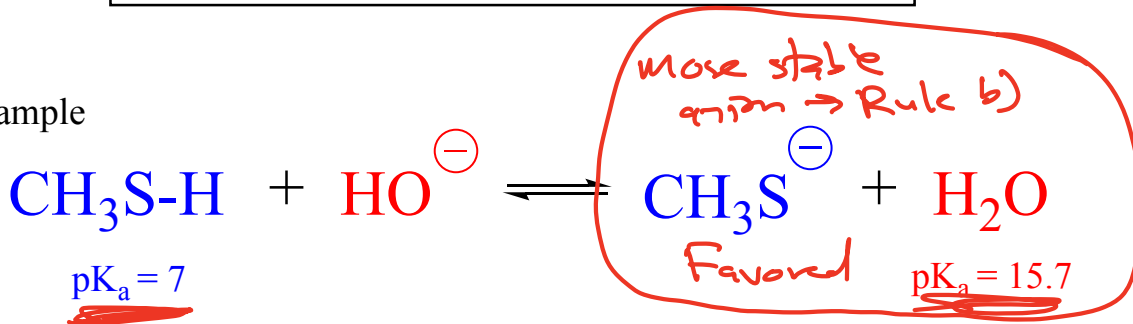


$$K_{\text{eq}} = 10^{(\text{pK}_a \text{H-B} - \text{pK}_a \text{H-A})}$$

Example



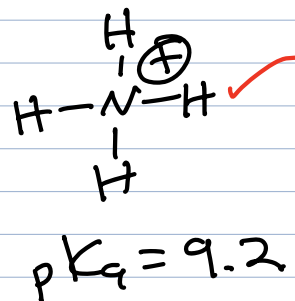
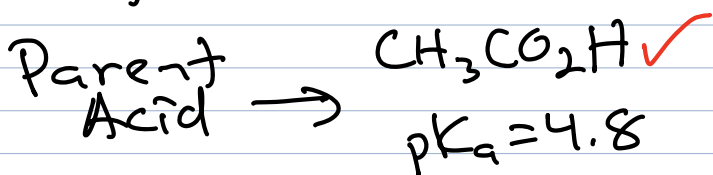
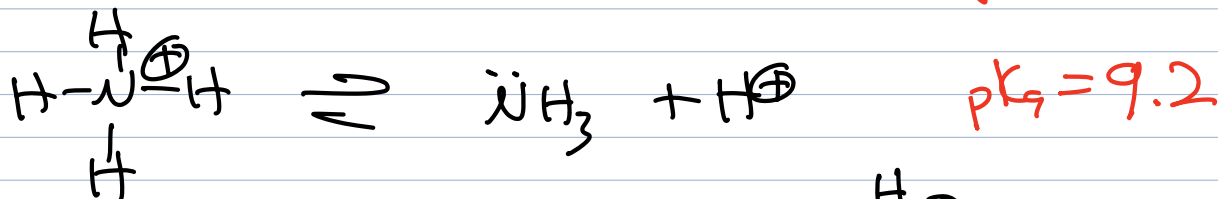
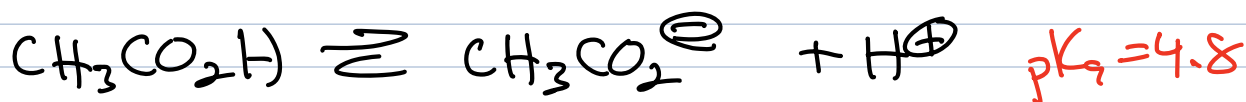
$$K_{\text{eq}} = 10^{(15.7 - 7.0)} = 10^{(8.7)}$$

New idea: How do pK_a and pH work together to determine predominant protonation states of acids?

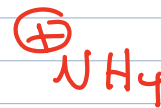
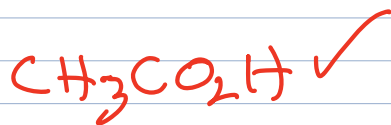
An acid is mostly **protonated** at a pH that is **below** its pK_a

An acid is mostly **deprotonated** at a pH that is **above** its pK_a

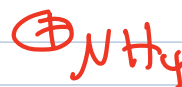
Examples:



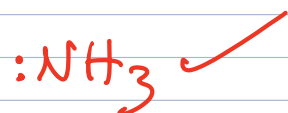
Forms Present
at $pH = 2.0$



Forms Present
at $pH = 7.0$

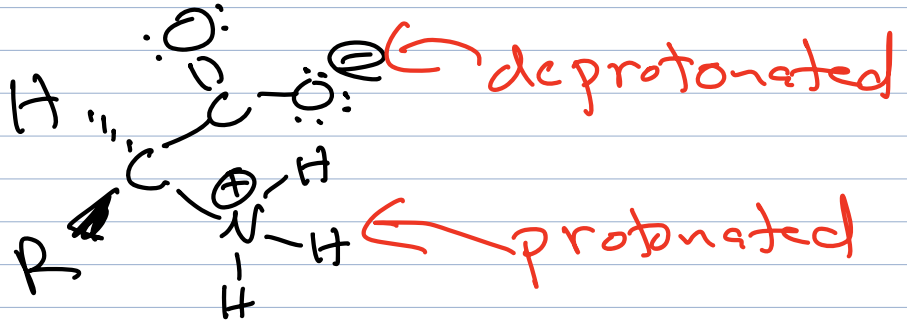


Forms Present
at $pH = 12.0$

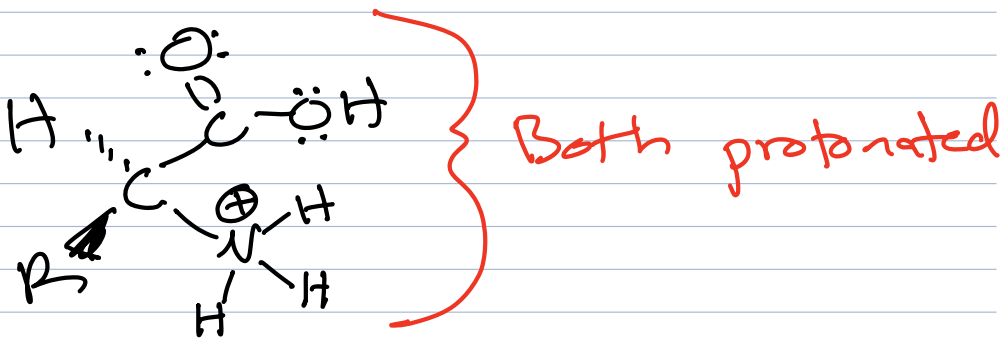


Application to important molecules \rightarrow amino acids \rightarrow same two groups we just saw

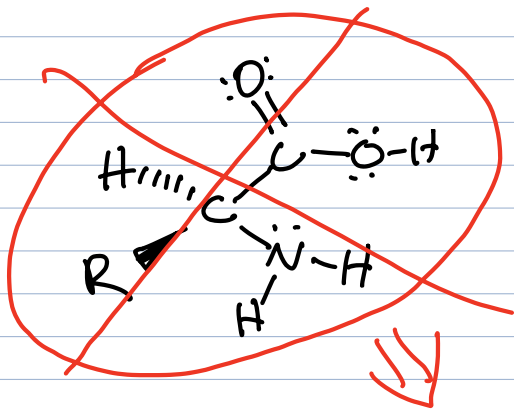
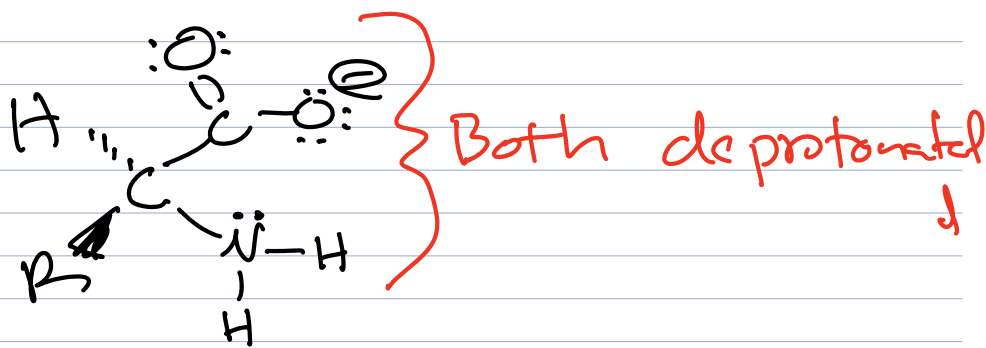
Form present at pH=7.0



Form present at pH=2.0




Form present at pH=12.0



\Rightarrow The $-\text{CO}_2\text{H}$ and $-\text{NH}_2$ forms are NEVER both present at any pH \rightarrow Not possible!!

protonated deprotonate

Why doesn't anyone get this right?



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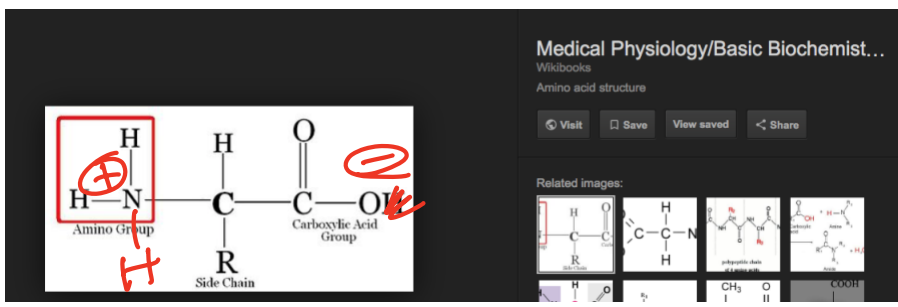
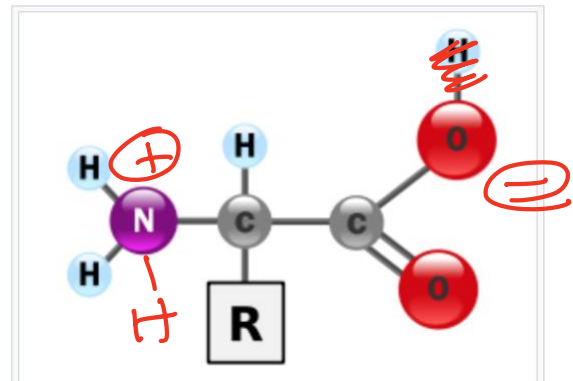
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Amino acid

From Wikipedia, the free encyclopedia

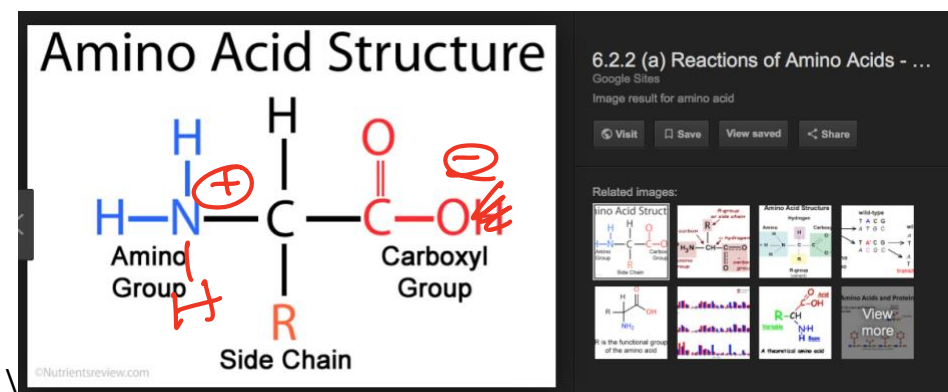
This article is about the class of chemicals. For the structures and properties of the standard proteinogenic amino acids, see [Proteinogenic amino acid](#).

Amino acids are **organic compounds** containing **amine** (-NH₂) and **carboxyl** (-COOH) functional groups, along with a **side chain** (R group) specific to each amino acid.^{[1][2][3]} The key elements of an amino acid are **carbon** (C), **hydrogen** (H), **oxygen** (O), and **nitrogen** (N), although other elements are

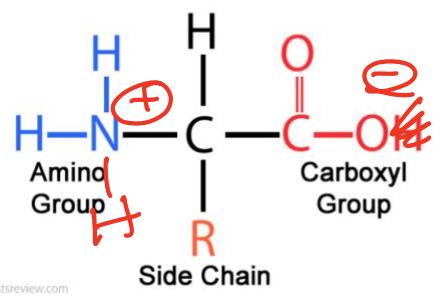


Medical Physiology/Basic Biochemist...
Wikibooks
Amino acid structure

Related images:



Amino Acid Structure

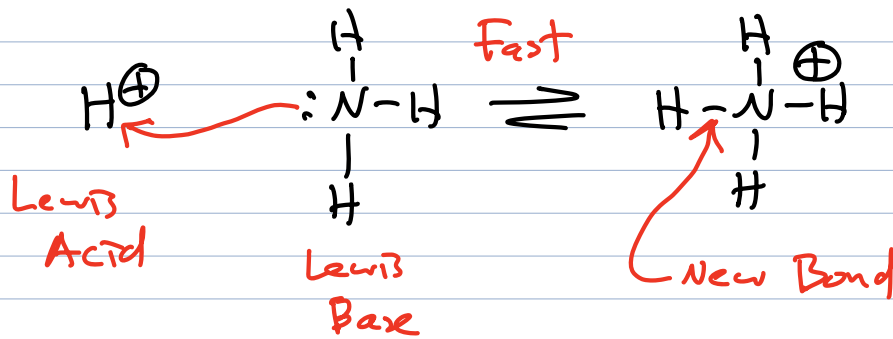


6.2.2 (a) Reactions of Amino Acids - ...
Google Sites
Image result for amino acid

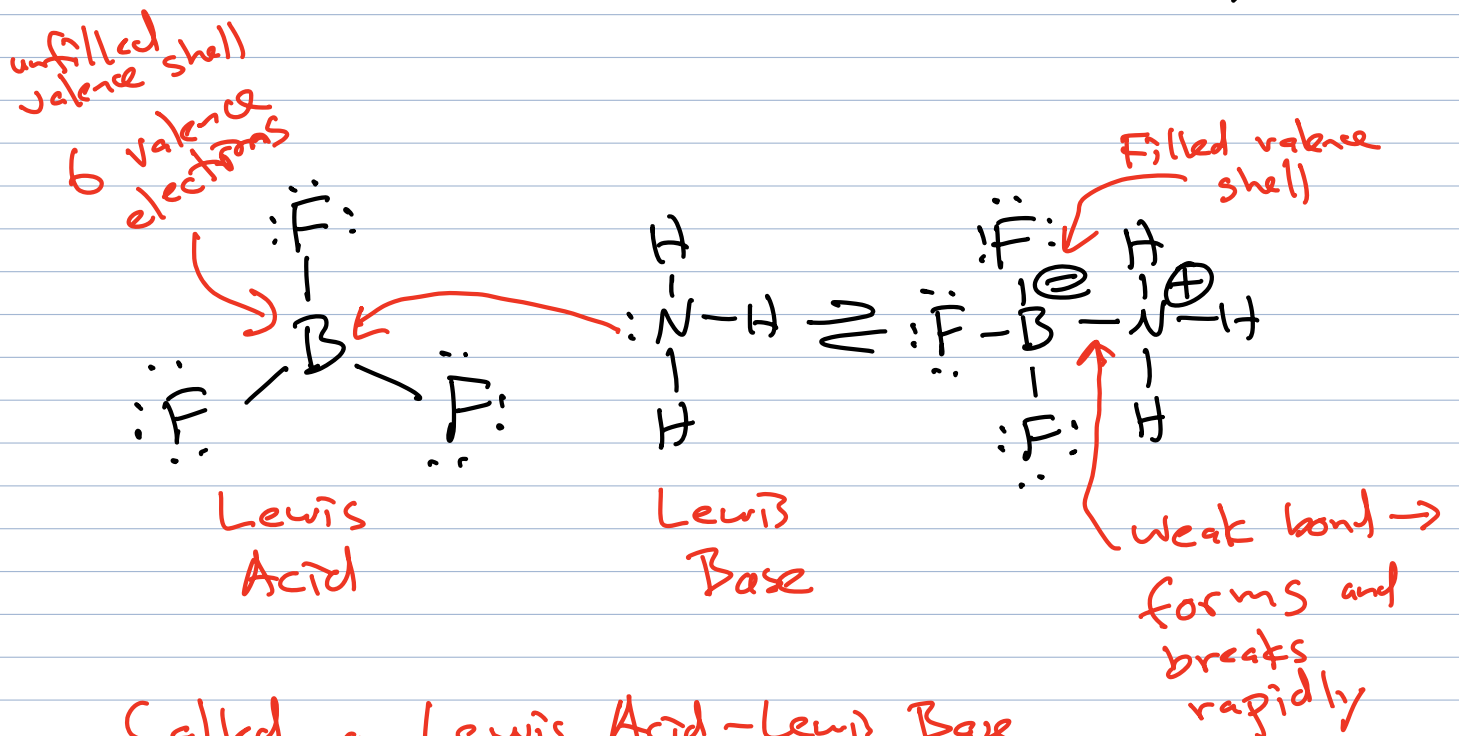
Related images:

Lewis Acid \rightarrow accepts an electron pair

Lewis Base \rightarrow donates an electron pair



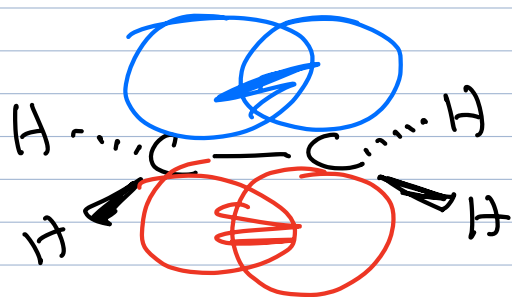
H^+ can be thought of as a Lewis acid, \rightarrow
but so can other species \rightarrow atoms with an
incomplete valence shell \Rightarrow B, Al, Zn, Fe



Called a Lewis Acid-Lewis Base
complex or "dative bond"
or "coordinate covalent bond"



Electronic Structure of Alkenes



π bonds \rightarrow
overlap of
2p orbitals

Consequences of π bonds

1) π bonds cannot rotate

2) Electron density is above and below the bond axis so it is "available" to react with electron deficient atoms/molecules

Like a
Lewis
base

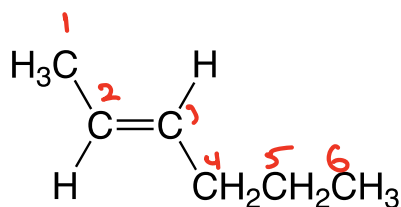
Naming Alkenes

General Directions:

1. Locate longest continuous chain. ✓
2. Number the chain so the double bond gets the lowest possible number. ✓
3. For the parent chain name, use "-ene" not "-ane" as suffix and place a number to indicate the location of the double bond before the main chain name.
4. Make the suffix "-adiene", "-atriene", etc. if multiple double bonds are present.

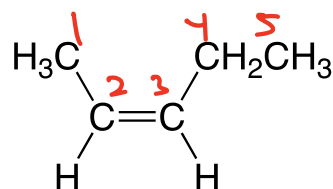
cis/trans nomenclature – older chemical nomenclature, but still used commonly in biochemistry – most useful when each sp^2 atom of the double bond has an H atom.

1. Track the longest chain through the double bond ✓
 - a. ***cis*** if whole main chain is on the same side of the double bond. ✓
 - b. ***trans*** if chain emerges on opposite sides of the double bond. ✓



trans-2-hexene

same side



cis-2-pentene

NOT IUPAC

Naming Alkenes

General Directions:

1. Locate longest continuous chain. ✓
2. Number the chain so the double bond gets the lowest possible number. ✓
3. For the parent chain name, use "-ene" not "-ane" as suffix and place a number to indicate the location of the double bond before the main chain name.
4. Make the suffix "-adiene", "-atriene", etc. if multiple double bonds are present.

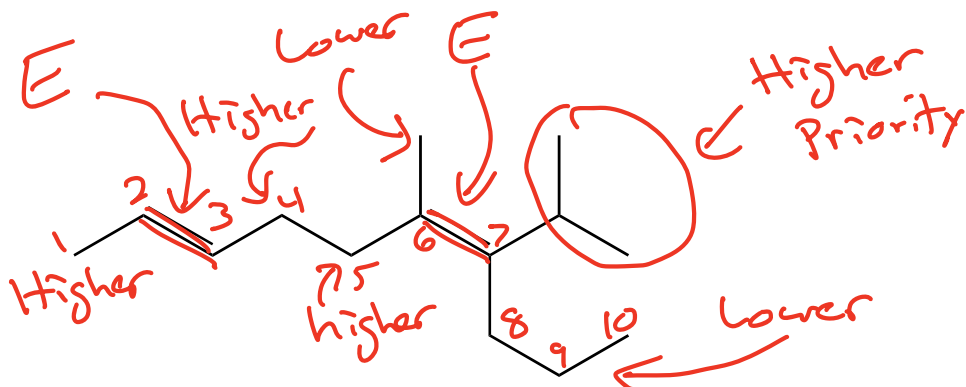
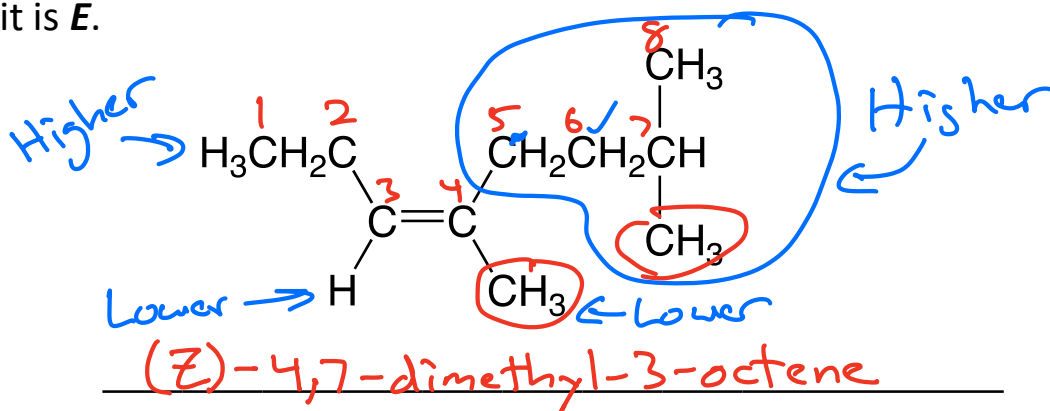
E,Z nomenclature - A general IUPAC nomenclature to names alkenes.



Z (zusammen) = same side, same side, same side

E (entgegen) = opposite side

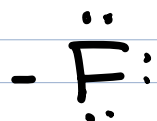
5. On each carbon of the double bond rank the two groups according to the Cahn, Ingold, Prelog priority rules (*R* vs. *S* rules).
6. If both of the highest-ranking groups are on the same side of the double bond it is **Z**.
7. If both of the highest-ranking groups are on opposite sides of the double bond it is **E**.



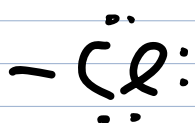
(2E,6E)-7-isopropyl-6-methyl-2,6-decadiene

or (2E,6E)-6-methyl-7-(1-methylethyl)-2,6-decadiene

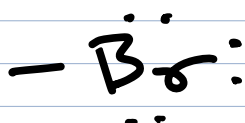
Today's Special Bonus Feature: Halogen Nomenclature



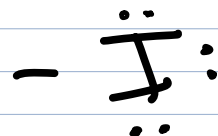
fluoro-



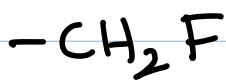
chloro-



bromo-



iodo-



fluoromethyl
group



(R)-2-bromohexane