

Welcome ... back!

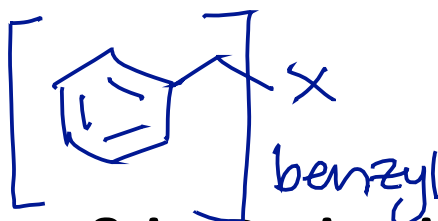
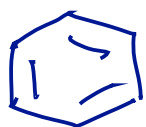
Office hours:

M: 1:30 - 2:30

Tu-F: after lecture

116B

Week 4

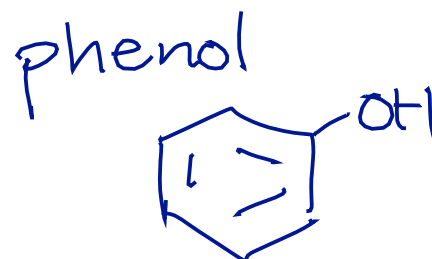


July 14, 2014

## Benzene & its Derivatives

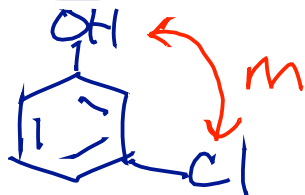
Draw the structures of the following molecules:

toluene methylbenzene

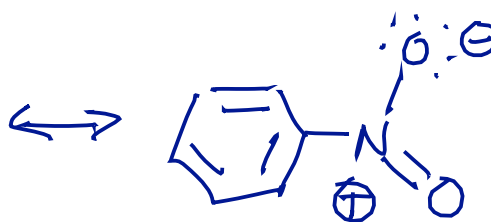
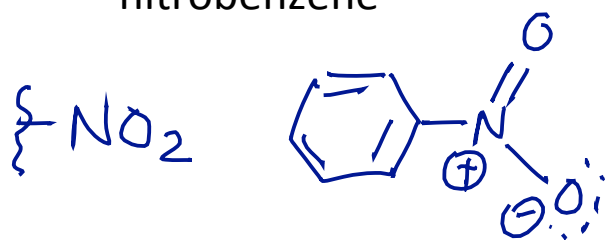


"meta" m-chlorophenol

1,3-

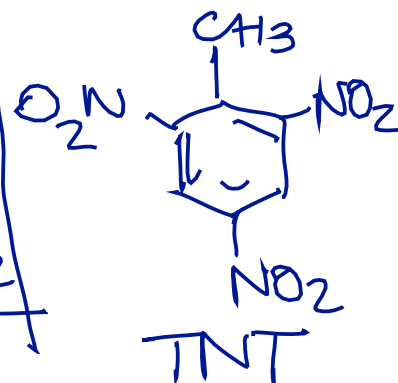
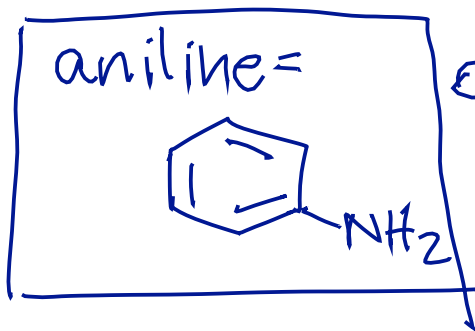
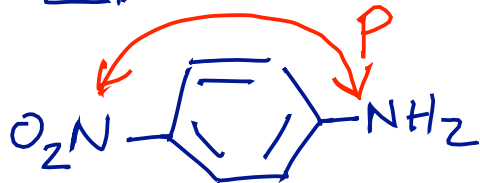


nitrobenzene



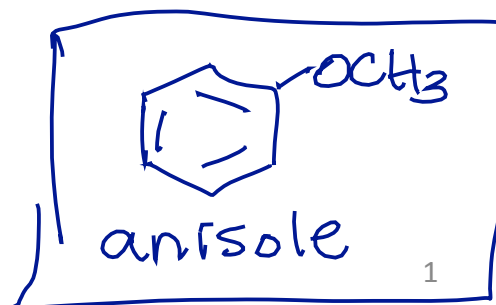
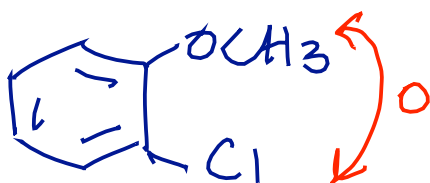
"para" p-nitroaniline

1,4-



"ortho" o-chloroanisole

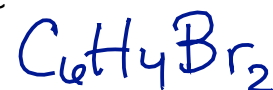
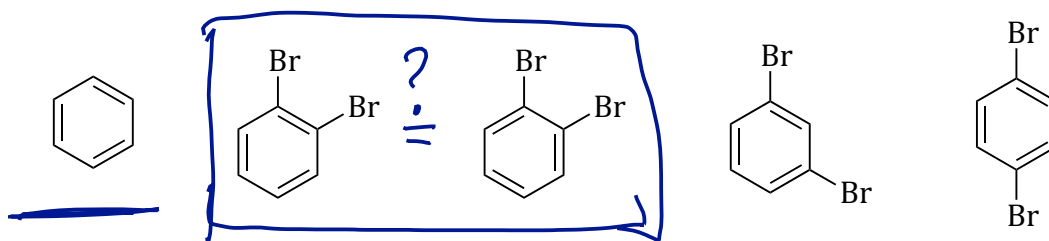
1,2-



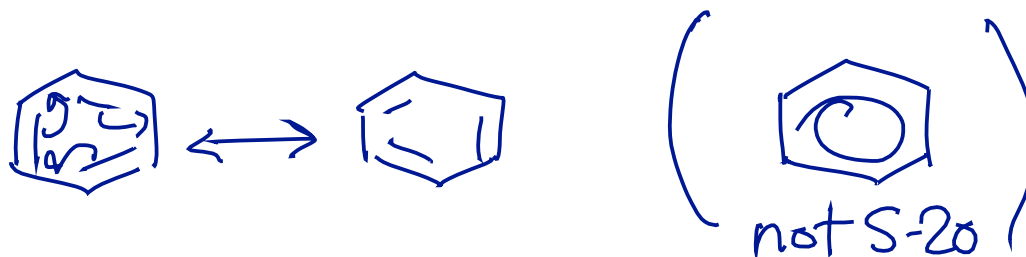
Reading: Section 16.1

## Aromaticity: More than just conjugation

In the 19<sup>th</sup> century, benzene was quite a puzzle. In 1865, Kekulé proposed the now-familiar 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 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:

*more than 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  $n = 0, 1, 2, \dots$ )

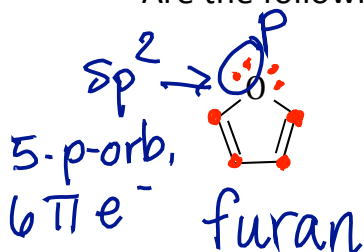
2, 6, 10, 14

Aromatic compounds are **especially stable**.

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

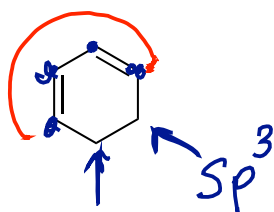
4, 8, 12, 16...

Are the following molecules *aromatic*, *antiaromatic*, or neither?

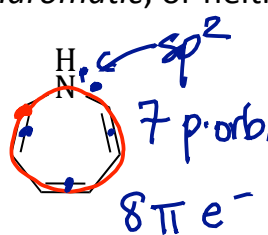


Reading: Section 15.7

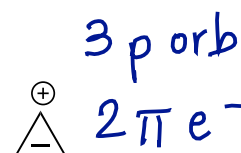
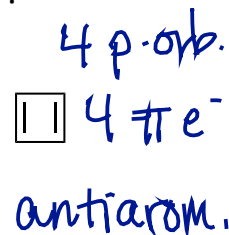
aromatic



nonaromatic



antiaromatic



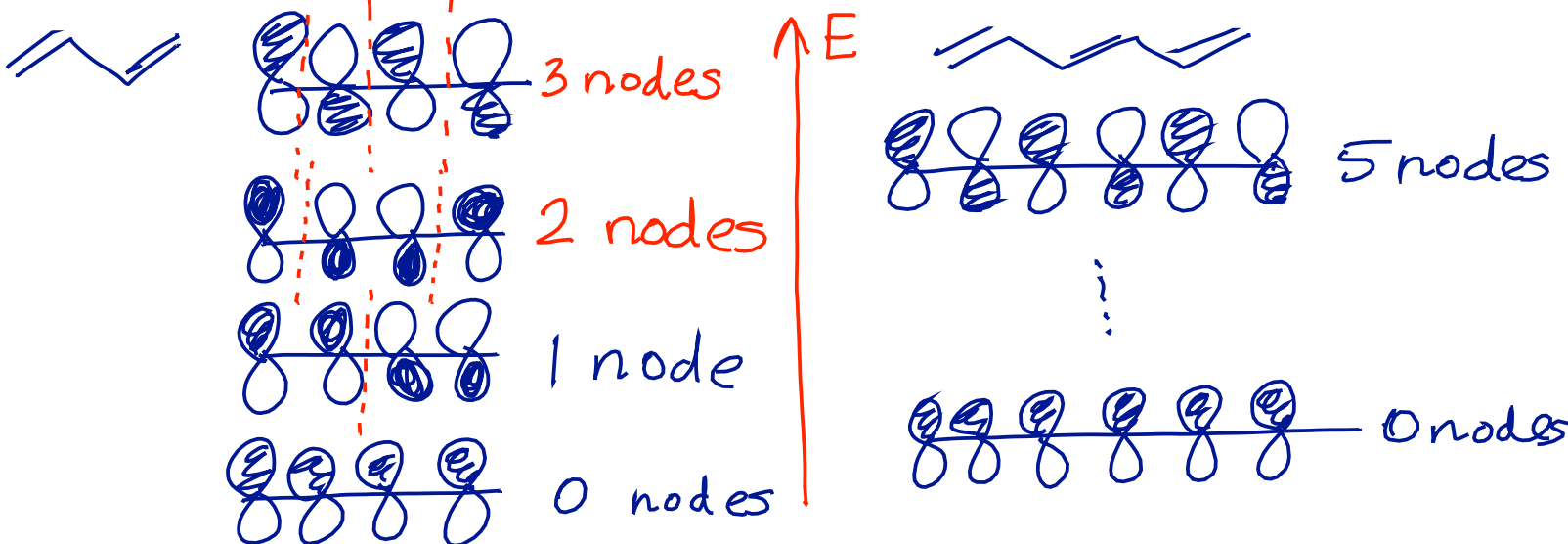
aromatic!

# Molecular Orbitals and Aromaticity 1:

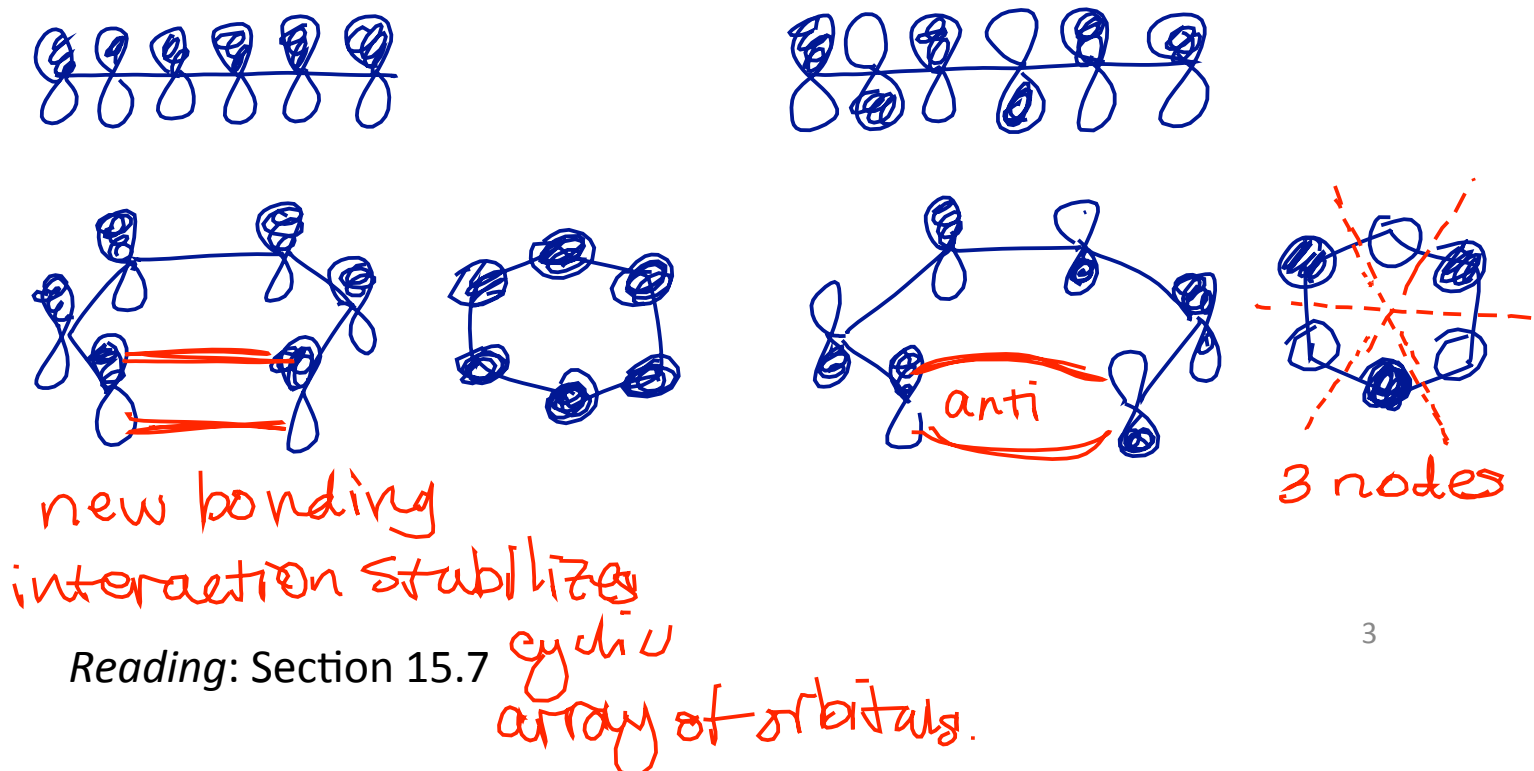
## The Hückel Rule & Frost's Circle

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  $n$   $\pi$ -molecular orbitals, with up to  $n-1$  nodes:



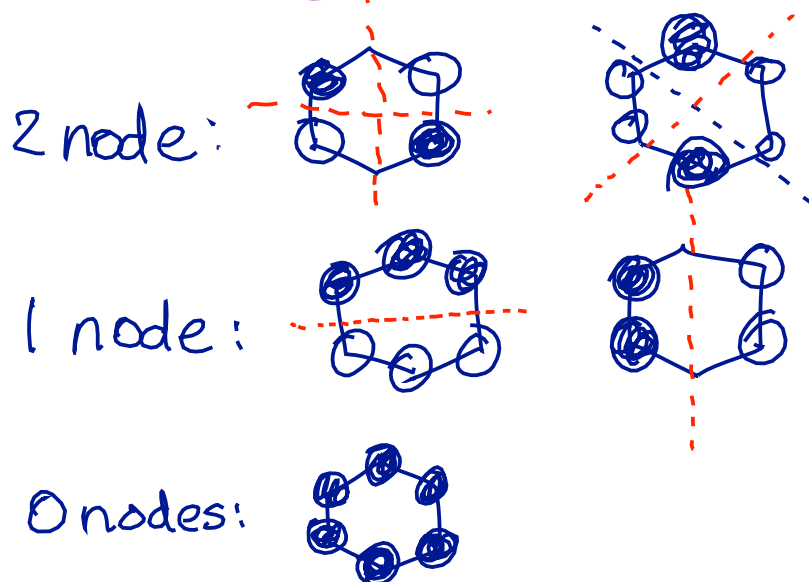
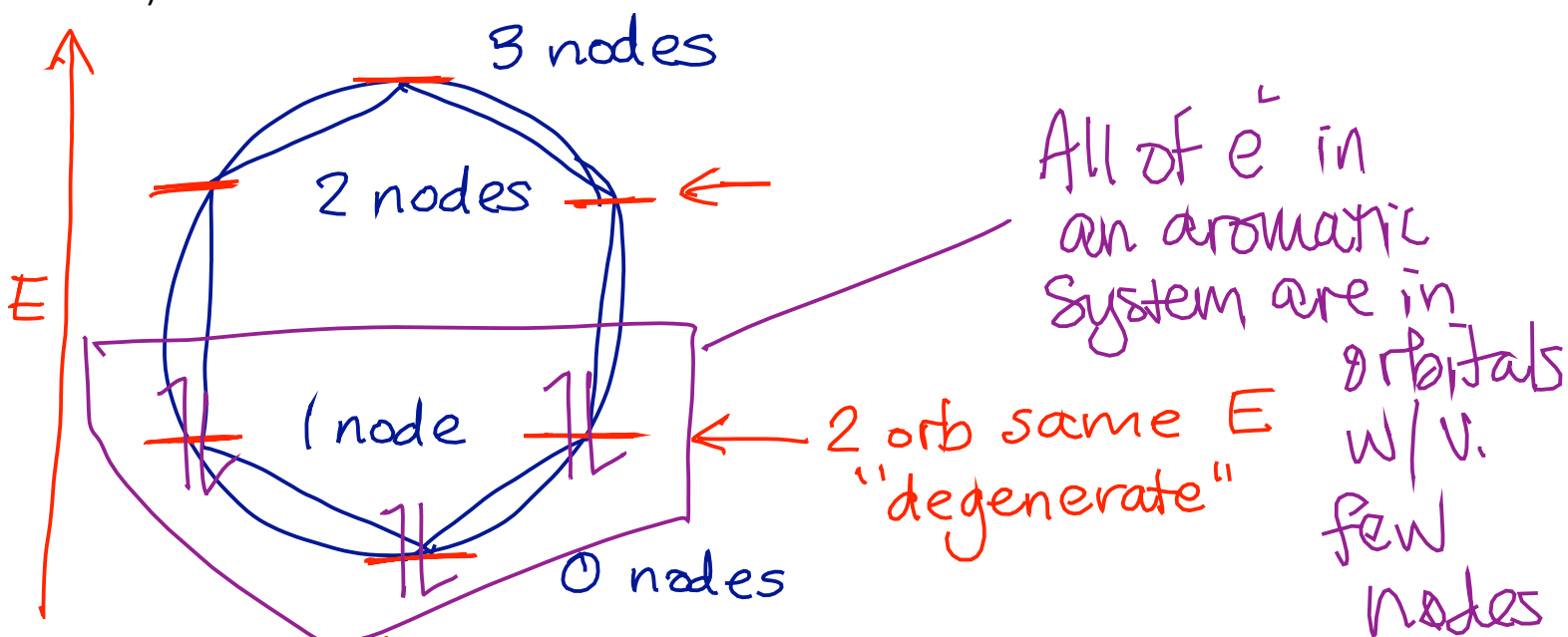
For *cyclic* conjugated systems, we still have  $n$   $\pi$ -molecular orbitals, but where do the nodes go, and how many are there? The highest and lowest energy orbitals look like what we might expect – the “all-bonding” and “all-antibonding” combinations:



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

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

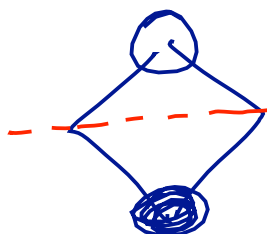
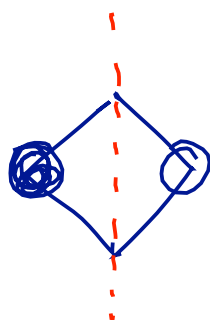
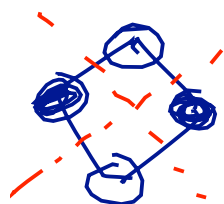
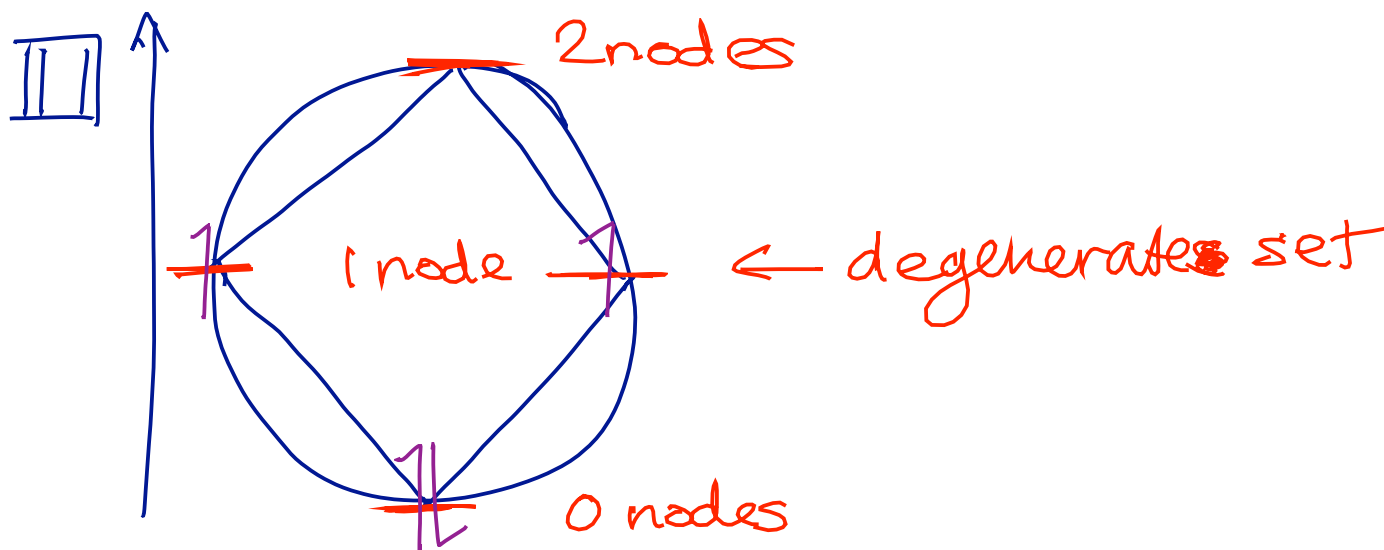
- 1) Pick your polygon and draw it with one vertex at the bottom.
- 2) Place an MO at each vertex. Note that we have *pairs* of MO's at equal energies!
- 3) As we go from bottom to top, we increase the number of nodes from 0 to  $n/2$ .
- 4) For degenerate pairs of orbitals, nodes are orthogonal.
- 5)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



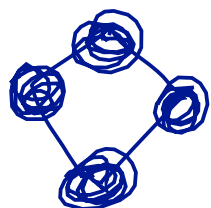
Reading: Section 15.7

## 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.



not stabilized!



0 nodes

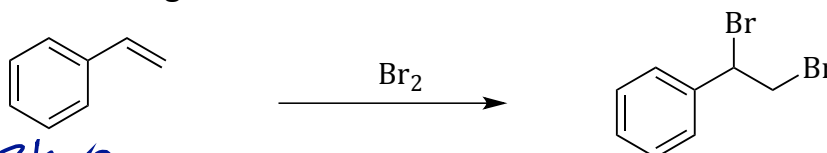
Antiaromatic molecules  
electrons in non-bonding  
orbitals AND  
are diradicals

# The Effect of Aromaticity on Reactivity

The rule is simple: aromaticity = (very!) good, antiaromaticity = bad

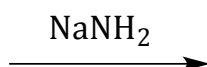
Explain the following observations:

- 1) When styrene is treated with  $\text{Br}_2$ , the bromine adds only to the terminal alkene and not to the benzene ring.

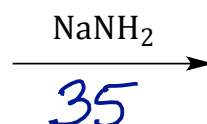


styrene

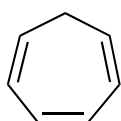
- 2) Only one of the following hydrocarbons can be deprotonated by  $\text{NaNH}_2$ .



5 p-orb's  
6  $\pi$  e's  
aromatic



no reaction

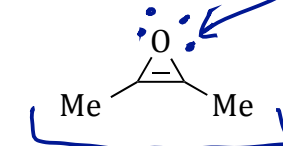
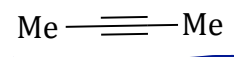
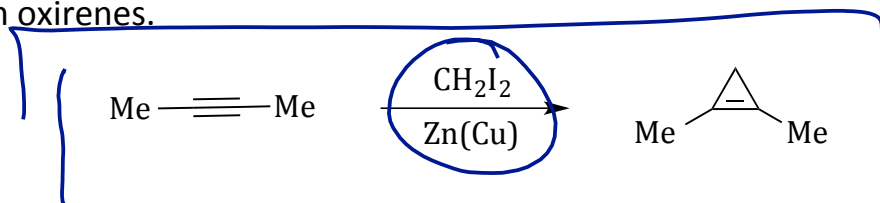


no reaction



7 p-orb's  
8  $e^-$   
antiaromatic

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



unstable

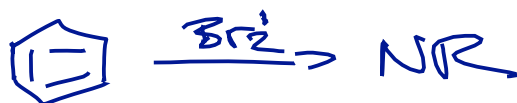
antiaromatic

lp in p-orbital  
4  $\pi e^-$



alkenes are nucleophiles

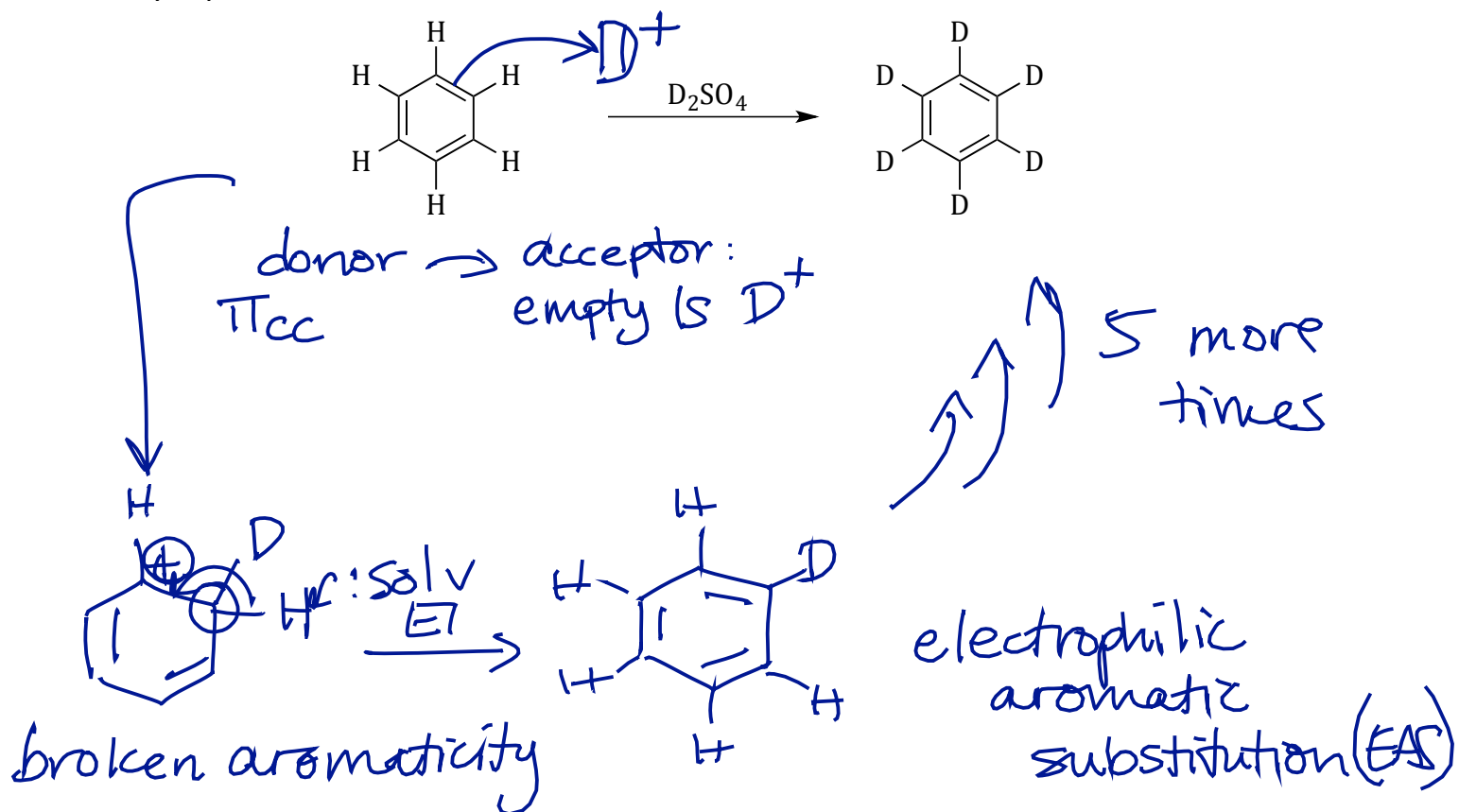
Week 4



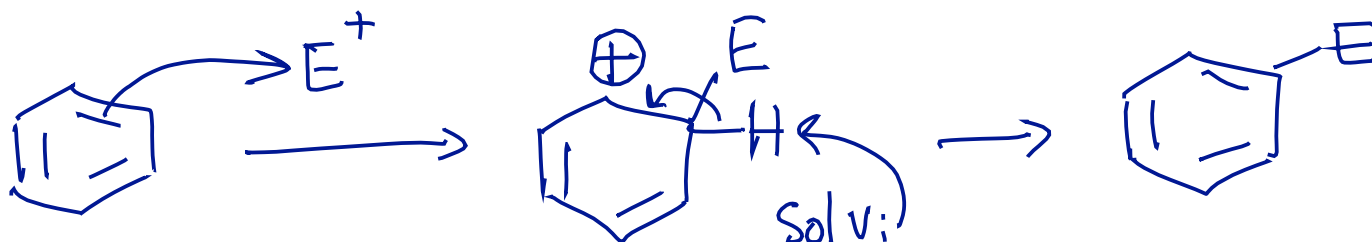
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## Electrophilic Aromatic Substitution: General Mechanism

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.



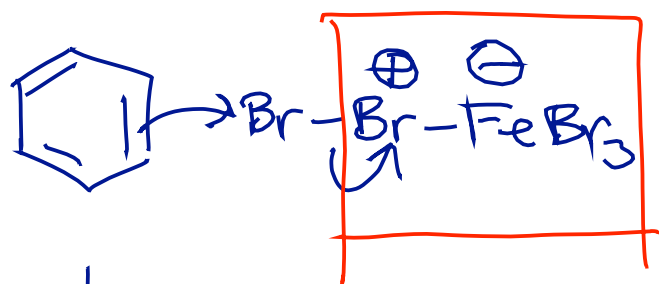
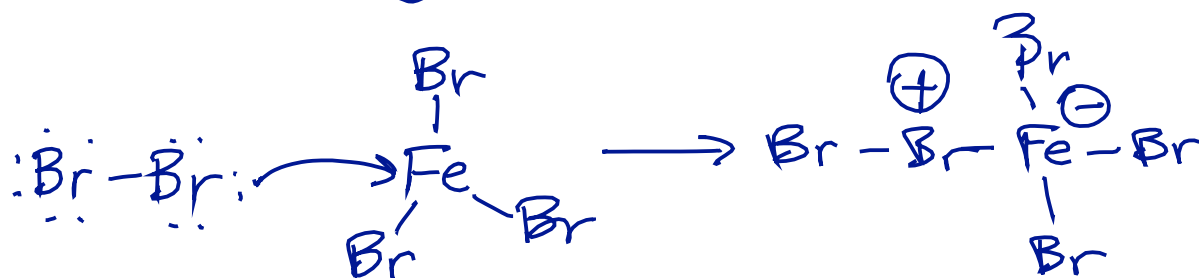
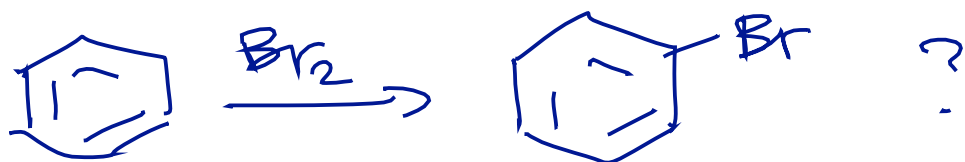
This is an example of *electrophilic aromatic substitution*. What is the general reaction for a generic electrophile,  $E^+$ ?



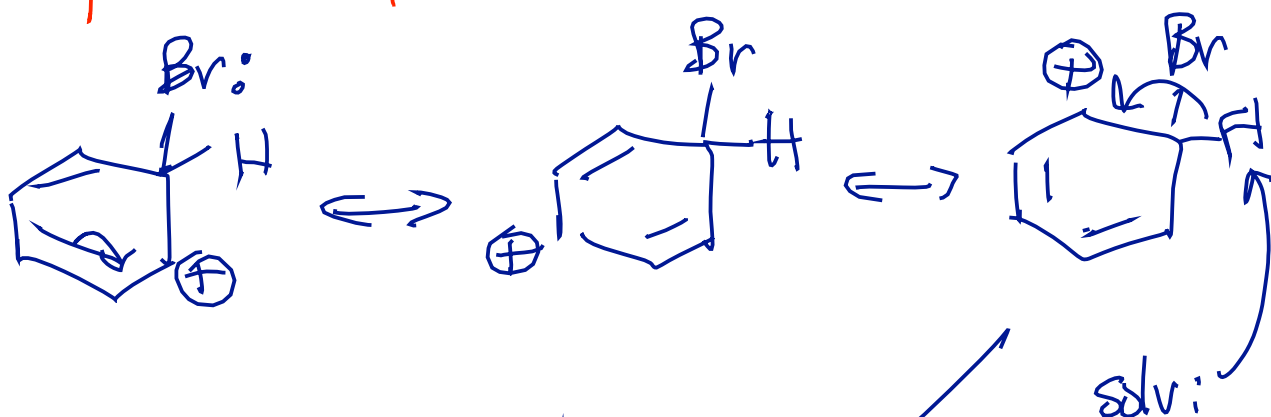


# Electrophilic Aromatic Substitution: Halogenation

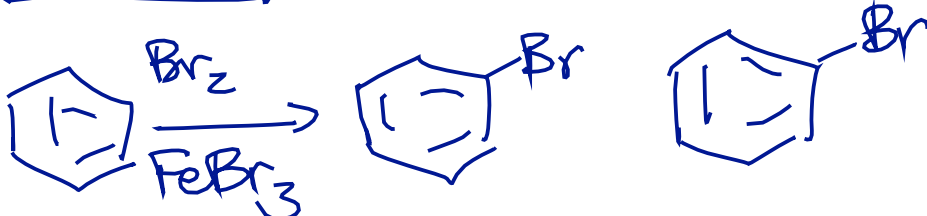
How can we replace a hydrogen ( $H^+$ ) with chlorine ( $Cl^+$ ) or bromine ( $Br^+$ )?



$FeBr_4^-$  a good LG

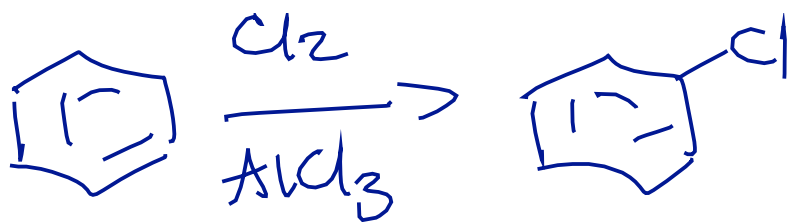
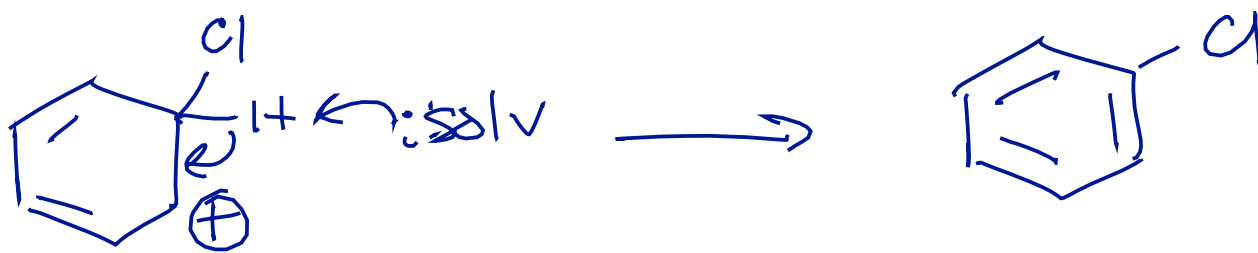
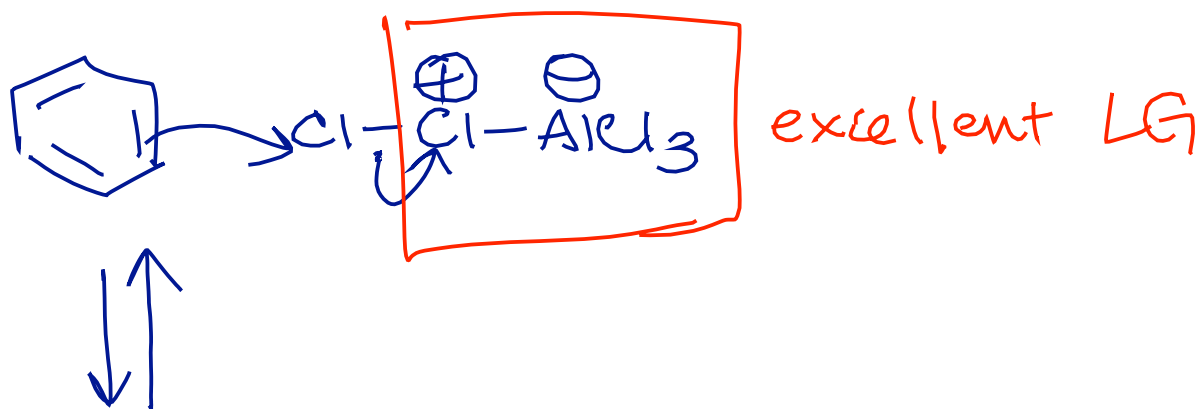
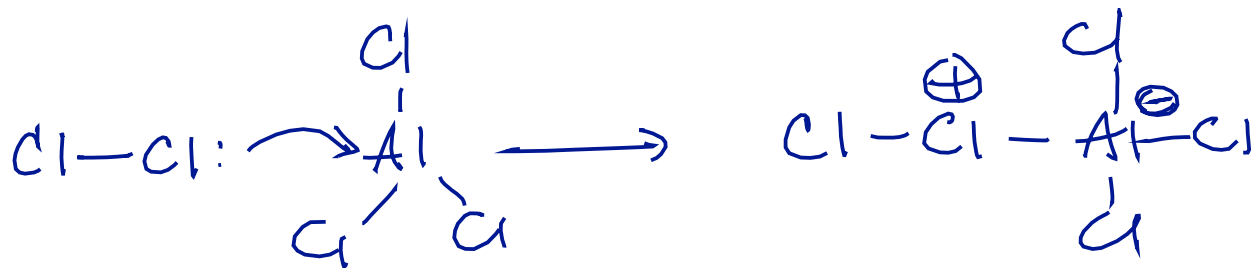
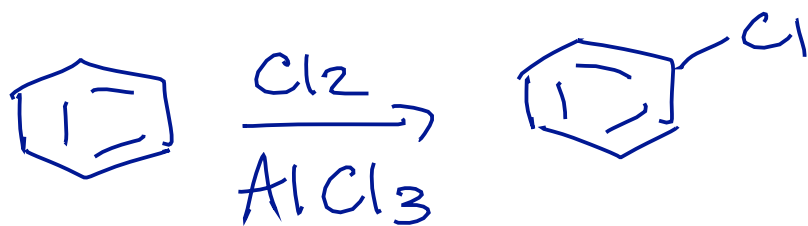


New Rxn



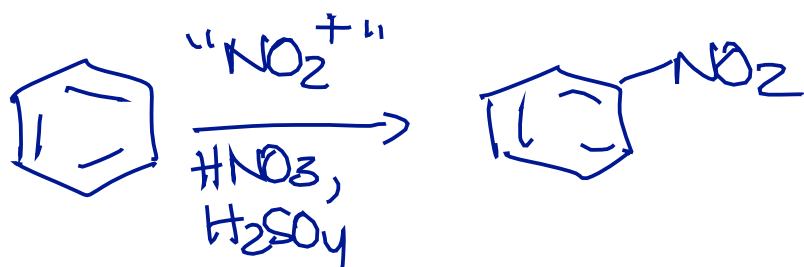
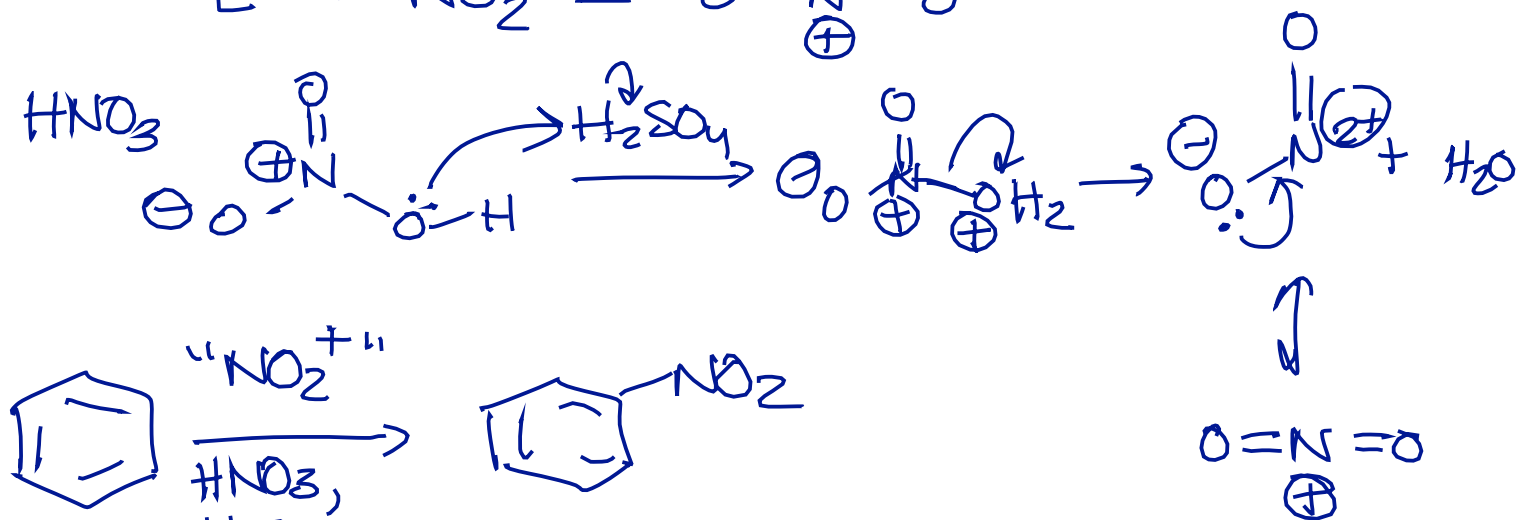
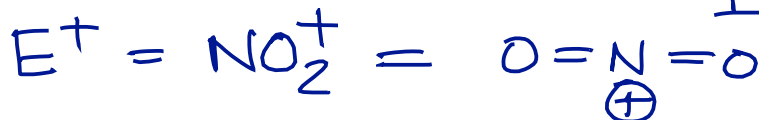
Reading: Section 16.4

Chlorination:



## Electrophilic Aromatic Substitution: Nitration and Sulfonation

How can we replace a hydrogen ( $H^+$ ) with a nitro group ( $NO_2^+$ )?



How can we replace a hydrogen ( $H^+$ ) with a sulfonic acid group ( $SO_3H^+$ )?