### Your lecturers







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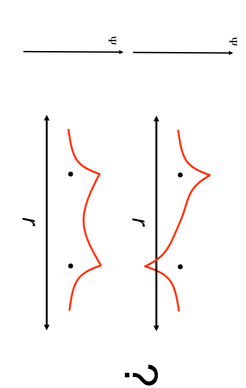
#### **Chemistry 2**

#### **Quantum Mechanics in Chemistry** Lecture 1



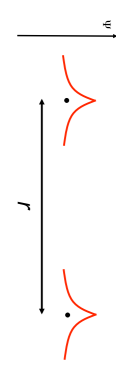
#### Revision – H<sub>2</sub><sup>+</sup>

At equilibrium, we have to make the lowest energy possible using the 1s functions available



#### Revision $-H_2^+$

- Near each nucleus, electron should behave as a 1s electron.
- At dissociation, 1s orbital will be exact solution at each nucleus



#### П П 1s<sub>A</sub> anti-bonding anti-bonding bonding bonding Revision – H<sub>2</sub><sup>+</sup> 1s<sub>B</sub> $1s_B$ ¥ Ψ Ψ ¥ $1s_A$ $\Psi = 1s_A - 1s_B$ $\Psi = 1s_A + 1s_B$ $\Psi = 1s_A + 1s_B$ $\Psi = 1s_A - 1s_B$ 1s<sub>B</sub> 1s<sub>B</sub> $1s_B$ $1s_B$

# 2<sup>nd</sup> row homonuclear diatomics

 $Revision - H_2$ 

Now what do we do? So many orbitals!

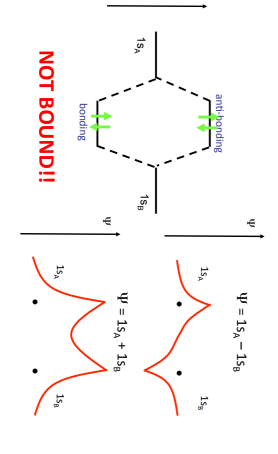
2p 2s

2p

**2**s

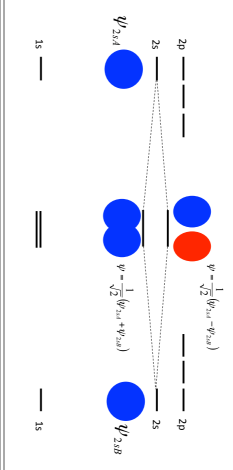


### Revision – He<sub>2</sub>



### Interacting orbitals

 Orbitals interact proportionally to the inverse of their energy difference. Orbitals of the same energy interact completely, yielding completely mixed linear combinations.



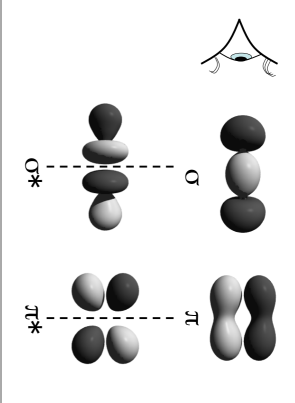
### Interacting orbitals

Orbitals can interact and combine to make new approximate solutions to the Schrödinger equation. There are two considerations:

1.Orbitals interact **inversely** proportionally to their **energy difference**. Orbitals of the same energy interact completely, yielding completely mixed linear combinations. In quantum mechanics, energy and frequency are related (E=hv). So, energy matching is equivalent to the phenomenon of **resonance**.

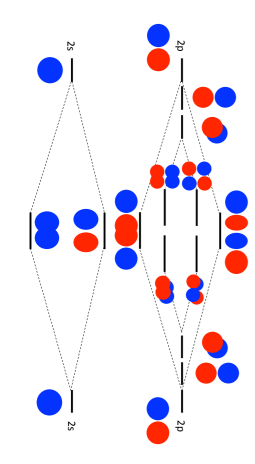
2.The extent of orbital mixing is given by the **resonance integral**  $\beta$ . We will show how beta is calculated in a later lecture.

# **Molecular Orbital Theory - Revision**



## (First year) MO diagram

Orbitals interact *most* with the corresponding orbital on the other atom to make perfectly mixed linear combinations. (we ignore core).



### Interacting orbitals

1. The extent of orbital mixing is given by the integral

$$\beta$$
 = something

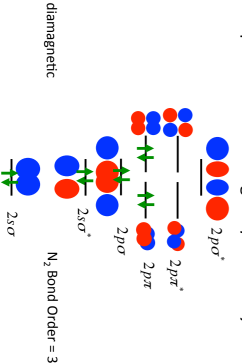


The 2s orbital on one atom *can* interact with the 2p from the other atom, but since they have different energies this is a smaller interaction than the 2s-2s interaction. We will deal with this later.

**1**s

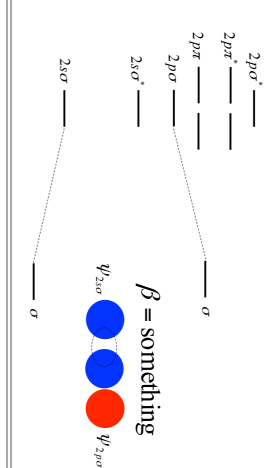
# **Molecular Orbital Theory - Revision**

Can predict bond strengths qualitatively



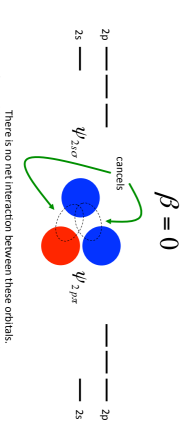
# More refined MO diagram

 $\boldsymbol{\sigma}$  orbitals can now interact



### Interacting orbitals

1. The extent of orbital mixing is given by the integral



**1**s

The positive-positive term is cancelled by the positive-negative term

15

# More refined MO diagram

 $\boldsymbol{\pi}$  orbitals do not interact ۹

 $2p\pi^*$ 

 $2p\sigma^*$ 

$$2p\sigma$$
  $\sigma$   $\sigma$   $\sigma$ 

 $2s\sigma^*$ 

# $2s\sigma$

# More refined MO diagram

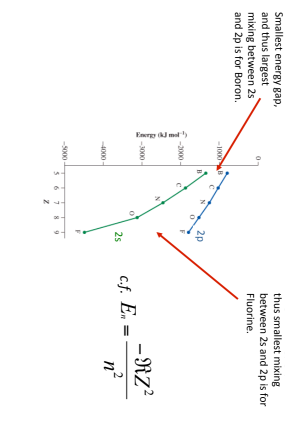
 $\sigma^*$  orbitals can interact  $\sigma^*$ 

 $2p\sigma$ 

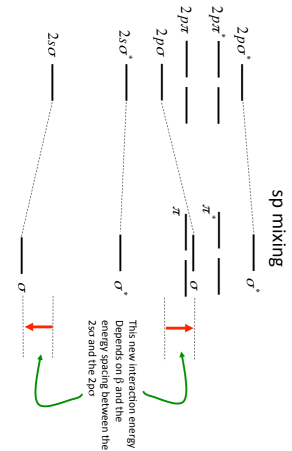
$$2s\sigma^*$$
  $\sigma^*$ 

#### sp mixing

Largest energy gap, and



# More refined MO diagram



### Learning outcomes



- energy difference, and the resonance integral,  $\beta$ . •Use the principle that the mixing between orbitals depends on the
- structure in simple organic molecules. •Apply the separation of  $\sigma$  and  $\pi$  bonding to describe electronic
- molecules in terms of s-p mixing. • Rationalize differences in orbital energy levels of diatomic

#### sp mixing

### **Practice Questions**

- Why is s-p mixing more important in Li<sub>2</sub> than in F<sub>2</sub>?
- How many core,  $\sigma\text{-bonding,}$  and  $\pi\text{-electrons}$  are there in
- a) acetylene
- ethylene

benzene

d) buckminsterfullerene

Check that your total number of electrons agrees with what is expected (6 per carbon, 1 per hydrogen).

#### Next lecture

 Particle in a box approximation solving the Schrödinger equation.

### Week 10 tutorials

 Wavefunctions and the Schrödinger equation.

weakly bound

paramagnetic

diamagnetic

B<sub>2</sub>

 $\frac{\mathsf{N}}{\mathsf{N}}$