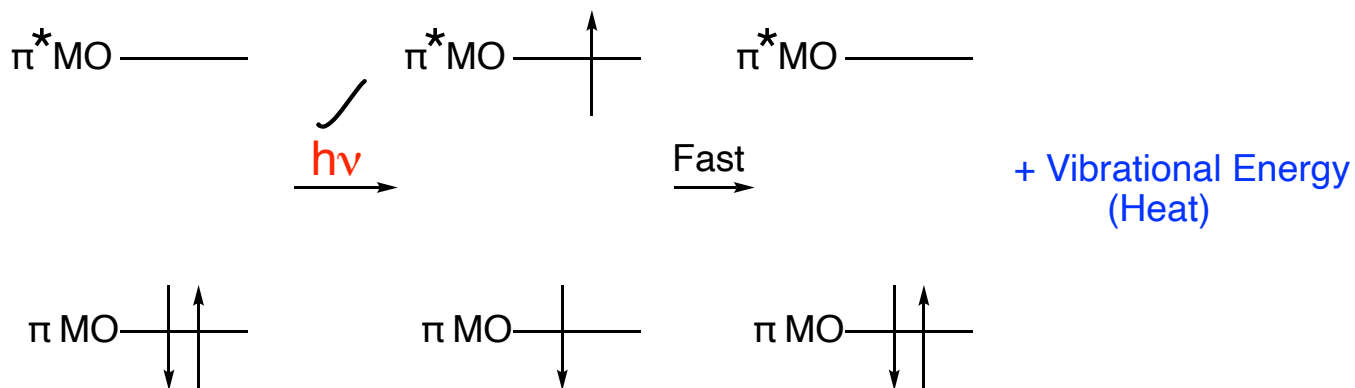
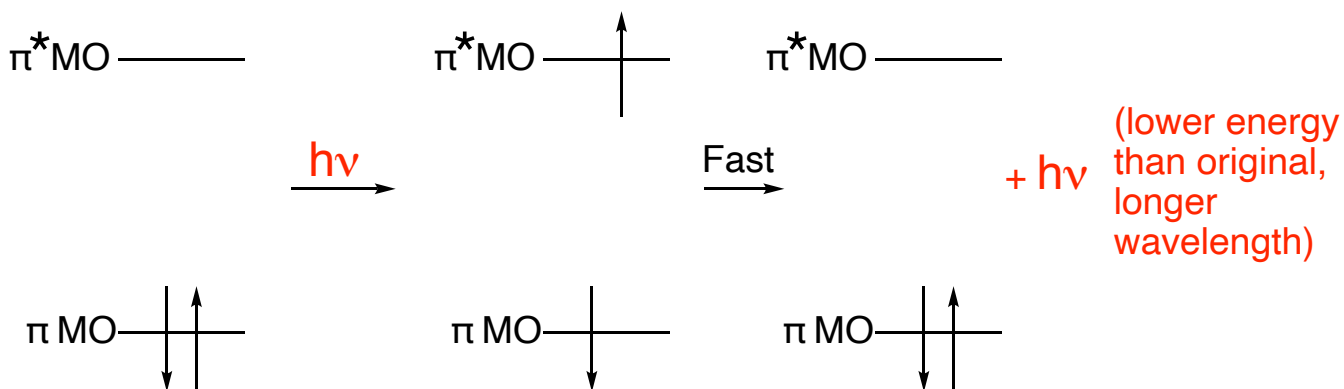


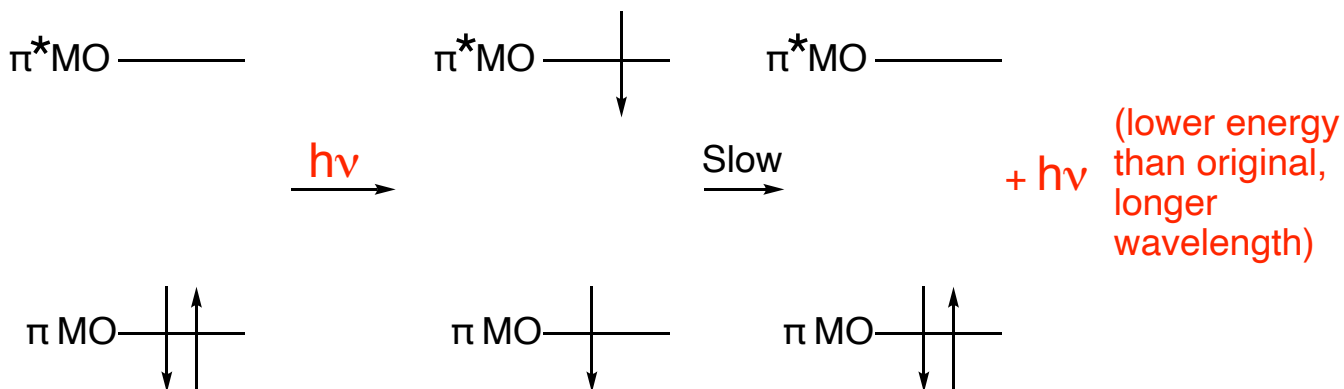
### Generation of heat, Most molecules



### Flourescence - Rigid Molecules, Not uncommon



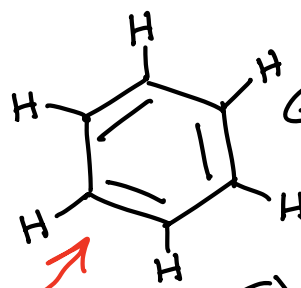
### Phosphorescence - "Glow in the Dark", Rare



# Preview

Called  
"aromaticity"

"aromatic"  
molecule



Benzene

This is  
A LOT

Extraordinarily  
Stable!

~36 kcal/mol  
more stable than  
expected

## Pericyclic Reactions →

$\pi$  bonds  
and  $\sigma$  bonds  
interchange

↳ Happens because  
the transition  
state is super  
stable

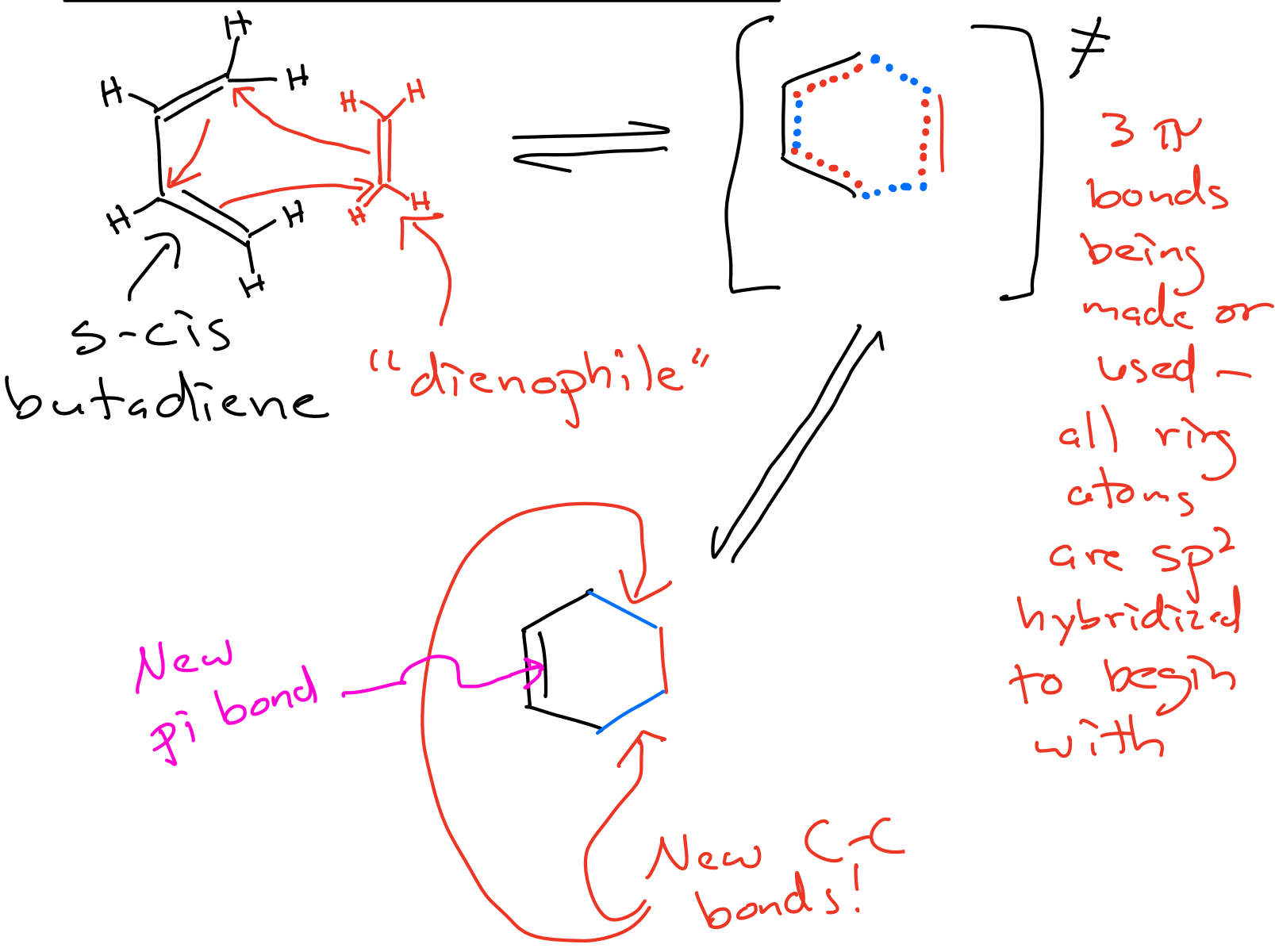
"aromatic" character  
of transition state

Otto!



# Diels-Alder Reaction

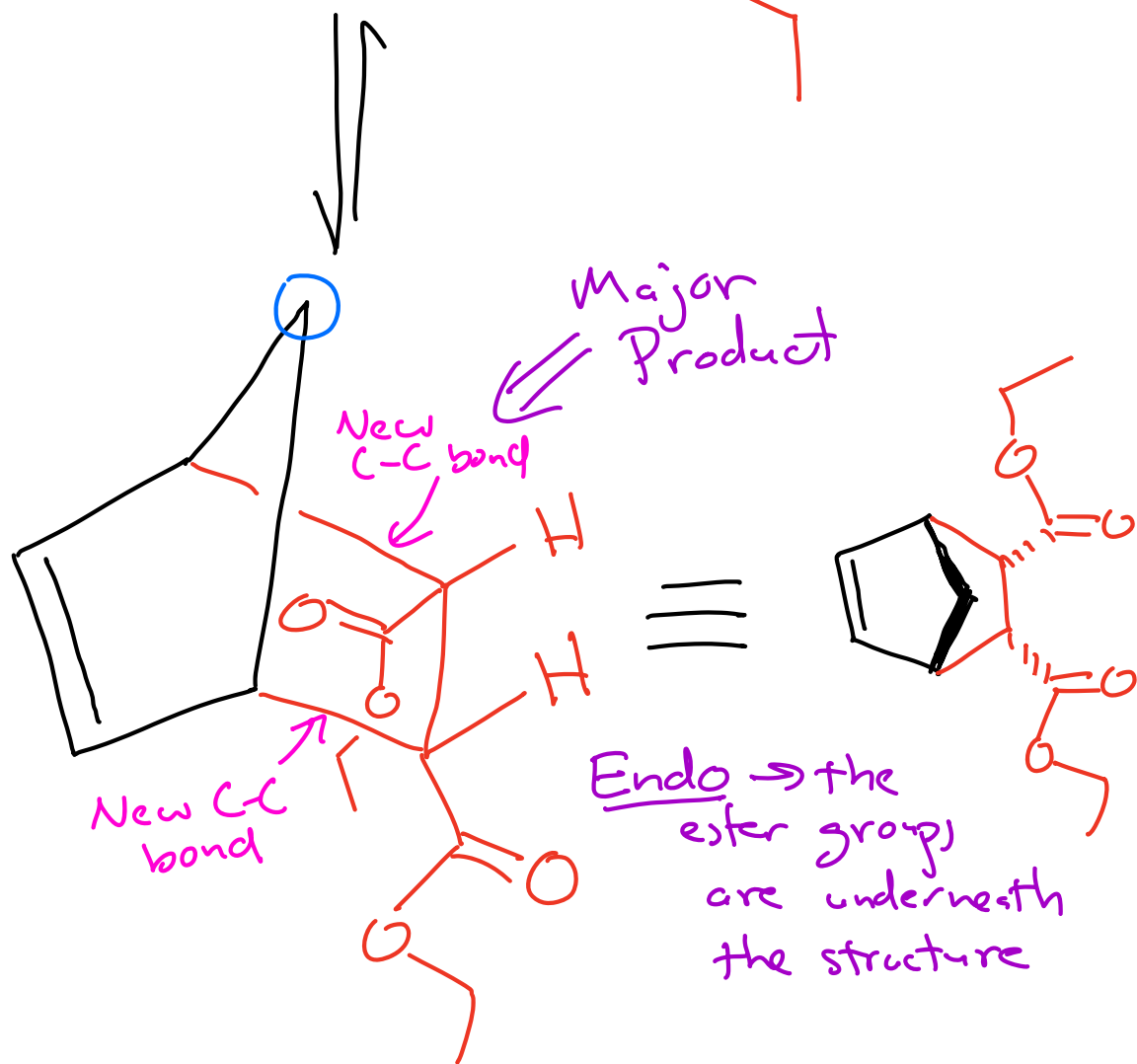
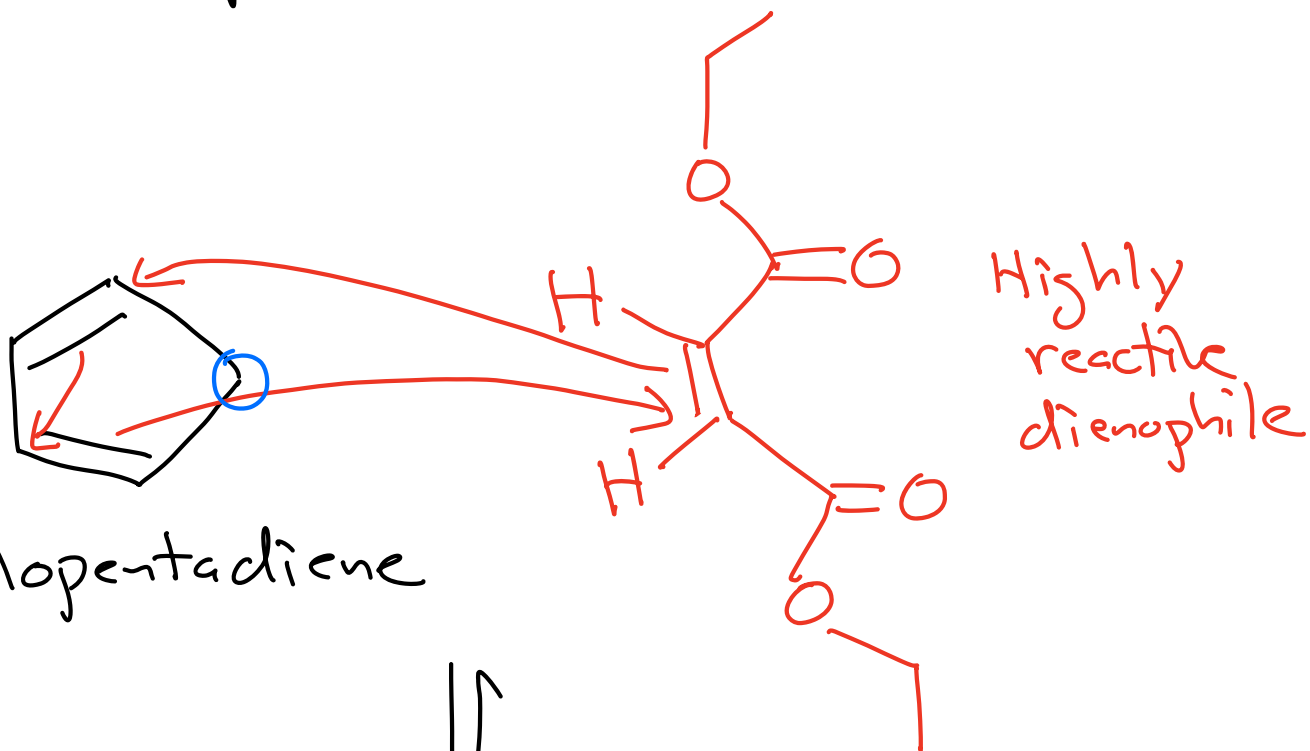
..... bonds being broken  
..... bonds forming



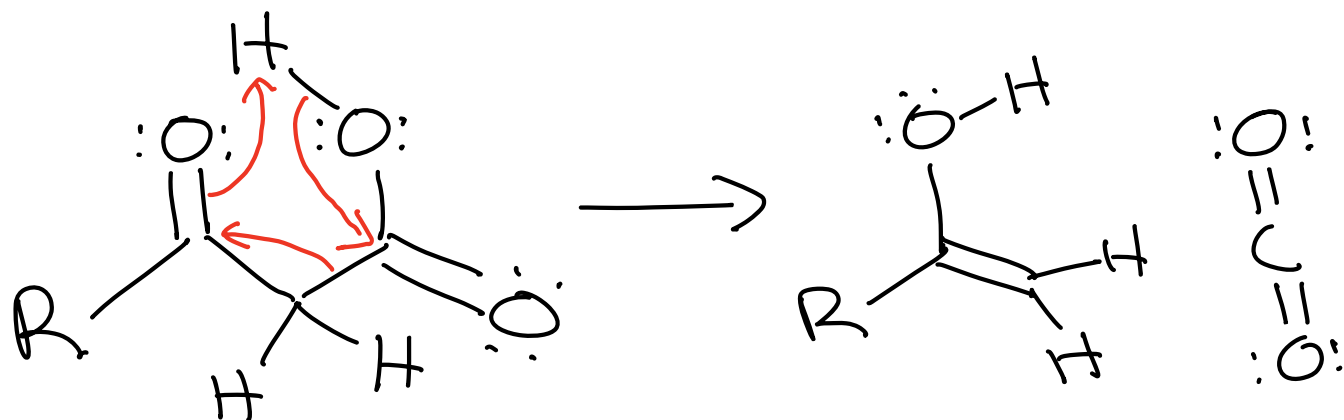
The above reaction gives a poor yield and was used only to illustrate the process  $\rightarrow$  there are many, many known examples of Diels-Alder reactions



The following is the only Diels-Alder reaction you are responsible for in this class



You have seen one other example of this type of reaction:



3  $\pi$  bonds being broken or formed in the transition state  $\rightarrow$  very stable transition state!

Transition state has aromatic character!

That is why  $\beta$ -keto acids and  $\beta$ -diacids decarboxylate when you heat them!

## *The Golden Rules of Organic Chemistry*

Your goal should be to understand, not memorize, the material presented in your organic chemistry course. The following principles should be learned as you begin your study of organic chemistry, then used as a solid foundation for building your understanding throughout the course. These simple ideas explain a great deal about the structures and properties of organic molecules, as well as the characteristic ways in which they react. Thoroughly understanding the following three key principles and related ideas will allow you to develop an intuitive feel for organic chemistry that avoids the necessity of resorting to the far less effective use of extensive memorization.

### *A. Predicting Structure and Bonding*

1. In most stable molecules, all the atoms will have filled valence shells.
2. Five- and six-membered rings are the most stable.
3. There are two possible arrangements of four different groups around a tetrahedral atom.

### *B. Predicting Stability and Properties*

4. The most important question in organic chemistry is "Where are the electrons?"
5. Delocalization of charge over a larger area is stabilizing.
6. ~~Delocalization of unpaired electron density over a larger area is stabilizing.~~
7. Delocalization of pi electron density over a larger area is stabilizing.

### *C. Predicting Reactions*

8. Reactions will occur if the products are more stable than the reactants and the energy barrier is low enough.
9. Functional groups react the same in different molecules.
10. A reaction mechanism describes the sequence of steps occurring during a reaction.
11. Most bond-making steps in reaction mechanisms involve nucleophiles reacting with electrophiles.

All conjugated systems are extra stable, but there is a certain class that is particularly stable:

Aromatic Rings  $\Rightarrow$  Hückels Rules  
(definition)

1) All ring atoms are  $sp^2$  hybridized  
(have a  $2p$  orbital)

2) Ring is flat

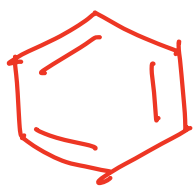
3) Monocyclic

4) 2, 6, 10, 14, 18, 22....  $\pi$  electrons



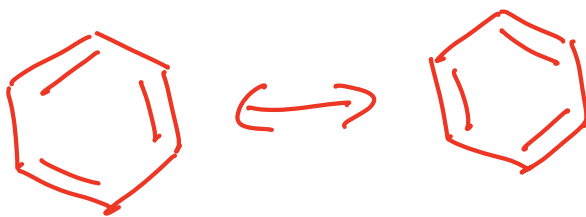
$4n + 2 \pi$  electrons

$n = 0, 1, 2, 3, 4, 5, \dots$

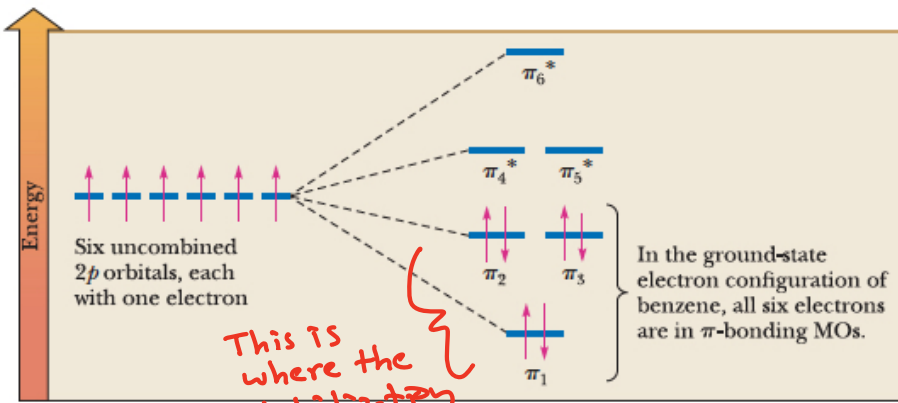


Benzene

← The  $\pi$  electrons of benzene are much less reactive than normal alkenes  $\rightarrow$  benzene only reacts under harsh conditions

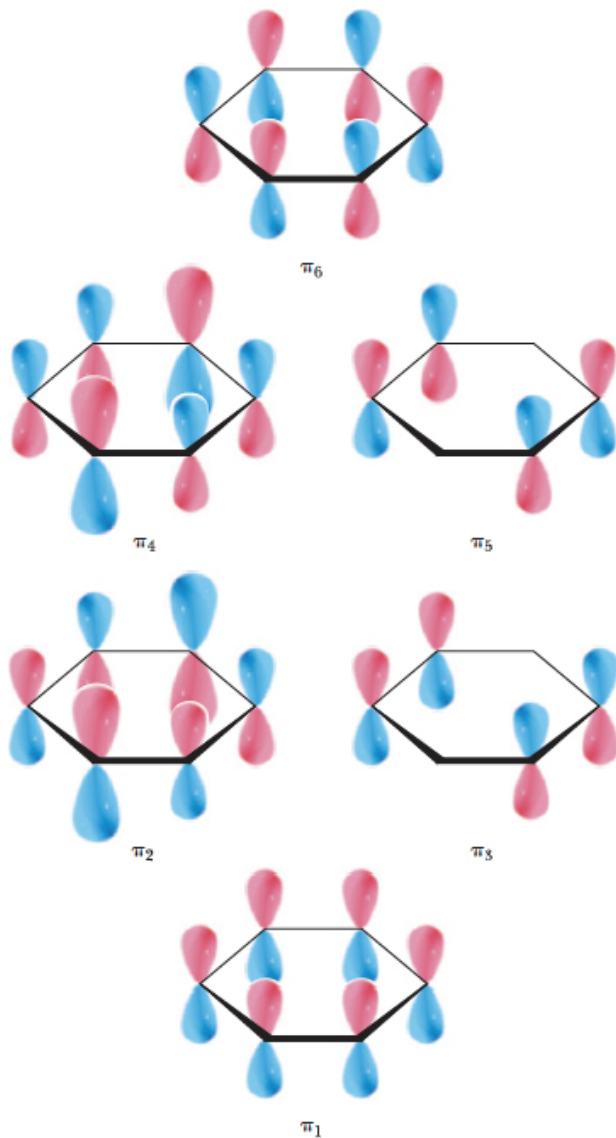


All bonds are the same length!

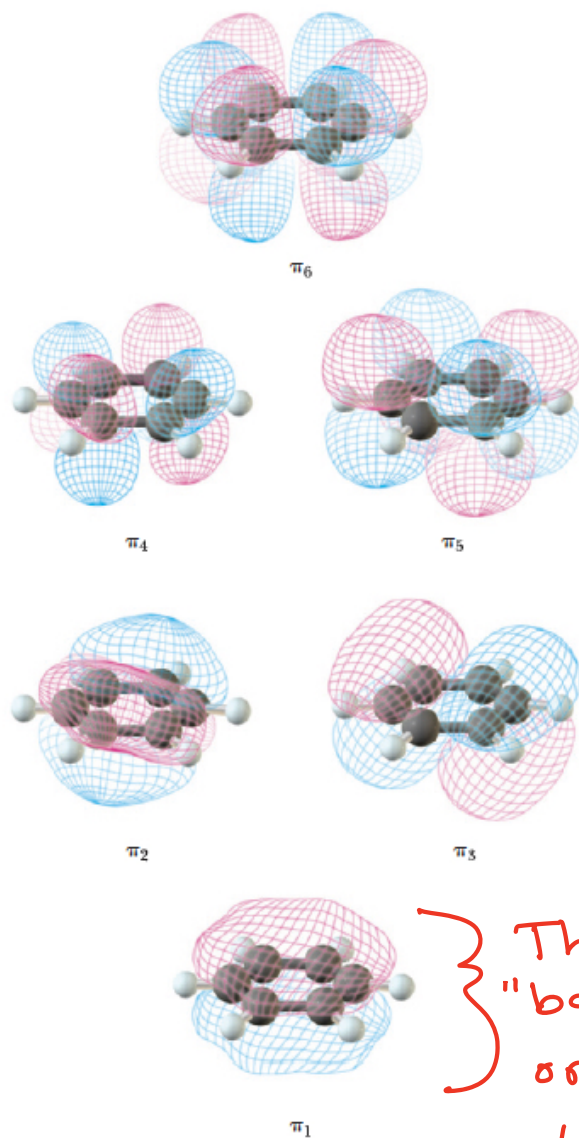


**FIGURE 21.2** The molecular orbital representation of the  $\pi$  bonding in benzene.

(a) Cartoon orbitals



(b) Calculated orbitals



The "bagel" orbital



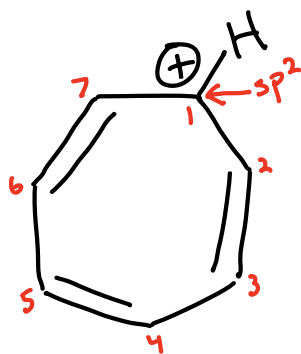
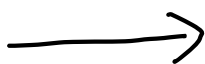
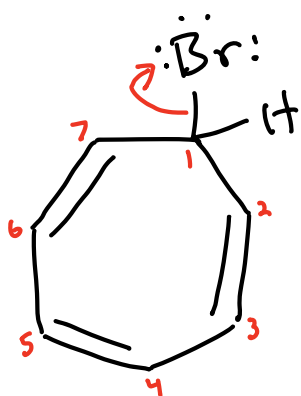
a super stable circular "pi-way"

**FIGURE 21.3** Orbitals for the  $\pi$  system of benzene. (a) Cartoon representations of the six calculated orbitals that chemists routinely draw. These pictures accentuate the fact that various combinations of parallel  $2p$  orbitals lead to the  $\pi$  system of benzene. (b) Calculated orbitals. The three lowest in energy are occupied with electrons (see Figure 21.2). The lowest of these orbitals is the image most chemists use for the  $\pi$  system of benzene: a torus of electron density above and below the ring.

# Two Important Consequences of Aromaticity

- 1) Aromaticity stabilizes ions  $\rightarrow$  anions and cations
- 2) Atoms in molecules will be  $sp^2$  if that produces aromaticity

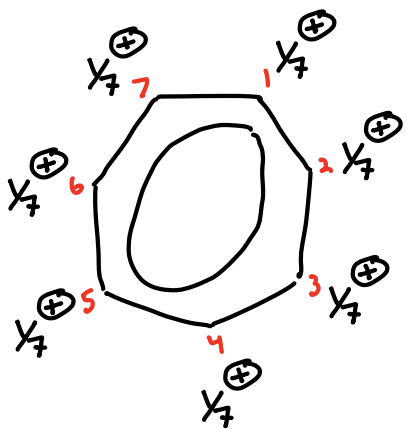
## Tropylium Ion



all ring atoms  
 $sp^2$   
flat  
monocyclic  
6  $\pi$  electrons

Aromatic!

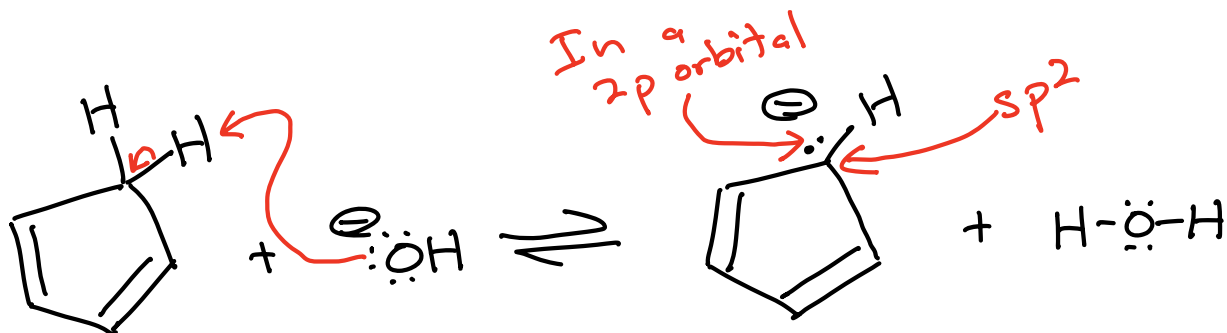
Super stable  
cation



All atoms are  
equivalent  $\rightarrow$   
7 equal contributing  
structures!



# Cyclopentadienyl Anion



$\text{pK}_a = 16$

Aromatic!  
Super stable anion

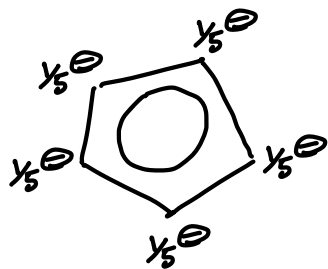
all ring atoms  
sp<sup>2</sup>

flat  
monocyclic

6  $\pi$  electrons

→ 2 electrons  
of lone pair

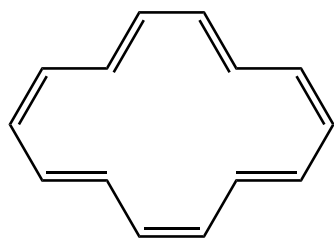
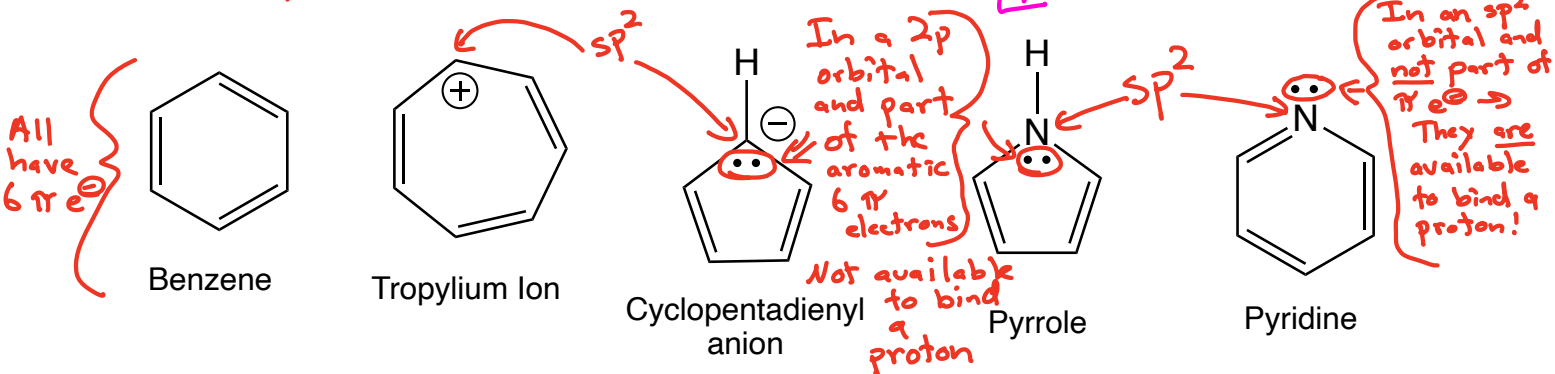
→ 4 electrons  
in the  
2  $\pi$  bonds



All atoms are  
equivalent →  
5 equal contributing  
structures!

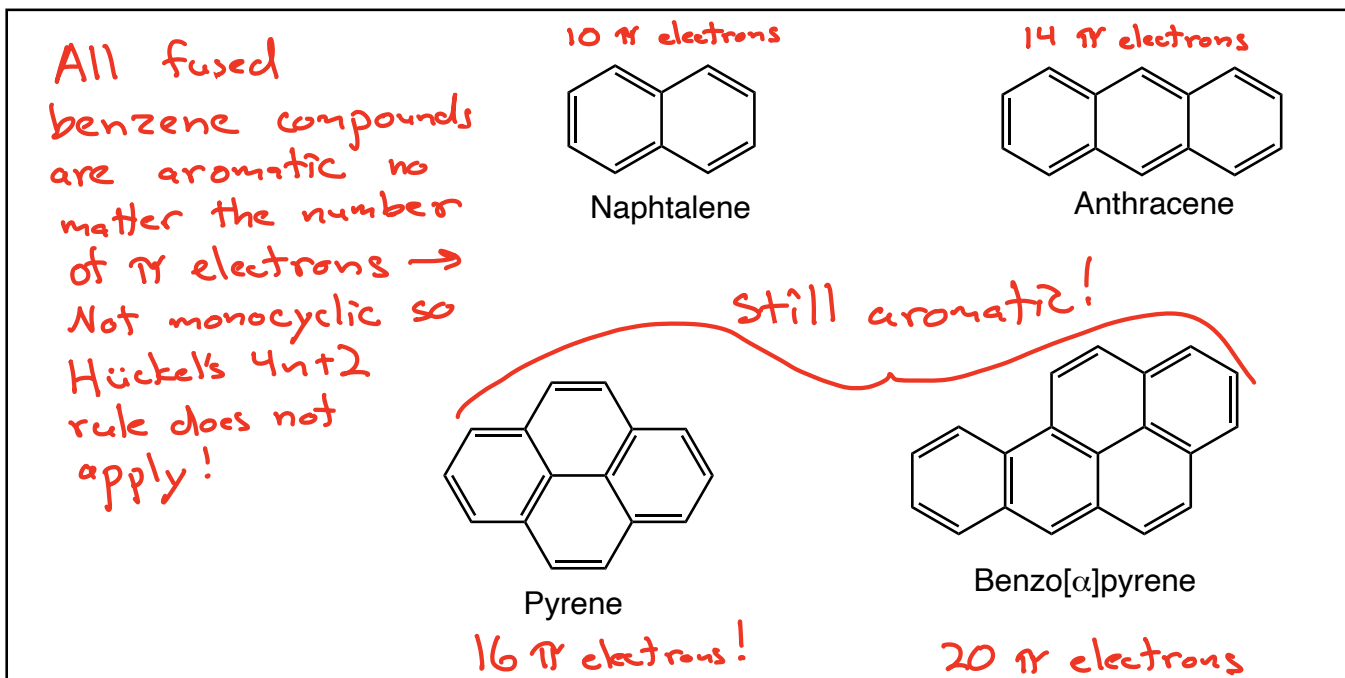
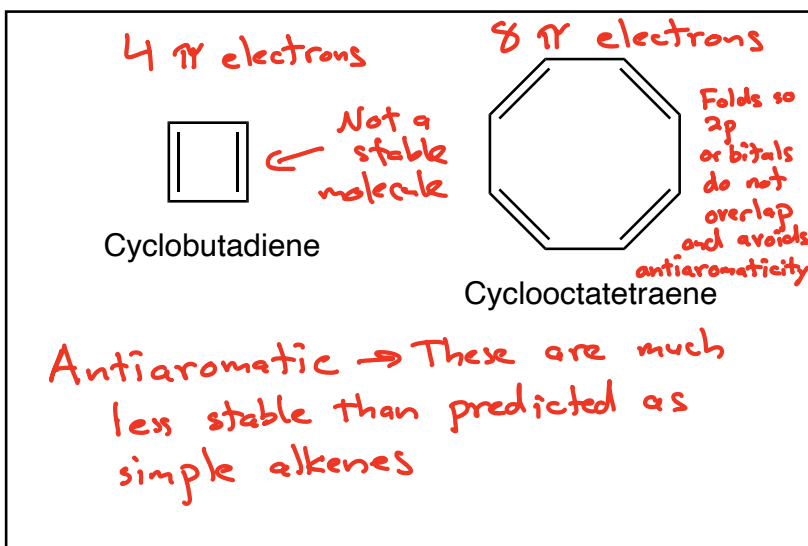
# Hückel's Aromaticity Criteria

- 1) All ring atoms are  $sp^2$  (they have a 2p orbital)
- 2) Flat (so the 2p orbitals overlap)
- 3) Monocyclic (Rule 4) only applies to single rings)
- 4)  $4n+2$  pi electrons (2,6,10,14....)



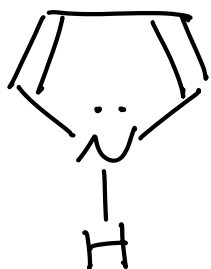
Annulene

14  $\pi$  electrons

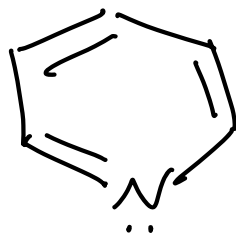


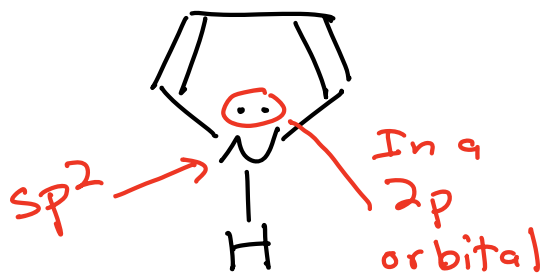


Pyrrrole

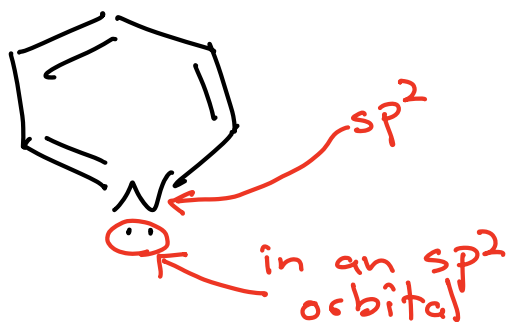


Pyridine





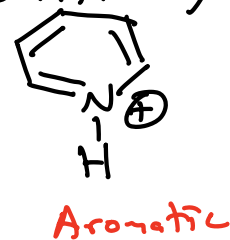
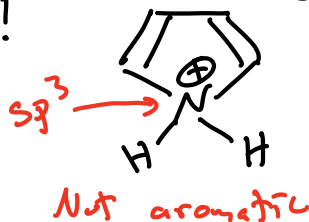
The "lone pair" on N is actually part of the 6  $\pi$  electrons so they are delocalized and not able to bond to a proton



The lone pair on N is available to bond to a proton

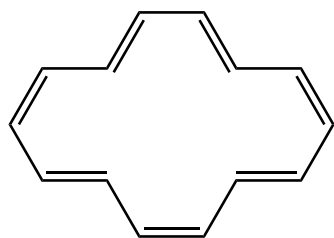
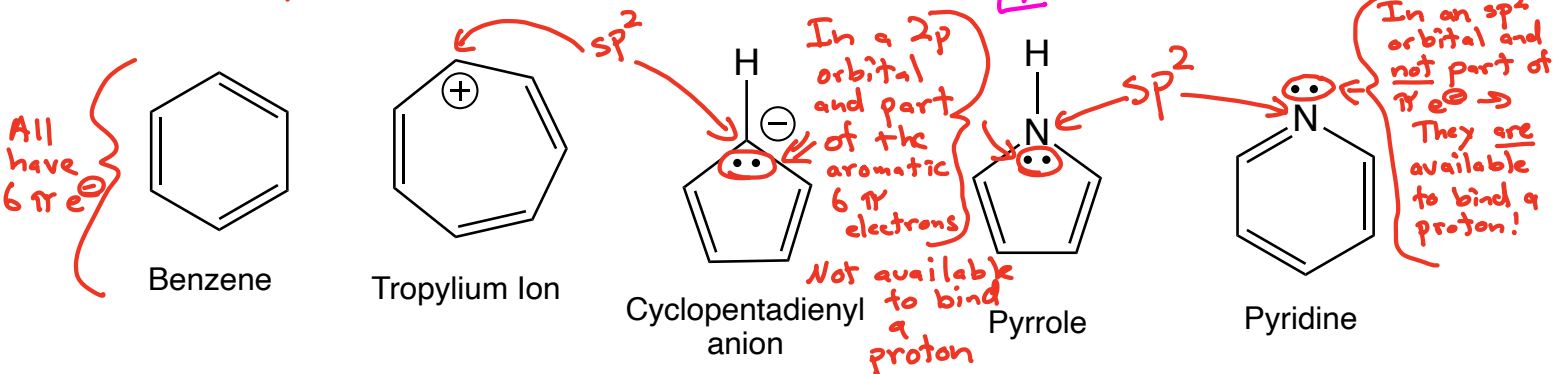
This is the base!

Another way to look at it: Upon protonation, the pyrrole would be forced to lose aromaticity because the N atom would be forced to be  $sp^3$  and only 4  $\pi$  electrons would remain. Losing aromaticity costs far too much energy! Protonated pyridine is still aromatic!



# Hückel's Aromaticity Criteria

- 1) All ring atoms are  $sp^2$  (they have a 2p orbital)
- 2) Flat (so the 2p orbitals overlap)
- 3) Monocyclic (Rule 4) only applies to single rings)
- 4)  $4n+2$  pi electrons (2,6,10,14....)



Annulene

14  $\pi$  electrons

