

## Learning outcomes from Lecture 1

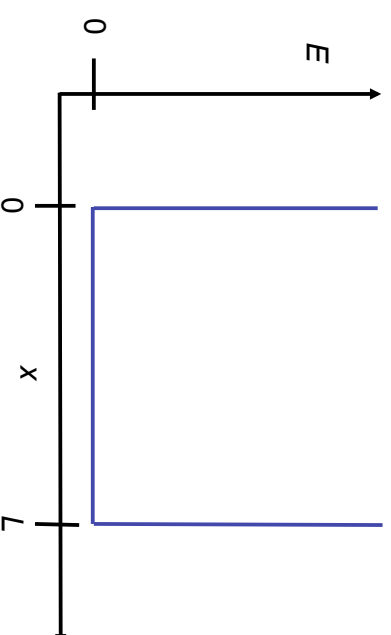
- Use the principle that the mixing between orbitals depends on the energy difference, and the resonance integral,  $\beta$ .
- Apply the separation of  $\sigma$  and  $\pi$  bonding to describe electronic structure in simple organic molecules.
- Rationalize differences in orbital energy levels of diatomic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

## Assumed knowledge for today

Be able to predict the geometry of a hydrocarbon from its structure and account for each valence electron. Predict the hybridization of atomic orbitals on carbon atoms.

## “The particle in a box”

- The box is a 1d well, with sides of infinite potential, where the particle cannot be ...



see worksheet

## The de Broglie Approach

- The wavelength of the wave associated with a particle is related to its momentum:

$$p = mv = h / \lambda$$

- For a particle with only kinetic energy:

$$E = \frac{1}{2} mv^2 = p^2 / 2m = h^2 / 2m\lambda^2$$

- For a free particle,  $\lambda$ , can have any value:  
 $E$  for a free particle is *not* quantized

## Chemistry 2

### Lecture 2

### Particle in a box approximation



## The Schrödinger Equation Approach

- The total energy is extracted by the Hamiltonian operator.
- These are the “observable” energy levels of a quantum particle

$$\hat{H}\Psi(x) = \epsilon_i \Psi(x)$$

Hamiltonian operator

Energy eigenfunction

Energy eigenvalue

## “The particle in a box”

- Energy is quantized:  

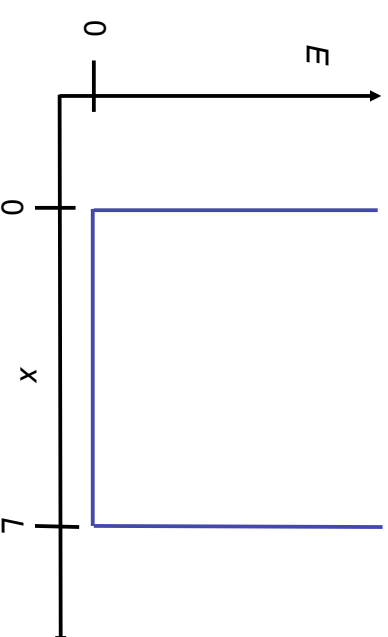
$$E_n = h^2 n^2 / 8mL^2$$
- Lowest energy (zero point) is not zero:  

$$E_{n=1} = h^2 / 8mL^2$$
- Allowed levels are separated by:  

$$\Delta E = E_{n+1} - E_n = h^2(2n+1) / 8mL^2$$

## “The particle in a box”

- The box is a 1d well, with sides of infinite potential, where the electron cannot be ...



## The Schrödinger equation

- The Hamiltonian has parts corresponding to Kinetic Energy and Potential Energy. In one dimension, x:  

$$\hbar = h / 2\pi \text{ ("h bar")}$$

$$\hat{H}\Psi = \left( -\frac{\hbar^2}{2m} \frac{\partial^2}{\partial x^2} + V(x) \right) \Psi$$

Hamiltonian operator

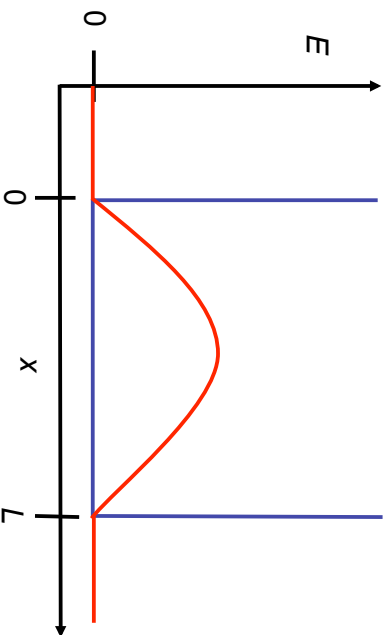
Kinetic Energy

Potential Energy

## "The particle in a box"

- Let's try some test solutions

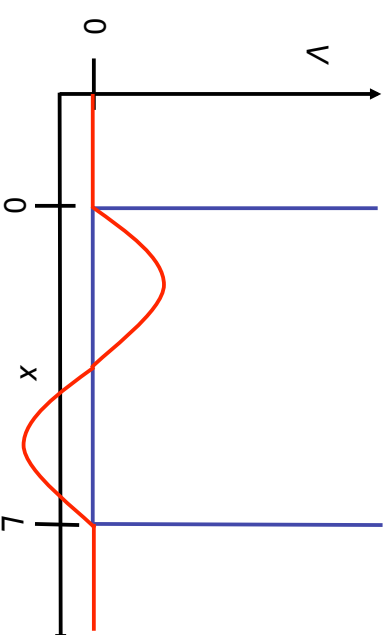
$$\psi = \sin(\pi x/L) \quad \{x>0; x<L$$



## "The particle in a box"

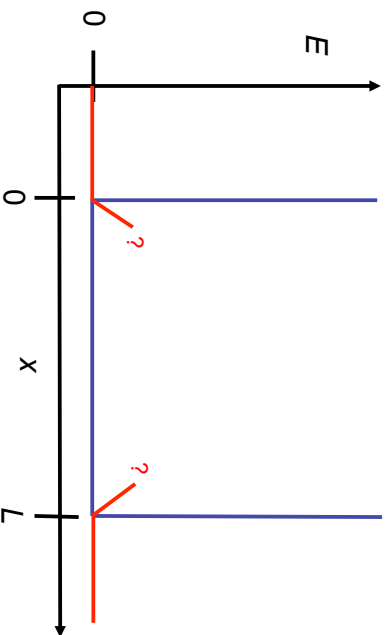
- Other solutions?

$$\psi = \sin(2\pi x/L) \quad \{x>0; x<L$$



## "The particle in a box"

- The particle cannot exist outside the box...  
 $\psi = 0 \quad \{x<0; x>L$  (boundary conditions)



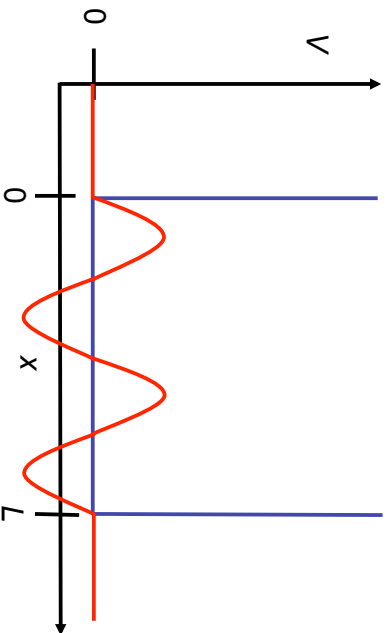
## "The particle in a box"

$$\begin{aligned} \hat{H}\Psi &= -\frac{\hbar^2}{2m} \frac{\partial^2}{\partial x^2} \Psi + V(x)\Psi \\ &= -\frac{\hbar^2}{2m} \frac{\partial}{\partial x} \left( \frac{\partial}{\partial x} \Psi \right) + 0\Psi \quad \text{Zero potential inside box} \\ &= -\frac{\hbar^2}{2m} \frac{\partial}{\partial x} \left( \frac{\partial}{\partial x} \sin(\pi x/L) \right) \\ &= -\frac{\hbar^2}{2m} \frac{\partial}{\partial x} (\pi/L \cos(\pi x/L)) \\ &= -\frac{\hbar^2}{2m} (-\pi^2/L^2 \sin(\pi x/L)) \\ &= \frac{\hbar^2 \pi^2}{2mL^2} \sin(\pi x/L) = \frac{\hbar^2 \pi^2}{2mL^2} \Psi = \epsilon \Psi !!! \end{aligned}$$

## "The particle in a box"

- Other solutions?

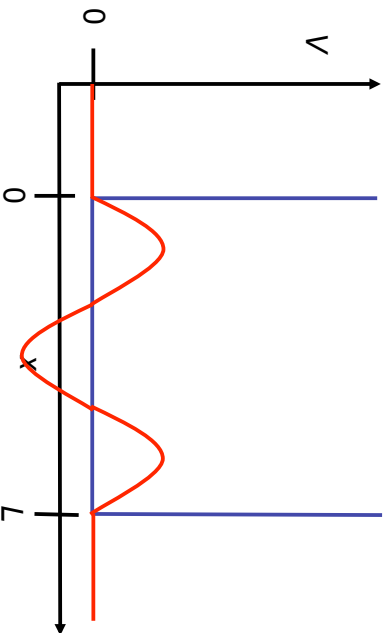
$$\psi = \sin(4\pi x/L) \quad \{x>0; x<L$$



## "The particle in a box"

- Other solutions?

$$\psi = \sin(3\pi x/L) \quad \{x>0; x<L$$



## "The particle in a box"

$$\psi = \sin(n\pi x/L) \quad \{x>0, x<L; n>0$$
$$\epsilon_n = \hbar^2 n^2 \pi^2 / 2mL^2$$

Philosophical question: why is  $n = 0$  not an appropriate solution?

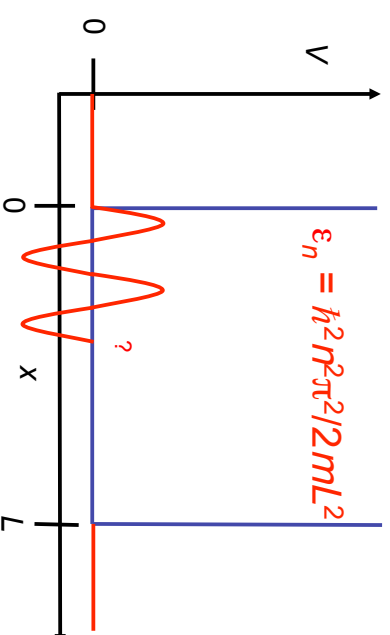
Hint: what's the probability of observing the particle?

## "The particle in a box"

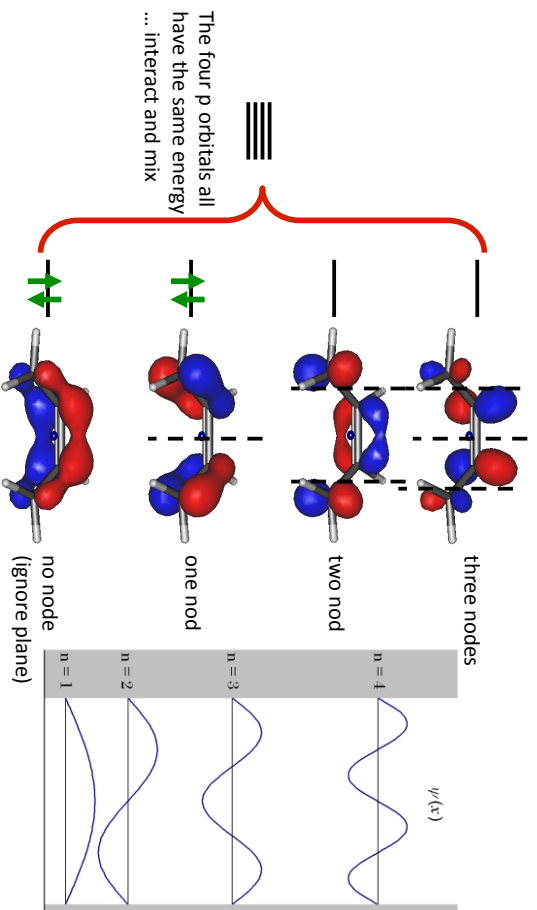
- Other solutions?

$$\psi = \sin(n\pi x/L) \quad \{x>0; x<L$$

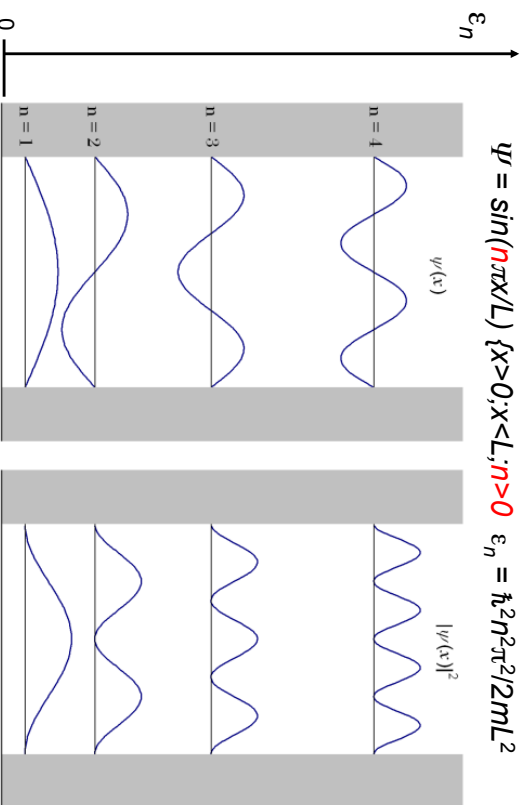
$$\epsilon_n = \hbar^2 n^2 \pi^2 / 2mL^2$$



## *$\pi$ orbitals of cis-butadiene*

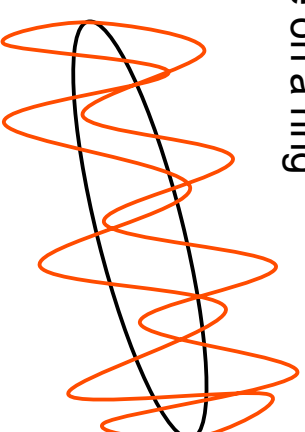


## *“The particle in a box”*



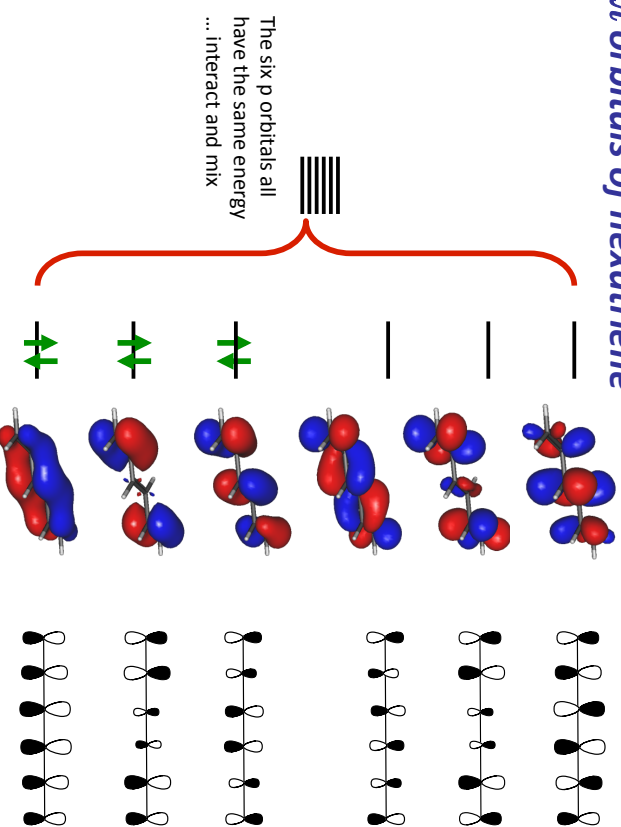
## *Something to think about*

- Particle on a ring



Must fit even wavelengths into whole cycle

## *$\pi$ orbitals of hexatriene*



## Learning outcomes



- Be able to explain why confining a particle to a box leads to quantization of its energy levels
- Be able to explain why the lowest energy of the particle in a box is not zero
- Be able to apply the particle in a box approximation as a model for the electronic structure of a conjugated molecule (given equation for  $E_n$ ).

## Next lecture

- Particle-on-a-ring model

## Week 10 tutorials

- Schrödinger equation and molecular orbitals for diatomic molecules

## Practice Questions

1. The energy levels of the particle in a box are given by  $\epsilon_n = \hbar^2 n^2 p^2 / 2mL^2$ .
  - (a) Why does the lowest energy correspond to  $n = 1$  rather than  $n = 0$ ?
  - (b) What is the *separation* between two adjacent levels?  
(*Hint:  $\Delta\epsilon = \epsilon_{n+1} - \epsilon_n$* )
  - (c) The  $\pi$  chain in a hexatriene derivative has  $L = 973$  pm and has 6  $\pi$  electrons. What is energy of the HOMO – LUMO gap? (*Hint: remember that 2 electrons are allowed in each level.*)
  - (d) What does the particle in a box model predicts happens to the HOMO – LUMO gap of polyenes as the chain length increases?