Welcome... back!

Office hours:

M: 1:30-2:30

Tu-F: after lecture

1168

Week 4

Reading: Section 16.1



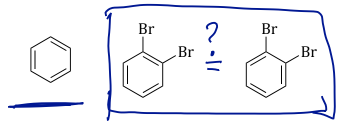
Draw the structures of the following molecules:

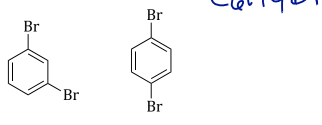
methylbenzene toluene *m*-chlorophenol nitrobenzene *p*-nitroaniline aniline= o-chloroanisole 1,2-

١L

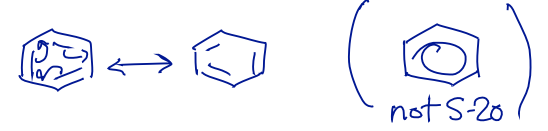
Aromaticity: More than just conjugation

In the 19th century, benzene was quite a puzzle. In 1865, Kekulé proposed the nowfamiliar structure for benzene that we might call "1,3,5-cyclohexatriene." This structure was quite controversial because it suggests that there should be four isomeric CoHyBra dibromobenzenes, while in fact there are only three.





Of course, we know that there is only one ortho-dibromobenzene. How can that be?



Benzene is an example of an **aromatic** compound. The criteria for aromaticity are:

morethan just resonance 1) The molecule (or ion) must contain a contiguous, planar, cyclic array of p-orbitals

2) The array of p-orbitals must contain 4n+2 electrons (where n=0,1,2...) $2, \frac{1}{2}, \frac{1}{2}, \frac{1}{2}$

Aromatic compounds are especially stable.

4,8,12,16... If a molecule (or ion) satisfies the first criterion, but contains 4n electrons, it is **antiaromatic**, and especially *unstable*.

Are the following molecules aromatic, antiaromatic, or neither?



7 p.orb. 4 p.orb. 3 p orb 8 Tre-antiarom.

Reading: Section 15.7

nonaromatic

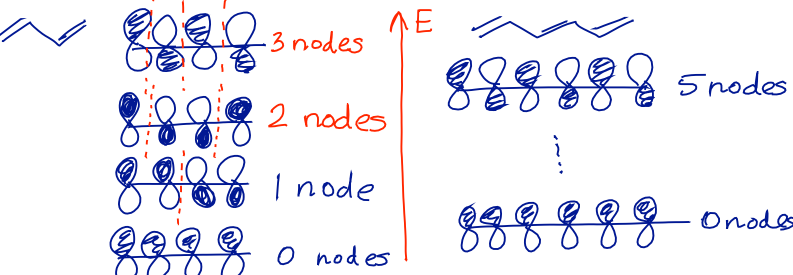
antiaromatic

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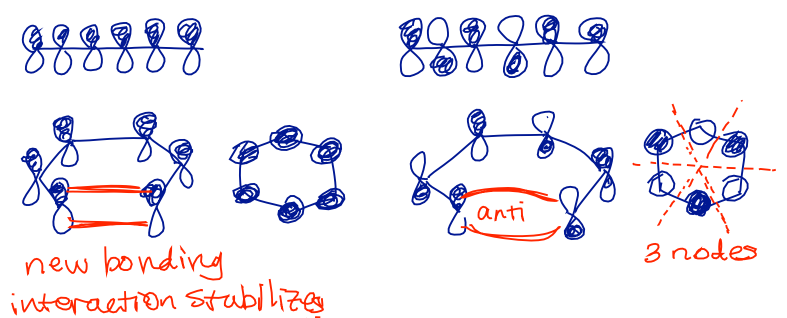
Molecular Orbitals and Aromaticity 1: The Hückel Rule & Frost's Circle

Why **4n + 2**? Is there anything special about the numbers 2, 6, 10, 14, etc? To answer that question, we need to look at the **molecular orbitals**!

For *linear* conjugated systems, we have n π -molecular orbitals, with up to n-1 nodes:



For *cyclic* conjugated systems, we still have n π -molecular orbitals, but where do the nodes go, and how many are there? The highest and lowest energy orbitals look like what we might expect – the "all-bonding" and "all-antibonding" combinations:



Reading: Section 15.7

Tala

3

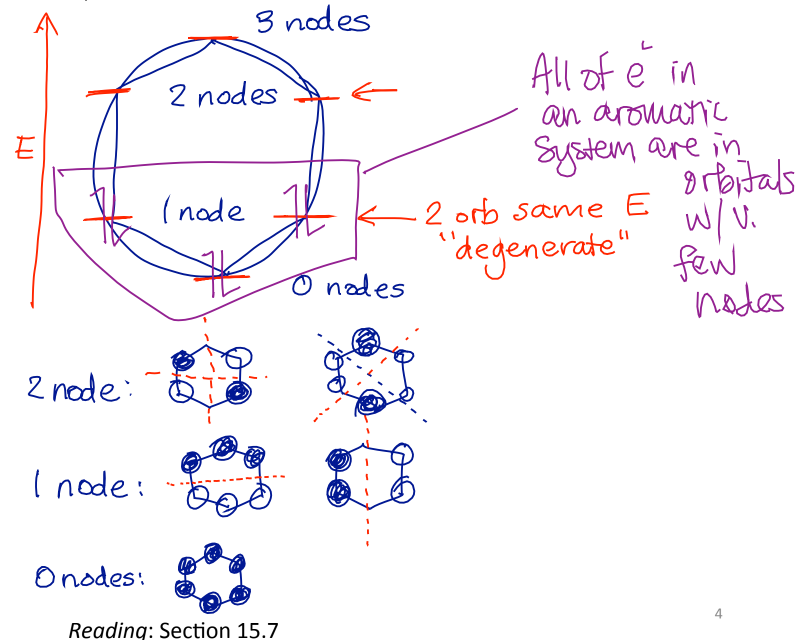
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Molecular Orbitals and Aromaticity 2: The Hückel Rule & Frost's Circle

To figure out what the other MO's look like:

- 1) Pick your polygon and draw it with one vertex at the bottom.
- 2) Place an MO at each vertex. Note that we have *pairs* of MO's at equal energies!
- 3) As we go from bottom to top, we increase the number of nodes from 0 to n/2.
- 4) For degenerate pairs of orbitals, nodes are orthogonal.
- 5) Fill with electrons.

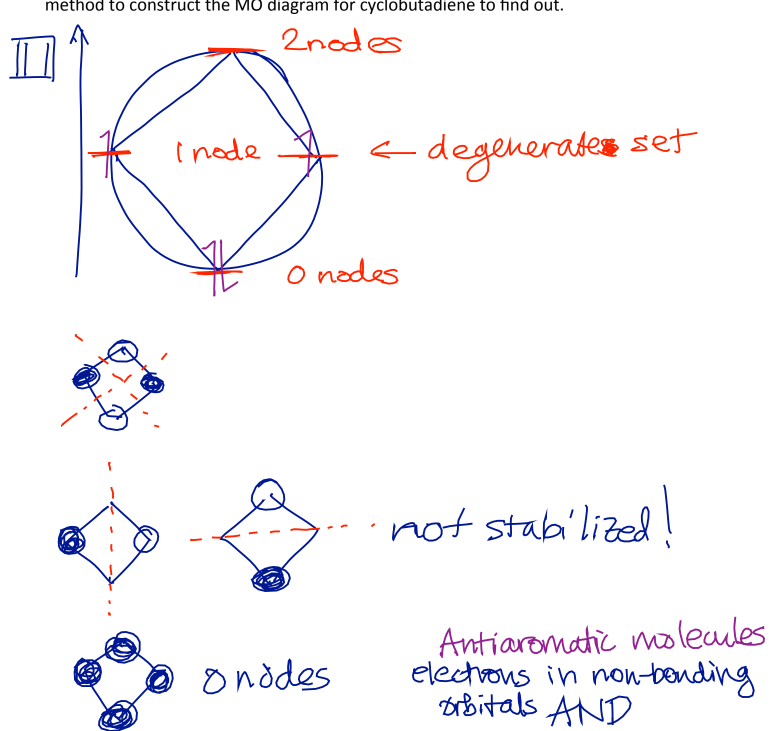


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Molecular Orbitals and Aromaticity 3:

The Hückel Rule & Frost's Circle

OK, but why is antiaromatic bad? What's wrong with 4n electrons? Use the Frost's Circle method to construct the MO diagram for cyclobutadiene to find out.



Reading: Section 15.7

arediradi cals

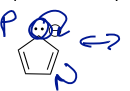
The Effect of Aromaticity on Reactivity

The rule is simple: aromaticity = (very!) good, antiaromaticity = bad Explain the following observations:

1) When styrene is treated with Br₂, the bromine adds only to the terminal alkene and not to the benzene ring.

$$Br_2$$
 Br_2
 Br_3

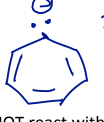
2) Only one of the following hydrocarbons can be deprotonated by NaNH₂.



6 portos

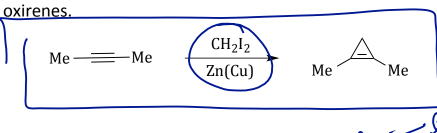






geantiaronal

3) Alkynes react with carbenes to form cyclopropenes, but do NOT react with mCPBA to form oxirenes.



Me — _____ Me _____ mC

Me Me

4 Tre-

Reading: Section 15.7

antraromatil

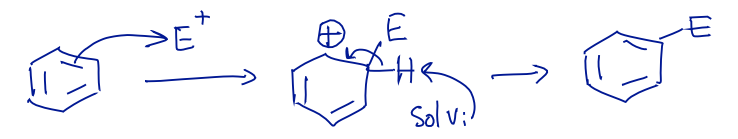
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Electrophilic Aromatic Substitution:

General Mechanism

When benzene is treated with D_2SO_4 (D = deuterium, ²H), the protons of benzene are slowly replaced with deuterium. Provide a curved-arrow mechanism.

This is an example of *electrophilic aromatic substitution*. What is the general reaction for a generic electrophile, E⁺?

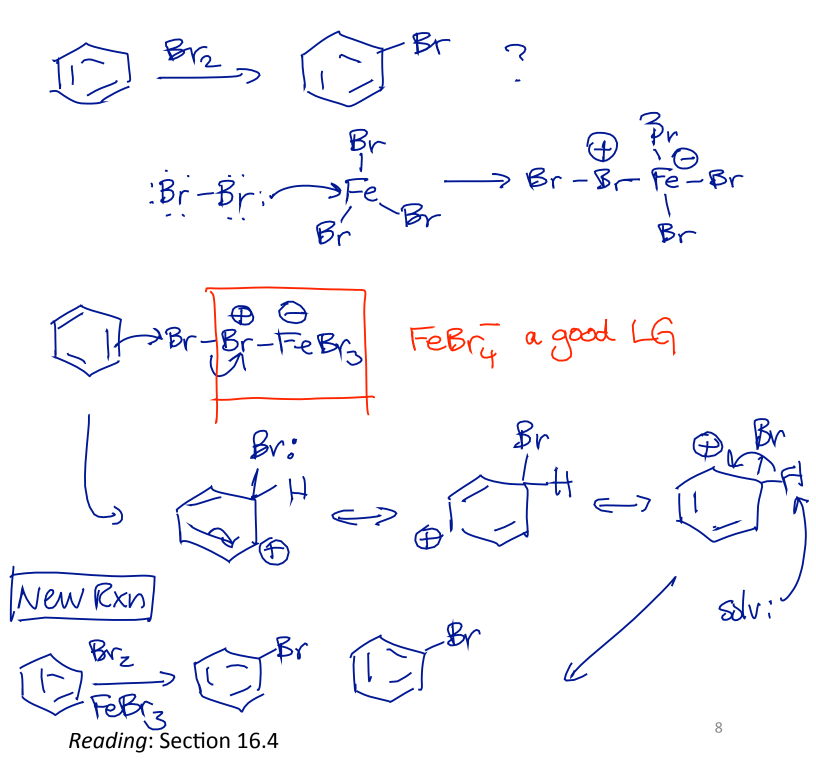


Reading: Section 16.4

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Electrophilic Aromatic Substitution: Halogenation

How can we replace a hydrogen (H+) with chlorine (Cl+) or bromine (Br+)?



Chlorination: Cl2 Cl2 AICl3 $CI-CI: \longrightarrow AI \longrightarrow CI-CI-AI-CI$ Styci-Gi-Airia excellent LG

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Electrophilic Aromatic Substitution:

Nitration and Sulfonation

How can we replace a hydrogen (H+) with a nitro group (NO_2^+) ? $E^+ = NO_2^+ = O = N = O$ HNO_3 O = N = O $HNO_2^+ = NO_2^+ = NO_$

How can we replace a hydrogen (H⁺) with a sulfonic acid group (SO₃H⁺)?