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Substitution and Elimination - Page 1 of 2

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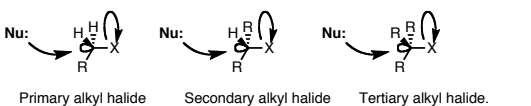
Nucleophilic Substitution Bimolecular (S_N2)

Key points:
 • Nucleophilic substitution is swapping of one functional group for another
 • Structure of alkyl halide is key: primary > secondary >> tertiary
 • Mechanism proceeds through backside attack
 • Stereochemistry proceeds with **inversion**
 • Rate depends on concentration of **both** nucleophile and alkyl halide
 • Reaction occurs faster in **polar aprotic solvents**



The S_N2 Reaction is Sensitive to Steric Effects.

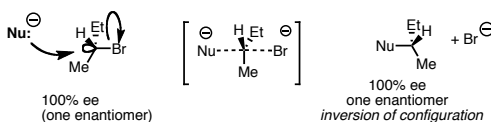
In the S_N2 the electrophile is the antibonding orbital, as you populate the antibonding orbital with electrons you weaken the bond to the leaving group



Fast **Intermediate** **So slow it doesn't happen.**

The S_N2 Mechanism: The Backside Attack Leads to Inversion

The nucleophile (electron pair) attacks the empty antibonding orbital which is on the **backside** of the carbon-leaving group bond



The Rate Limiting Step of the S_N2 Is Bimolecular

$$\text{Rate} = k [\text{Substrate}] [\text{Nucleophile}]$$

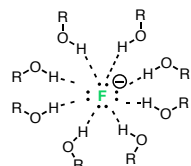
Rate	[R-X]	[Nu]	
x	1	1	<i>Doubling the concentration of either component will double the rate.</i>
2x	2	1	
2x	1	2	
4x	2	2	

Polar aprotic solvents are the best for the S_N2

Because of the charges involved in the S_N2 mechanism polar solvents are better than nonpolar solvents.

Two types of polar solvents: polar protic (have O-H or N-H bonds)
 Polar aprotic (has dipoles but cannot hydrogen bond).

Why? **Polar protic solvents make a "jacket" around nucleophiles through hydrogen bonding, decreasing the reactivity of the nucleophile.**

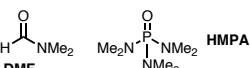


Examples of polar protic solvents:

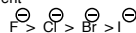
H₂O, CH₃OH, CH₃CH₂OH, CH₃COOH, i-PrOH, CH₃CH₂CH₂NH₂

Polar aprotic solvents:

DMSO Acetone Acetonitrile



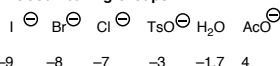
Note that in polar aprotic solvent reactivity of halides **reverses** because there is no hydrogen bonding to solvent



What Makes A Good Leaving Group?

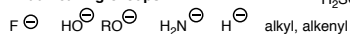
One main thing: **Good Leaving Groups are Weak Bases**

Good Leaving Groups



pKa of conjugate acid

Bad Leaving Groups

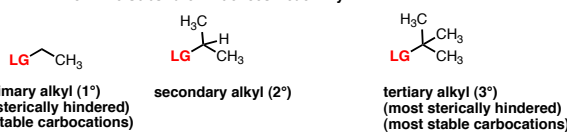


pKa of conjugate acid

With the exception of F[⊖], these are all **strong bases** *Fluorine forms very strong bonds and is thus a very bad leaving group*

Note how HO[⊖] is not a good leaving group but H₂O is: therefore, HO can be made into a good leaving group by adding a strong acid

How The Substrate Influences Reactivity



S_N2 (unhindered) E₂ (provided a proton is on the β carbon)

S_N1/E1: not observed

S_N2 (slower than primary) E₂ (provided a proton is on the β carbon)

S_N1/E1: can occur when LG is OH and a strong acid is added **watch out for rearrangements!** Generally poor when LG is a halide.

S_N2: Not observed E₂ (provided a proton is on the β carbon)

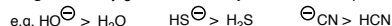
S_N1/E1: strongly favored if LG = halide, good reaction if LG = OH, requires acid

Note that hybridization in all cases here is sp³

These reactions do not occur for alkenyl and alkynyl halides under normal conditions.

What Factors Influence Nucleophilicity?

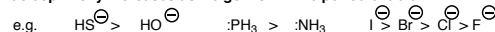
1. **Charge:** The conjugate base is always a better nucleophile than the conjugate acid



2. **Electronegativity:** The less tightly held a pair of electrons are, the more readily they can be donated.

Nucleophilicity increases as we go to the LEFT along the periodic table: C > N > O > F:

Nucleophilicity increases as we go DOWN the periodic table:



Another term for how "tightly held" electrons are is **polarizability.**

Nucleophilicity increases with polarizability.

3. **Solvent:** Polar protic solvents will hydrogen bond with nucleophiles (see column on left) which decreases their nucleophilicity.

• More polarizable nucleophiles will be less affected by polar protic solvents, since hydrogen bonding ability is greatest for most electronegative nucleophiles.

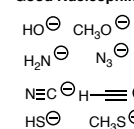
• Polar aprotic solvents **will not** hydrogen bond; therefore they are good solvents for the S_N2

Note that in polar aprotic solvent reactivity of halides **reverses** because there is no hydrogen bonding to solvent

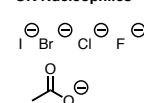
4. **Steric bulk:** If a sterically bulky group surrounds the nucleophilic atom, this will decrease its nucleophilicity.

Example: CH₃O[⊖] is more nucleophilic than for this reason

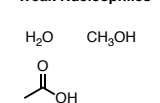
"Good Nucleophiles"



"OK Nucleophiles"



"Weak Nucleophiles"



"Non-Nucleophiles"



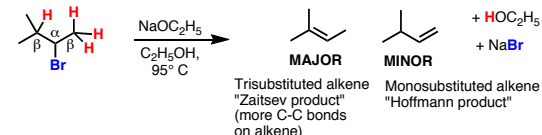
These are examples only, not comprehensive

Elimination, Bimolecular (E₂)

Key points:

- Elimination is the loss of a leaving group from the α-carbon and a hydrogen from the β-carbon resulting in the **formation of a double bond**
- The major product is the more substituted alkene (Zaitsev product) and the bulkiest groups will be trans
- Use of a bulky base will result in formation of more of the less substituted alkene
- Requires strong base
- The leaving group and the hydrogen must be oriented **anti-periplanar**
- Can occur with primary, secondary, tertiary substrates **provided there is a hydrogen on the β-carbon that can orient itself anti to the leaving group**
- Rate determining step is **bimolecular**
- Solvent is generally polar protic
- Reaction is favored with heat

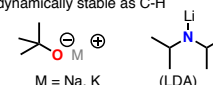
Example: *Elimination of HBr*



The Major Product is the Most Substituted Alkene

Why? Because alkenes become more thermodynamically stable as C-H bonds are replaced with C-C bonds

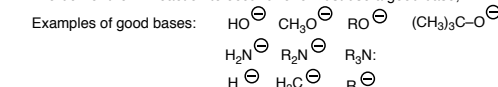
Exception: Bulky bases will give less of the Zaitsev product because deprotonation on the less substituted carbon becomes favored.



Exception #2 is when the leaving group is bulky (e.g. NR₃[⊖])

A Strong Base Is Required

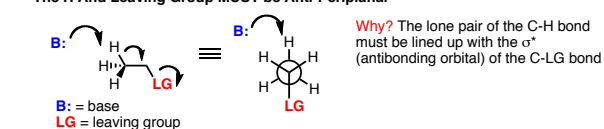
In order for the E₂ reaction to occur you must use a good base,



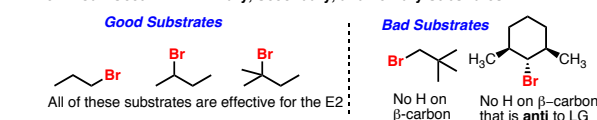
Basicity increases going to the left along the periodic table: C > N > O > F:

Basicity increases going up the periodic table: F[⊖] > Cl[⊖] > Br[⊖] > I[⊖]

The H And Leaving Group MUST be Anti-Periplanar



The E₂ Can Occur With Primary, Secondary, and Tertiary Substrates



The Rate Limiting Step of the E₂ Is Bimolecular

$$\text{Rate} = k [\text{Substrate}] [\text{Base}]$$

Rate	[R-LG]	[B]	
x	1	1	<i>Doubling the concentration of either component will double the rate.</i>
2x	2	1	
2x	1	2	
4x	2	2	

Solvent is Generally Polar Protic - not in all instances, but polar protic will favor E₂ in cases where it competes with the S_N2 (see left)

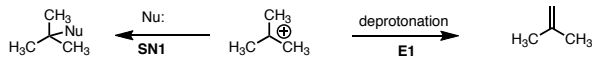
The E₂ Is Favored By Heat

- Keep this in mind when considering substrates that could go E₂ or S_N2
- Heat favors elimination due to higher entropy

Substitution and Elimination - Page 2 of 2

Substitution Nucleophilic Unimolecular (S_N1) and Elimination, Unimolecular (E1)

- These reactions have a **lot** in common and often occur together.
- They represent different fates of the same carbocation



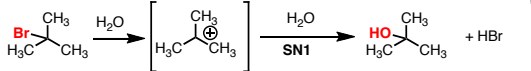
The S_N1 is nucleophilic attack at the carbocation

The E1 is deprotonation of the carbon adjacent to the carbocation

What the S_N1 and E1 Have In Common

- The rate-limiting step is **formation of a carbocation**
- The rate-limiting step is **unimolecular** and **only** depends on the concentration of the **substrate** (e.g. alkyl halide)
- The rate is also proportional to the **stability of the carbocation** (see column on right)
- Tertiary carbocations > secondary carbocations >> primary carbocations
- Carbocations are **ions** (polar) so according to "like dissolves like", polar solvents are favored. In particular, **polar protic solvents** help

S_N1 Example:

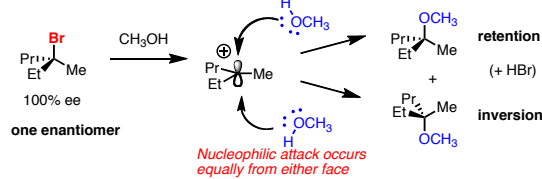


The S_N1 involves two steps that occur in sequence:

- 1) departure of the leaving group 2) attack of the nucleophile at the carbocation
- The reaction proceeds with **weak/moderate nucleophiles** (e.g. H₂O,
 - If the carbon is chiral, the chiral carbon becomes **racemized**
 - **The S_N1 is always accompanied by some E1** (~95:5)

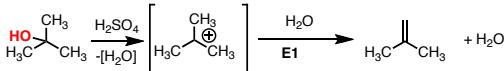
Why does the S_N1 lead to a mixture of stereoisomers?

Because the carbocation is flat!



A racemic mixture is obtained

E1 Example:



The E1 also involves two steps that occur in sequence:

- 1) departure of the leaving group 2) deprotonation of the carbon adjacent to the carbocation

- The proton is removed by a **weak base**
- The **more substituted double bond** is favored (Zaitsev product)
- The major product will have the bulkiest groups *trans* to each other across the double bond
- The reaction is favored by **heat**
- The base is the solvent or the counterion of the aci

The Rate Limiting Step of the S_N1 and E1 Is Unimolecular

$$\text{Rate} = k [\text{Substrate}] [\text{Nucleophile/Base}]$$

Rate	[R-X]	[Nu]	
x	1	1	Doubling the concentration of the substrate doubles the rate.
2x	2	1	Doubling the concentration of the nucleophile/base has no effect
x	1	2	
2x	2	2	

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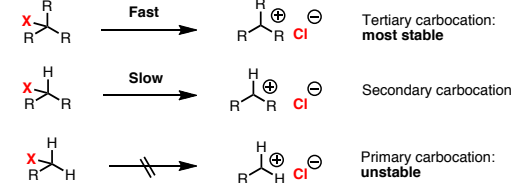
All About Carbocations

- Carbocations are flat (trigonal planar) - the central carbon is sp² hybridized
- They are stabilized by 1) increasing substitution at carbon and 2) resonance
- Tertiary carbocations > secondary carbocations >> primary carbocations
- They can be formed in several ways: 1) dissolving a tertiary alkyl halide in a polar protic solvent 2) adding acid to a secondary or tertiary alcohol
- They are more stable in polar solvents
- Secondary carbocations can rearrange to tertiary carbocations through hydride or alkyl shifts

What influences carbocation stability?

1) Carbon substituents stabilize carbocations

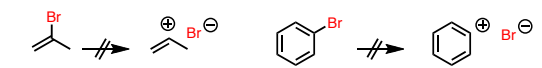
Tertiary > secondary >> primary



2) Resonance stabilizes carbocations



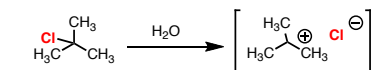
3) Vinyl and aryl carbocations are very unstable



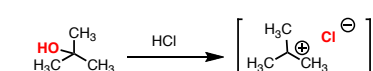
How are Carbocations formed?

Three ways.

1) dissolve a tertiary alkyl halide in a polar protic solvent.

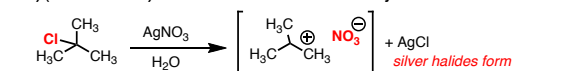


2) Add an acid to a secondary or tertiary alcohol:



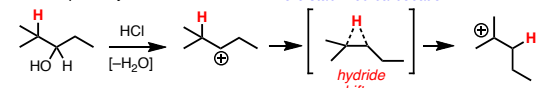
Can also be done for secondary alcohols - watch out for rearrangements!

3) (Less common) - addition of silver salts to an alkyl halide

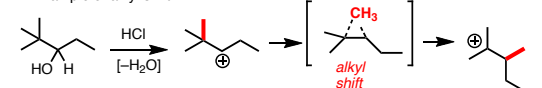


Secondary Carbocations Can Rearrange

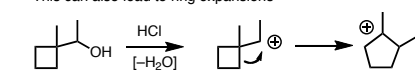
Driving force is formation of more stabilized carbocation



Example of alkyl shift:

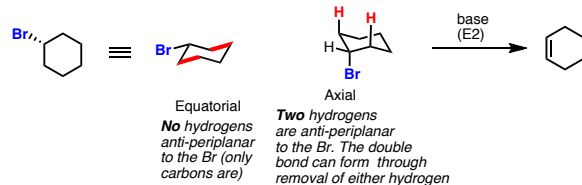


This can also lead to ring expansions

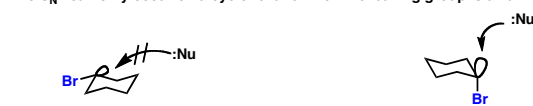


Further Notes on the S_N2 / E₂ (more advanced topics)

The E2 can only occur when the leaving group is axial:



The S_N2 can only occur on a cyclohexane when the leaving group is axial

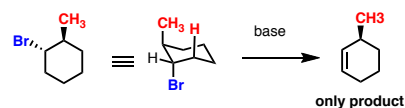


Why?

When the leaving group is equatorial, the antibonding orbital points *into* the center of the ring, making it sterically inaccessible

When the leaving group is **axial**, the antibonding orbital points **above** the ring, where it can be attacked by a nucleophile

If there is no hydrogen *anti* to the leaving group, then the double bond cannot form



One hydrogen are anti-periplanar to the Br. Therefore **only one** double bond can form

Relative Rate of S_N2 Reactions where X is a leaving group

CH ₃ X	30
CH ₃ CH ₂ CH ₂ -X	1
CH ₃ CH ₂ -X	0.4
	0.025
	1 x 10 ⁻⁵

rate decreases with increasing steric bulk

note that even though X is on a primary carbon, it is adjacent to a quaternary carbon, which results in a very slow reaction

Additions to Alkenes

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Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

Reaction		"Regiochemistry"	"Stereochemistry"	
Hydroboration		Anti-Markovnikov	syn addition	Sometimes you might see BH ₃ ·THF or B ₂ H ₆ used here: it's the same reagent in a slightly different form. The base (can be NaOH, KOH, identity unimportant) helps make H ₂ O ₂ more reactive. The reaction is <i>anti</i> -Markovnikov because the H-B bond is polarized toward H (electronegativity of H = 2.2, B = 2.0) - the H adds to the carbon best able to stabilize positive charge (i.e. the most substituted one).
Oxymercuration		Markovnikov	stereorandom	This reaction goes through 3-membered "mercurinium" ion. The NaBH ₄ step removes the mercury. While the addition is <i>anti</i> , the overall reaction is <i>stereorandom</i> because this step involves a carbon based free radical (usually not discussed). Alternatively, an alcohol used in place of water will produce an ether.
Acid-catalyzed addition of H ₂ O (hydration)		Markovnikov	stereorandom	Strong acid protonates the alkene, generating free carbocation. Watch out for possibility of rearrangements when a tertiary carbocation could be generated through a 1,2 shift. HSO ₄ ⁻ anion is not strongly nucleophilic, hence it does not add. Stereorandom because of the free carbocation.
Addition of HX		Markovnikov	stereorandom	HCl and HBr (as well as HI, not pictured) protonate the alkene to give a free carbocation which can then be trapped by the halide anion. Stereorandom
Addition of HX		Markovnikov	stereorandom	
Bromination		N/A	anti addition	<p>Bromonium ion mechanism Br⁺ or H₂O/ROH depending on solvent</p> <p>The key detail in these reactions is solvent: water and alcohol solvents will form the <i>halohydrin</i> products (the ones containing the OH and Br). All other solvents (you might see CCl₄, CHCl₃, hexane, etc.) provide the dibromide.</p>
Halohydrin Formation		Markovnikov	anti addition	
Chlorination		N/A	anti addition	As with bromination, above. Although not depicted, use of water or alcohol as solvent will also lead to formation of the halohydrin product (also anti).
Dihydroxylation		N/A	syn addition	Osmium is a transition metal. The tools won't be given in this course to fully understand how this reaction works. Occasionally a second reagent like NaHSO ₃ , H ₂ S, or Na ₂ S ₂ O ₃ is also given as a reactant in this reaction - minor detail, it's used to remove the osmium from the hydroxyl groups.
Dihydroxylation		N/A	syn addition	Keywords are "cold, dilute". NOTE: If "heat" or "acid" is mentioned in the conditions, the diol will be cleaved to provide carbonyl compounds (same reaction as ozonolysis with oxidative workup , below).
Epoxidation		N/A	syn addition	RCO ₃ H is a per-acid. A common peracid for this reaction is <i>m</i> -CPBA (<i>m</i> -chloroperbenzoic acid). If H ₃ O ⁺ , heat is written afterwards, this is opening of the epoxide to give the diol (<i>anti</i> -selective)
Hydrogenation		N/A	syn addition	The catalyst can vary - you might see Pd/C or Ni as well. All provide the same product with the same stereochemistry.
Radical addition of HBr		Anti-Markovnikov	stereorandom	Peroxides generate the Br· radical, which adds to the double bond in the way that will generate the most stable radical (i.e. the radical will go on to the most substituted carbon). This explains the selectivity for the anti-Markovnikov product. It's stereorandom because it goes through a free radical process.
Ozonolysis (Reductive workup)				Reductive workup: Zinc (Zn), or dimethyl sulfide (DMS, Me ₂ S) is a reducing agent. It reduces excess ozone, allowing for isolation of the aldehyde.
Ozonolysis (Oxidative Workup)				Oxidative workup: Hydrogen peroxide is used to obtain the carboxylic acid instead of the aldehyde.
Cyclopropanation		N/A	syn addition	This reaction goes through addition of a <i>carbene</i> (or <i>carbenoid</i>) to the double bond. The reaction is stereospecific. Another set of conditions to provide a cyclopropane are CHCl ₃ with strong base, which makes the dichlorocyclopropane.

Omissions, Mistakes, Suggestions?

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Summary of Alkene Reaction Patterns

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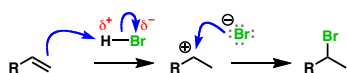
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Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

Pattern 1: The Carbocation Pathway

These reactions proceed through attack of an alkene upon an acid, which results in formation of a carbocation. The carbocation is then attacked by a nucleophile, which can occur from either face of the carbocation.

Example: addition of H-Br to alkenes



Regiochemistry: Markovnikov-Selective. In other words, **hydrogen** ends up bonded to the **less** substituted carbon of the alkene, and (in this case) Br ends up attached to the **most** substituted end of the alkene.

Stereochemistry: Random. In other words the H and the other group (e.g. Br) do not add exclusively "syn" or "anti"

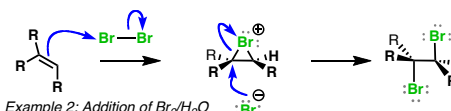
Reactions in this category: Addition of H-Cl
Addition of H-Br
Addition of H-I
Addition of H₃O⁺
(sometimes written as H₂O / H₂SO₄)

Note: since this reaction goes through a carbocation, **rearrangements are possible.**

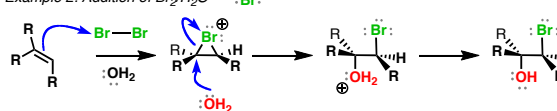
Pattern 2: The 3-Membered Ring Pathway

These reactions proceed through formation of a 3-membered ring cation, which is then attacked from the backside by a nucleophile, on the most substituted carbon.

Example 1: addition of Br₂ to alkenes



Example 2: Addition of Br₂/H₂O



Regiochemistry: Markovnikov-Selective (where appropriate). For example, with Br₂/H₂O, water adds to the **most** substituted carbon of the alkene.

Stereochemistry: Anti. The two groups add to opposite faces of the alkene.

Reactions in this category: Addition of halogens: Cl-Cl, Br-Br, I-I
Addition of halogens in the presence of nucleophilic solvents: Br₂/H₂O, Br₂/R-OH
Oxymercuration: Hg(OAc)₂, H₂O, then NaBH₄
Oxymercuration (with an alcohol as solvent) Hg(OAc)₂, ROH, then NaBH₄
Opening of epoxides under acidic conditions

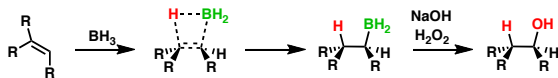
Pattern 3: The "Concerted" Pathway

Although these reactions do not share a common mechanism, they each form two new bonds on the same side of the alkene, a consequence of a "concerted" reaction mechanism.

Example 1 - hydrogenation



Example 2 - hydroboration



Regiochemistry: anti-Markovnikov-Selective (for BH₃). Note how hydrogen adds to the **most** substituted end of the alkene, and OH ends up on the **least** substituted end.

Stereochemistry: Syn. The two bonds are formed on the same side of the alkene

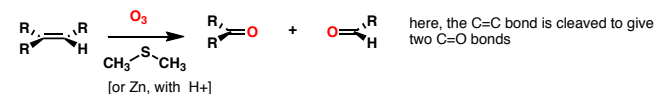
Reactions in this category: Hydroboration: BH₃, then H₂O₂/NaOH
Hydrogenation: Pd/C, H₂
Epoxidation: RCO₃H (e.g. mCPBA)
Dihydroxylation: OsO₄
Cyclopropanation: CH₂I₂, Zn-Cu
Dichlorocyclopropanation CHCl₃, NaOH

Pattern 4: Two miscellaneous reactions

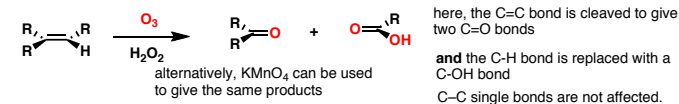
Two reaction patterns of note

Oxidative cleavage: Addition of an oxidant (such as ozone, O₃ leads to cleavage of the carbon-carbon double bond and formation of two new carbon-oxygen double bonds.

Ozonolysis (Reductive workup)

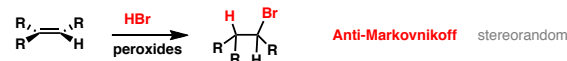


Ozonolysis (Oxidative Workup)



Radical addition of HBr

Here, addition of HBr to alkenes in the presence of a radical initiator (peroxides) leads to **anti-Markovnikov** addition of H-Br to the alkene; the Br adds to the least substituted carbon of the alkene, and the H adds to the most substituted.



Regiochemistry: anti-Markovnikov-Selective. Note that the Br adds to the less substituted end of the alkene.

Stereochemistry: Random. H and Br do not add exclusively "syn" or "anti" across the alkene.

Omissions, Mistakes, Suggestions?

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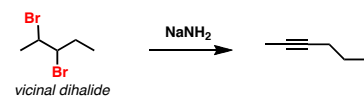
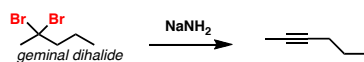
Introduction to Reactions of Alkynes

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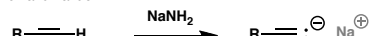
Synthesis of Alkynes

Elimination of Hydrogen halides



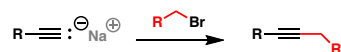
NaNH_2 is a strong base and will lead to elimination of HBr. According to Zaitsev's rule, the most substituted alkyne should form.

Deprotonation



Alkynes are relatively acidic [$\text{pK}_a = 25$]. A strong base like NaNH_2 will remove the C-H to provide the acetylide anion. The resulting anion is an excellent nucleophile.

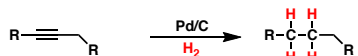
Reaction with Electrophiles



Deprotonated alkynes are great nucleophiles and will react with alkyl halides in $\text{S}_\text{N}2$ reactions

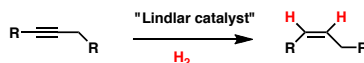
Addition Reactions to Alkynes

Hydrogenation



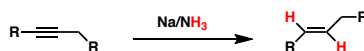
Can also use Pt, Ni, etc. as hydrogenation catalysts. These will reduce alkynes to alkanes.

Partial Hydrogenation



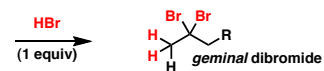
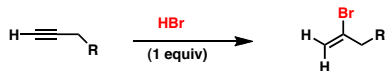
Lindlar catalyst is a "poisoned" palladium catalyst for the partial hydrogenation of alkynes. It contains palladium (Pd), lead (Pb), and quinoline, and is selective for the *cis*-alkene. Sometimes also see "Pd, CaCO_3 ".

Reduction (Dissolving metal reduction)



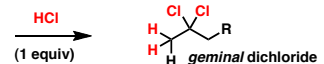
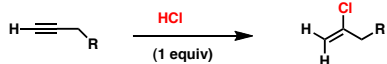
Sodium (Na) is a very strong reducing agent, and will reduce alkynes (but not alkenes). Selective for the *trans*-alkene. The NH_3 provides the hydrogens in the final product.

Addition of Hydrogen bromide to form gem-dibromides



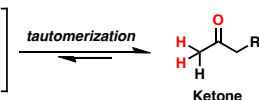
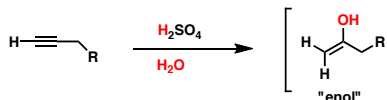
Markovnikov

Addition of Hydrogen chloride to form gem-dichlorides



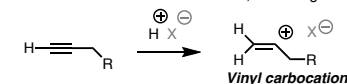
Markovnikov

Acid-catalyzed Hydration to provide ketones



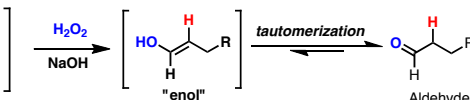
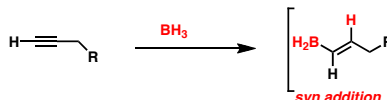
Markovnikov

The key intermediate in all three of these reactions is an intermediate *vinyl carbocation* which is formed upon protonation of the alkyne by strong acid. The carbocation forms on the more substituted side of the carbon, according to Markovnikov's rule.



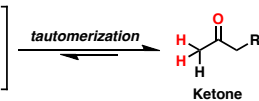
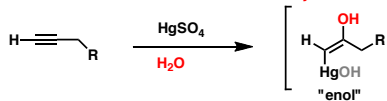
This carbocation will then be attacked by the nucleophile in question, leading to the formation of substituted products.

Hydroboration to form aldehydes



Anti-Markovnikov

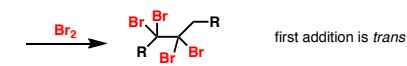
Oxymercuration to provide ketones



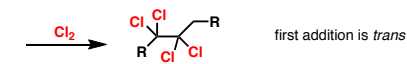
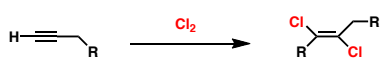
Markovnikov

The first intermediate in each of these three reactions is a cyclic ion, which undergoes attack by a nucleophile at the more substituted carbon (Markovnikov)

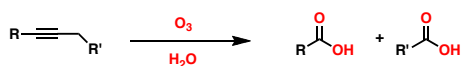
Bromination



Chlorination



Ozonolysis



If $\text{R} = \text{H}$, then one product will be CO_2 [$\text{O}=\text{C}=\text{O}$]

Omissions, Mistakes, Suggestions?

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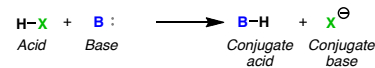
Common Reaction Patterns (Org 1)

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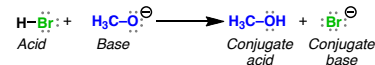
Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

Pattern #1 - Acid Base Reactions

Generic example:



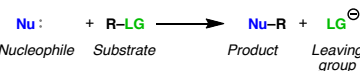
Specific example:



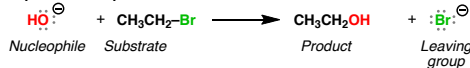
Notes: Favorable acid base reactions will involve a stronger acid and a stronger base reacting to give a weaker acid and weaker base. Relative strengths of acids and bases can be obtained through consulting a pKa table

Pattern #2 - Nucleophilic Substitution Reactions

Generic example:



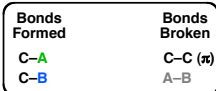
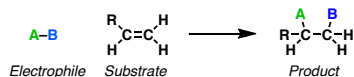
Specific example:



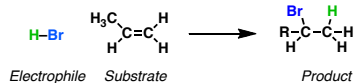
Notes: Substitution reactions can proceed through two pathways. The S_N2 pathway is concerted, and C-Nu bond formation occurs at the same time as C-LG bond breakage. Steric hindrance is paramount. In the S_N1 pathway, the leaving group leaves first to give a carbocation, which is then attacked by a nucleophile. In the S_N1 pathway rearrangements can occur depending on carbocation stability. Good nucleophiles tend to have a loosely held lone pair of electrons. Good leaving groups are weak bases.

Pattern #3 - Electrophilic Addition

Generic example:



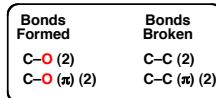
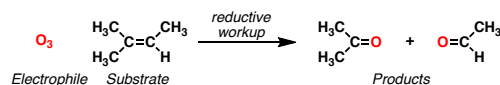
Specific example:



Notes: Note that every addition reaction involves breakage of a C-C π bond and formation of two new single bonds to C. For stepwise addition reactions, the tendency will be to place positive charge on the carbon best able to accept it (i.e. the carbon attached to the most carbon atoms). This gives rise to "Markovnikov's rule", where H always adds to the least substituted carbon. [Notable exception is hydroboration-oxidation, which is "anti-Markovnikov."] Stereochemistry of the two new bonds can be syn, anti, or a mixture of both, depending on mechanistic pathway.

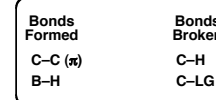
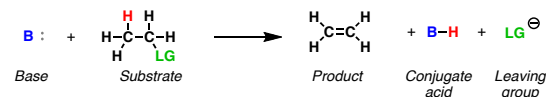
Pattern #7 - Oxidative Cleavage

Specific example:

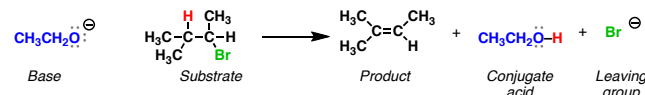


Pattern #4 - Elimination Reactions

Generic example:



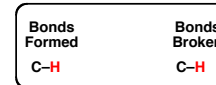
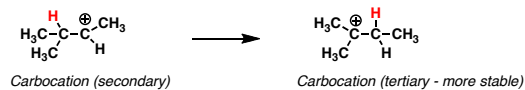
Specific example:



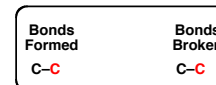
Notes: Note that every elimination reaction involves the formation of a new C-C π bond. Since alkene stability increases as the number of attached carbons increases, eliminations tend to favor formation of the more substituted alkene ("Zaitsev's rule"). An important exception is when more hindered bases (such as *t*-BuO⁻) are used, which gives the less substituted alkene. Two dominant mechanisms for elimination are the concerted pathway (E2) which occurs in one step, requiring *anti*-periplanar arrangement of H and leaving group. The second pathway is the stepwise E1 pathway, which occurs with loss of leaving group followed by deprotonation. A third (rare) pathway involves deprotonation, followed by loss of leaving group, called the E1-conjugate base (E1cb) pathway. All elimination reactions tend to be favored by heat.

Pattern #5 - Carbocation Rearrangements

Hydride shift example:



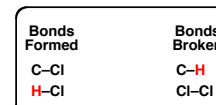
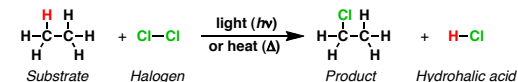
Alkyl shift example:



Notes: Rearrangements are favorable when the migration of hydride or alkyl leads to a more stable carbocation. Recall that carbocation stability proceeds in the order tertiary > secondary >> primary. Resonance can also stabilize carbocations. Hydrogen tends to migrate preferentially over carbon. Relief of ring strain can also be a major contributing factor with rearrangement reactions.

Pattern #6 - Free Radical Substitution

Generic example:



Notes: Free radical reactions proceed in three steps: initiation, propagation, and termination. Light (hν) or heat (Δ) is used to initiate the reaction. Chlorine is quite unselective relative to bromine, which will only react with tertiary and allylic/benzylic C-H bonds (which are weakest). Iodine does not perform this reaction. If multiple equivalents of chlorine are present, chlorination can occur multiple times until all C-H bonds are replaced with C-Cl.

Notes: Oxidation of C-C multiple bonds (π bonds) can be achieved with ozone (O₃) or potassium permanganate (KMnO₄). With ozone, reductive workup (using Zn or dimethyl sulfide) leaves all C-H bonds intact. With KMnO₄ or O₃ using oxidative workup, sp² hybridized C-H bonds are oxidized to C-OH.

A different type of oxidative cleavage can occur with vicinal diols using NaIO₄, HIO₄, or Pb(OAc)₄. These oxidants will cleave C-C single bonds and form C=O (π) bonds

Omissions, Mistakes, Suggestions?

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Introduction to Chemical Reactivity, Nomenclature, Boiling Points, and Water Solubility

<http://masterorganicchemistry.com>

Reactant #1	Reactant #2	Product	Example	Notes
Alkene	Pd/C + H ₂	Alkane		cis addition (hydrogens go on same side of alkene)
Amine	Acid	Ammonium salt		amines but NOT amides (amides are not basic on nitrogen)
Carboxylic acid	Base	Salt (Carboxylate salt)		
Carboxylic acid	Alcohol, acid, heat	Ester		Alcohols used as solvent
Ester	Water, acid, heat	Carboxylic acid		This is the reverse of the above reaction. Here we use water as solvent.
Ester	Water, base	Carboxylic acid		This is called ester hydrolysis or saponification
Alkene	Water, acid	Alcohol		alcohol forms on most substituted carbon (Markovnikov rule) proceeds through carbocation
Alkene	Strong acid	Alkyl halide		halide adds to most substituted carbon (Markovnikov rule) proceeds through carbocation
Alkene	Br ₂	Dibromide		Results in trans product
Alcohol	K ₂ CrO ₇	Carboxylic acid (primary alcohol) OR ketone (secondary alcohol)		Note that secondary alcohols stop at the ketone stage
Alcohol	KMnO ₄	Carboxylic acid (primary alcohol)		
Alkane	Cl ₂ , hv (or peroxides)	Alkyl chloride		Free-radical reaction (number of new C-Cl bonds depends on # of equivalents)
Benzene derivative	Br ₂ , FeCl ₃	Aryl bromide		Also gives 1,4 (para) product but never 1,3 (meta) product
Thiol	"Oxidant"	Disulfide		

The Four Types of Intermolecular Bonding in Organic Chemistry

Name	Type of Interaction	Found in	Example	Strength	Notes
Ionic	Attraction between point charges	Salts		Strongest	gives rise to greatest water solubility (most polar) also highest boiling points
Hydrogen Bonding	Attraction between positively charged H and negatively charged O, N or F.	Water, alcohols, carboxylic acids, amides, amines		2nd strongest	2nd greatest for effect on water solubility and boiling points
Dipole-Dipole	Attraction between dipole moments caused by differences in electronegativity	ketones, aldehydes, esters, alkyl halides, etc. - any molecule with a strongly electronegative element (O, N, F, Cl, Br)		2nd weakest	Increases as electronegativity difference increases 3rd greatest for effect on water solubility and boiling points
Van Der Waals (London forces)	Attraction between temporary dipoles	Hydrocarbons	Name	Weakest	Increases with surface area (increasing length of carbon chains) worst for water solubility (least polar) best for solubility in non-polar solvents (e.g. pentane)

# Carbons	Root
1	Meth-
2	Eth-
3	Prop-
4	But-
5	Pent-
6	Hex-
7	Hept-
8	Oct-
9	Non-
10-	Dec-

Primary carbon: attached to ONE carbon atom

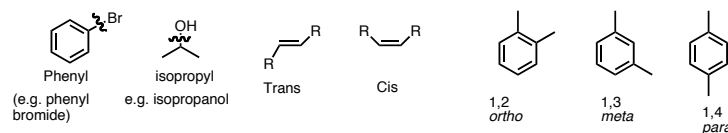
Secondary: attached to TWO carbon atoms

Tertiary: attached to THREE carbon atoms

Quaternary: attached to FOUR carbon atoms

Functional Group	Name	Example	Name
R-	Alkyl		Pentane
-OH	Hydroxyl		Pentanol or pentyl alcohol
-Cl, -Br, -F, -I	Halide		Pentyl chloride
-NH ₂	Amine		Pentylamine
R-O-R	Ether		Pentyl methyl ether
-SH	Thiol		Pentane thiol
C=C	Alkene		Pentene
R-C(=O)-H	Aldehyde		Pentanal
R-C(=O)-R	Ketone		Butyl methyl ketone OR 2-pentanone
R-C(=O)-OH	Carboxylic acid		Pentanoic acid
R-C(=O)-OR	Ester		Methyl pentanoate
R-C(=O)-NH ₂	Amide		N-methyl pentamide

Other important nomenclature terms to remember

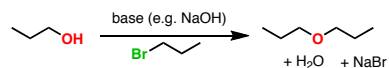


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Introduction to Alcohols and Ethers

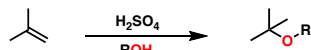
Ethers from alcohols and alkyl halides

The Williamson Ether synthesis: alcohol, base, alkyl halide (or tosylate)



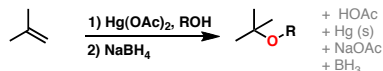
This is an S_N2 reaction; it works best for primary alkyl halides (and alkyl tosylates)

Ethers from alkenes



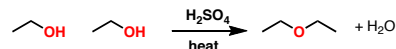
The reaction is similar to the hydration of alkenes with aqueous acid. Acid leads to the formation of a carbocation, which is then trapped by the alcohol as solvent.

Ethers from alkenes through oxymercuration



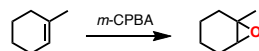
The reaction is similar to the hydration of alkenes with aqueous acid. Acid leads to the formation of a carbocation, which is then trapped by the alcohol as solvent.

Ethers from alcohols through dehydration



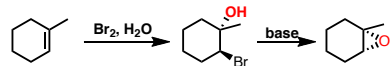
Strong acid (and heat) leads to protonation of the alcohol, followed by nucleophilic attack of a second molecule of alcohol to give the ether. Works only for the synthesis of symmetrical ethers.

Epoxides from alkenes



m-CPBA (meta-chloroperoxybenzoic acid, a peroxyacid) converts alkenes to epoxides, a cyclic ether.

Epoxides from halohydrins



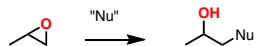
Formation of the halohydrin from the alkene is stereospecific for the trans product. Deprotonation of the alcohol by base results in S_N2 (with inversion) to give the epoxide.

Opening of epoxides

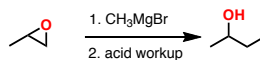
Due to ring strain, epoxides are highly reactive towards nucleophiles. They will react with nucleophiles under both acidic and basic conditions. However the patterns are different.

Under basic conditions

Under basic conditions, nucleophiles will attack epoxides at the **least sterically hindered position** (primary > secondary > tertiary). The reaction is essentially an S_N2 reaction



Example: reaction of epoxides with Grignard reagents

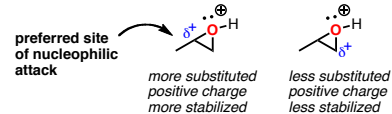


Under acidic conditions

Under acidic conditions, the epoxide oxygen is protonated:

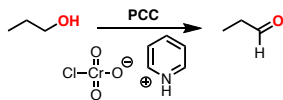


The nucleophile will attack the carbon best able to stabilize positive charge - which is the more substituted carbon. Just like Markovnikoff's rule!



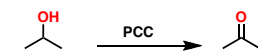
Oxidation of alcohols

Oxidation of primary alcohols to aldehydes



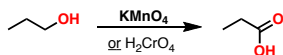
PCC = pyridinium chlorochromate

Oxidation of secondary alcohols to ketones



Can also use KMnO_4 or H_2CrO_4

Oxidation of primary alcohols to carboxylic acids



Another way of writing H_2CrO_4 is $\text{K}_2\text{Cr}_2\text{O}_7 / \text{H}_2\text{SO}_4$ or $\text{Na}_2\text{Cr}_2\text{O}_7 / \text{H}_2\text{SO}_4$

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Note - this sheet is not meant to be comprehensive. Your course may provide additional material, or may not cover some of the reactions shown here. Your course instructor is the final authority.

Conversion of alcohols to good leaving groups

Two main methods - alkyl halides and tosylates/mesyates

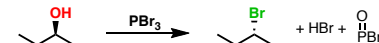
Alcohols to alkyl chlorides



Alcohols to alkyl bromides



One thing to note - these reactions occur with inversion of configuration. For example:



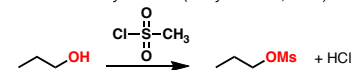
Alcohols to alkyl halides by using acids



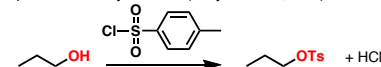
These reactions can proceed through an S_N1 or S_N2 pathway depending on the structure of the alcohol

Alcohols to tosylates and mesylates

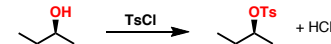
Methanesulfonyl chloride (mesyl chloride, MsCl)



p-toluenesulfonyl chloride (Tosyl chloride, TsCl)

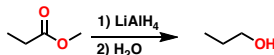


One thing to note - these reactions do not change the stereochemistry of the alcohol.

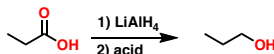


Alcohols through reduction

Reduction of esters by lithium aluminum hydride (LiAlH_4)

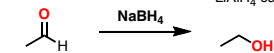


Reduction of carboxylic acids by lithium aluminum hydride (LiAlH_4)



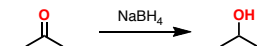
Reduction of aldehydes by sodium borohydride (NaBH_4)

LiAlH_4 can also do this reaction.



Reduction of ketones by sodium borohydride (NaBH_4)

LiAlH_4 can also do this reaction.



Omissions, Mistakes, Suggestions?

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