

Reaction Coordinate

New idea: How do pha and pH work together to determine predominant protonation states of acids? An acid is nostly protonated at q git that is below its pkg An acid is mostly deprotonablet q pt that is above its gkg Examples: $CH_3CO_2H \ge CH_3CO_2 + H^{\oplus} pK_q = 4.8$ $H - \lambda = H = \lambda H_3 + H = p K_4 = 9.2$ Parent CH3CO2HV H-N-HV Acid > pKa=4.8 H $pK_q = 9.2$ Forms Present at pH = 2.0 A Hy CH3CO2H Forms Present at pH=7.0 CH3CO2 @NHy Forms Present at ptt=12.0 CH3CO2 :NH3

Application to important molecules -> <u>amino</u> <u>acids</u> -> same two groups we just saw Form present H., C. .: Cort at pH=7.0 Cort H yrobnated Form present H., C-ÖH at pH=2.0 PH H forms are NEVER both present at any pH -> Not possible!!

why doesn't anyone get this right?



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Amino acid

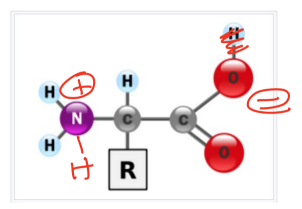
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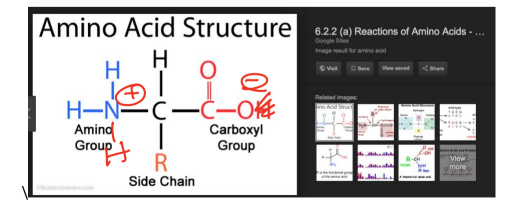
This article is about the class of chemicals. For the structures and properties of the standard proteinogenic amino acids, see Proteinogenic amino acid.

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



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Lewis Acid - accepts an electron pair Levis Base -> donates an electron pair ·N-H = H-N-H H Fast Levis Acro New Bond Lewis Bare H[@] can be thought of as a Lewis acid, but so can other species -> atoms with an unfilled shell 6 valence Filled valence shell ÷Fì N-H Z:F-B-N-H weak bond -> Lewis Lewis Base Acid forms and breaks rapidly Called a Lewis Acid - Lewis Base unglex or "detire band" or " coordinate covalent band"



Electronic Structure of Alkenes

Hand W bond, H overlap of 2p orbitals

Consequences of I bonds 1) It bonds cannot ratate

2) Electron density is above and Like 9 below the bond aris so it is "aucidable" to react with electron 1 eni deficient atoms/molecules

Naming Alkenes

General Directions:

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

<u>cis/trans nomenclature</u> – 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. \checkmark
 - b. *trans* if chain emerges on opposite sides of the double bond. \checkmark

Naming Alkenes

General Directions:

- 1. Locate longest continuous chain.
- 2. Number the chain so the double bond gets the lowest possible number. \checkmark
- 3. For the parent chain name, use "-<u>ene</u>" not "-<u>ane</u>" 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.

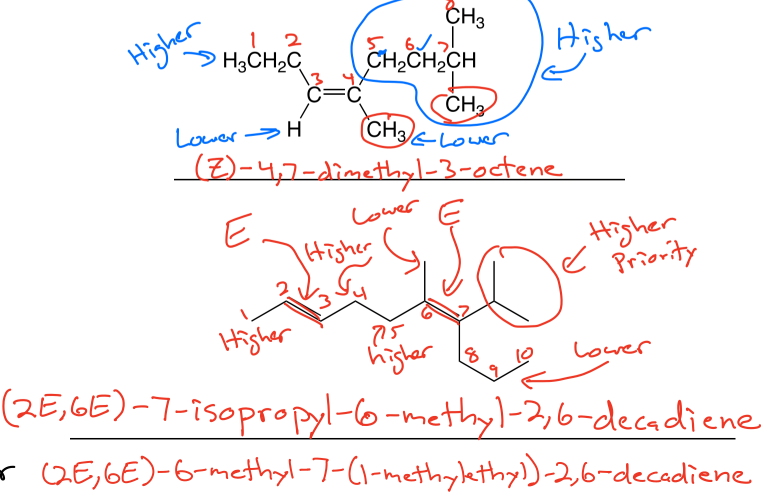
E,Z **nomenclature** - A general IUPAC nomenclature to names alkenes.



Z (zusammen) = zame zide, zame zide, zame zide

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 zame zide of the double bond it is **Z**.
- If both of the highest-ranking groups are on opposite sides of the double bond it is *E*.



Today's Special Bonnes Feature: Halogen Nomenclature -(l: -Br: -- <u>F</u>: fluoro-chloro-bromo-1000- $-CH_2F$ 6/543 fl-oromethy) Br gro-P (R)-2-brono hexane