

# Aromatics II

## MO Theory

## Hückel's Rule

## Spectroscopy

**Ref** 14: 6 – 9, 11 (both ed<sup>ns</sup>)

**Prob** 14: 23, 25 – 27, 29, 34 (8<sup>th</sup> ed)

14: 24, 26 – 28, 30, 35 (9<sup>th</sup> ed)

**Adv Rdg** 15: 1 – 9 (both ed<sup>ns</sup>)

## Review

- benzene, C<sub>6</sub>H<sub>6</sub>, stable & unreactive

- explained by

1.) resonance (*last lecture*)

2.) MO Theory

good starting point:

“MO's are formed by combination of similar AO's; similar in *energy, size, shape*”

recall: # of MO's = # of “originating” AO's

applied to benzene:

p AO's → π MO's

**6 p AO's → 6 π MO's**

Illustration:

“6 p AO's combine in various ways to

form 6 π MO's”

( *in-phase, out-of-phase, varying # of nodes ...* )

## Nodes in MO's

(areas w/ zero e<sup>-</sup> density)

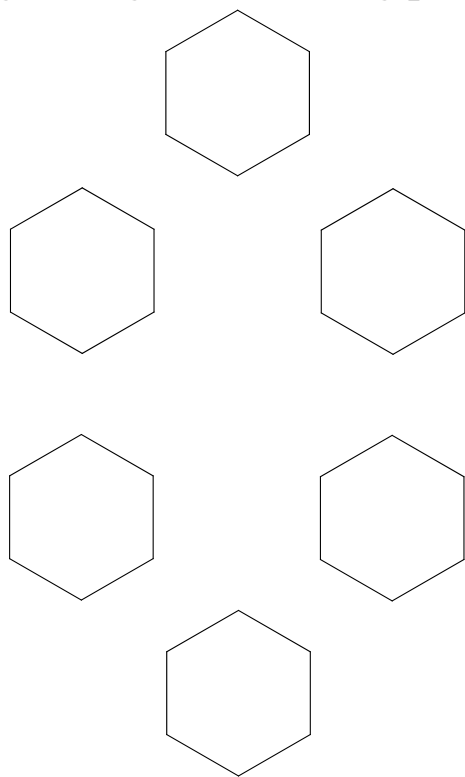
rule: “as # of nodes increases,  
the energy of the MO ↑ ”

$\left[ \begin{array}{l} \text{b/c less binding,} \\ \text{it is easier for the molecule to fall apart} \end{array} \right]$

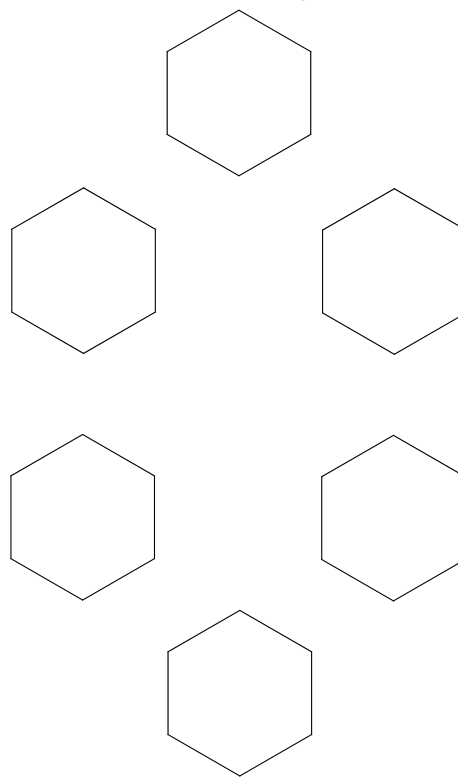
## MO Energy Diagram

*comment: the # of MO's at lvl 2 & 3 doubles up  
because of symmetry considerations*

## Originating/Contributing p AO's



## Resulting MO's



## Occupation of MO's by e<sup>-</sup>'s

In the ground state of the benzene molecule:

all e<sup>-</sup>'s are in lower energy orbitals ("bonding"),  
none in higher energy orbitals ("anti-bonding"),  
when compared with the originating p AO's

∴ reason for **aromatic stabilization**

(similar MO arguments apply to all other aromatic systems)

## Hückel's Rule

predicts aromaticity of org. cmpds.  
(containing rings w/ high level of unsaturation)

A. **Aromatic**, if

1.) monocyclic system

(rule can be extended to polycyclic systems,  
such as naphthalene)

2.) ring planar

3.) each atom in ring can be sp<sup>2</sup> hybridised;  
p AO's form π system ("sideways overlap")

4.) π system contains

## B. Anti-Aromatic (unstable), if

- |     |   |               |
|-----|---|---------------|
| 1.) | } | same as in A. |
| 2.) |   |               |
| 3.) |   |               |

## C. Non-Aromatic (behave “normally”)

all others;

esp., non-planar systems

## B.) Anti-aromatic (*very unstable*)

## C.) Non-aromatic

looks anti-aromatic: 8 e<sup>-</sup>'s, n = 2, “4n rule”;

but behaves like normal alkene;

conclusion: system must be non-planar,  
approx. “tub-like”:

# Examples

## A.) Aromatic

1.)            n = 1, 6 e<sup>-</sup>'s

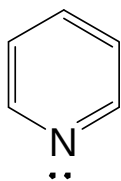
2.)            n = 4, 4n + 2 = 16 + 2 = 18 e<sup>-</sup>'s

# Variety of Aromatic Structures

- can contain heteroatoms (N, O S, ...)
- may be ionic

Examples:

## Pyridine



N is  $sp^2$  hybridised;

has 3  $sp^2$  orbitals and 1 “pure” p orbital:

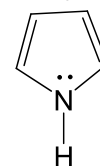
- p orbital involved in aromaticity
- 2  $sp^2$  orbitals form  $\sigma$  bonds w/ C atoms
- 1  $sp^2$  orbital contains lone pair of  $e^-$ 's

$\therefore$  lone pair not used in aromaticity

“remains exposed”

**remains quite basic**

## Pyrrole



N is  $sp^2$  hybridised;

has 3  $sp^2$  orbitals and 1 “pure” p orbital:

$sp^2$  orbitals form 3  $\sigma$  bonds;

p orbital ( w/ 2  $e^-$ 's )

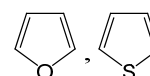
contributes to aromatic system

$\therefore$  lone pair “used up” in aromaticity

“not exposed”

**pyrrole much weaker base than pyridine**

similar arguments for furan, thiophen (HMWK)



## Anions

Ex. cyclopentadienyl anion,  $C_5H_5^-$

(forms easily from cyclopentadiene with bases such as  $OH^-$ )

planar, monocyclic system,  
all C's  $sp^2$  hybridised,  
conjugated  $\pi$  system  
6  $e^-$ 's (Hückel #)

**aromatic**

## Cations

Ex.1.: cyclopropenyl cation,  $C_3H_3^+$

(suitable Lewis acid, trityl cation:  $Ph_3C^+ BF_4^-$ )

aromatic requirements satisfied:

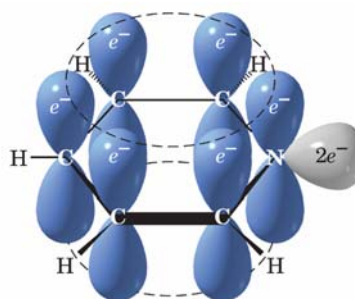
planar, monocyclic system,  
all C's  $sp^2$  hybridised,  
conjugated  $\pi$  system  
2  $e^-$ 's (Hückel #)

**aromatic**

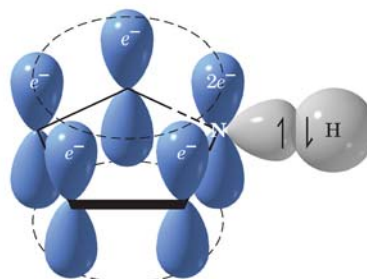
Ex.2.: cycloheptatrienyl cation,  $C_7H_7^+$ 

aromatic requirements satisfied, as before !

## Solomons Fig's: Pyridine &amp; Pyrrole

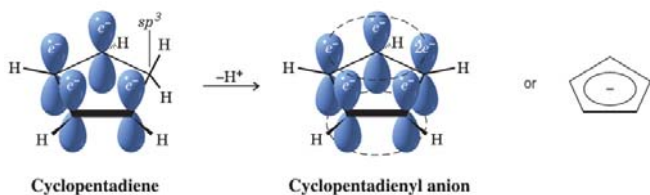


**FIGURE 14.20** The stylized  $p$  orbital structure of pyridine.

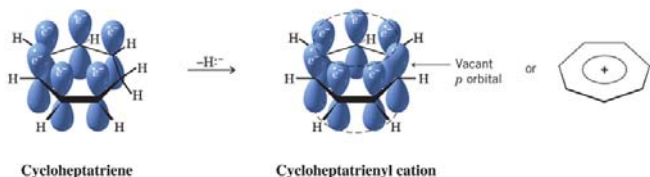


**FIGURE 14.21** The stylized  $p$  orbital structure of pyrrole. (Compare with the orbital structure of the cyclopentadienyl anion in Fig. 14.10.)

## Solomons Fig's: Cyclopentadienyl Anion &amp; Cycloheptatrienyl Cation



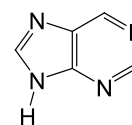
**FIGURE 14.10** The stylized  $p$  orbitals of cyclopentadiene and of the cyclopentadienyl anion.



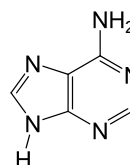
**FIGURE 14.12** The stylized  $p$  orbitals of cycloheptatriene and of the cycloheptatrienyl (tropylium) cation.

Complex Heterocyclic Aromatics  
*important in Biochemistry*

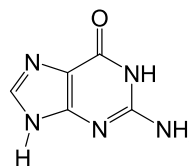
pyrimidine



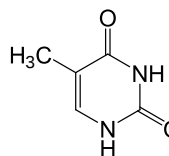
purine



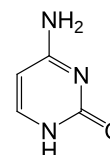
adenine



guanine



thymine



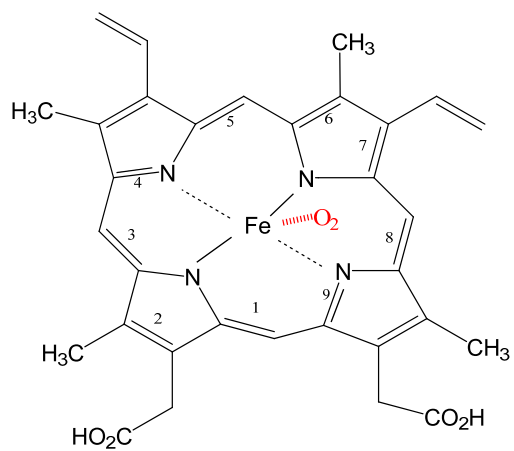
cytosine

DNA Bases

# Another Aromatic System

*of biological importance*

Heme, part of hemoglobin,  
oxygen carrier in red blood cells



numbers indicate double bonds that are part of aromatic system

$n = 4$ ,  $4n + 2 = 18$ ; 18  $e^-$ s = aromatic

b.) J values

ortho = 9 Hz

meta = 2 Hz

para ~ 0.5 Hz (negligible)

c.) effect of substituents

# Spectroscopy

1.) NMR

a.)  $\delta$  (aromatic) ~ 7.0 ppm

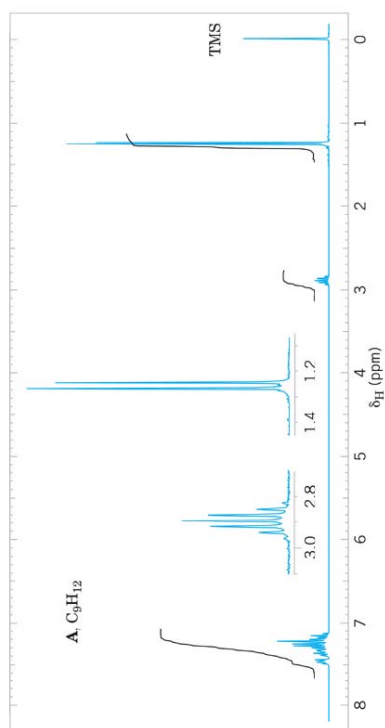
HMWK:

do similar calc<sup>ns</sup> for 60 and 500 MHz instruments.

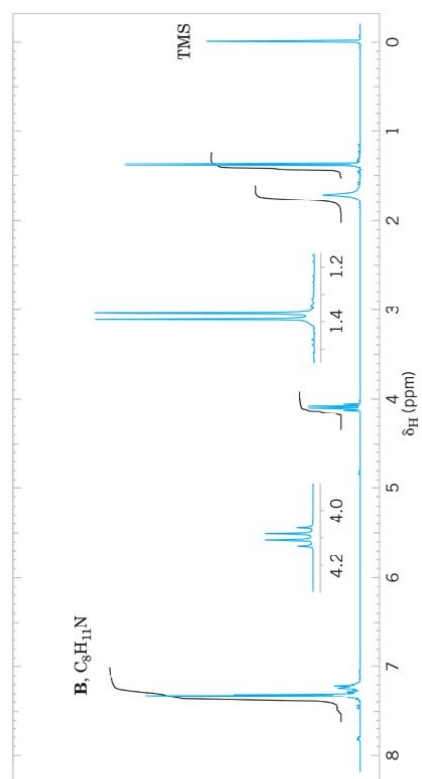
# 2.) IR

| <u>“aromatic bond”</u> | <u>type</u> | <u>wavenumber</u>     |
|------------------------|-------------|-----------------------|
| $= C - H$              | stretch     | ~ 3030 $cm^{-1}$      |
| $C = C$                | stretch     | 1600 – 1650 $cm^{-1}$ |
| $= C - H$              | bend        | 900 – 700 $cm^{-1}$   |

Solomons Fig. 14.27 Aromatic NMR Spectra #1



Solomons Fig. 14.27 Aromatic NMR Spectra #2



Solomons Fig. 14.27 Aromatic NMR Spectra #3

