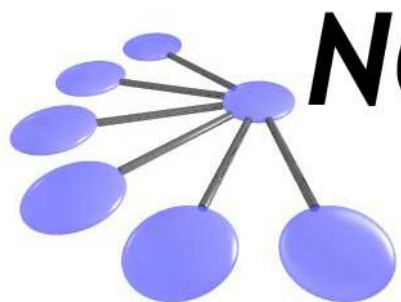
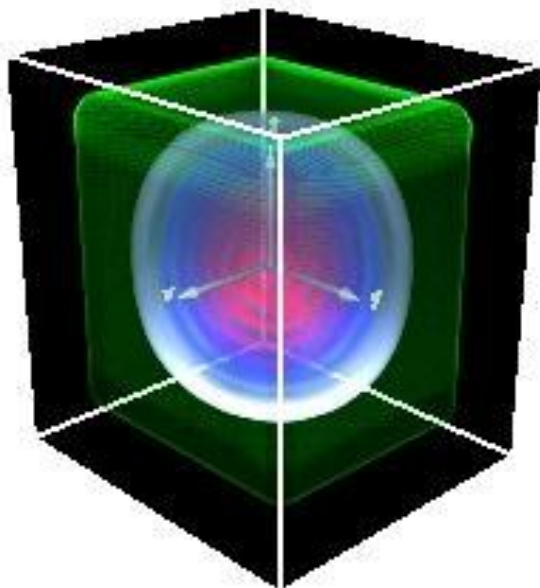


# Network for Computational Nanotechnology (NCN)

Purdue, Norfolk State, Northwestern, MIT, Molecular Foundry, UC Berkeley, Univ. of Illinois, UTEP



## NCN First Time User Guide to Quantum Dot Lab\*



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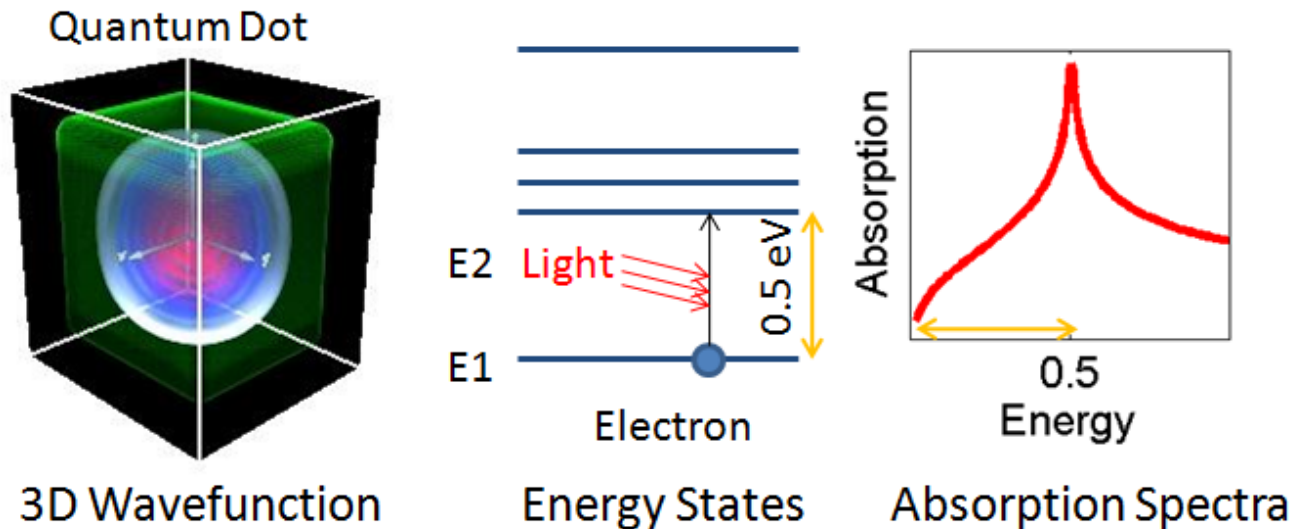
\*\*email:kim568@purdue.edu

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# Introduction

- The quantum dot lab is a tool that solves the Schrodinger equation for an electron in a quantum dot.
- The quantum dot lab yields the wavefunction, the electron energy levels, and the optical transition rates/absorption strength of an electron.



- A detailed introduction to the quantum dot lab also can be found at <https://www.nanohub.org/resources/4194>.

# First Look

1 Input → 2 Simulate ? About this tool  
Questions?

Number of States:

Surface passivation:  "yes"

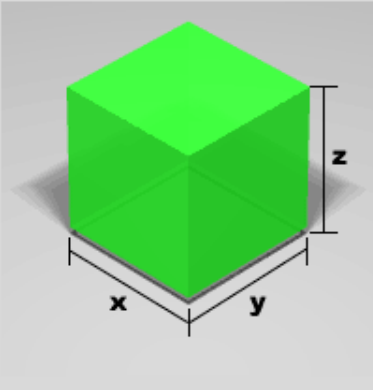
Device Structure | Light Source

Geometry:

X dimensions:

Y dimensions:

Z dimensions:



Effective Mass:

Discretization:

Energy gap:

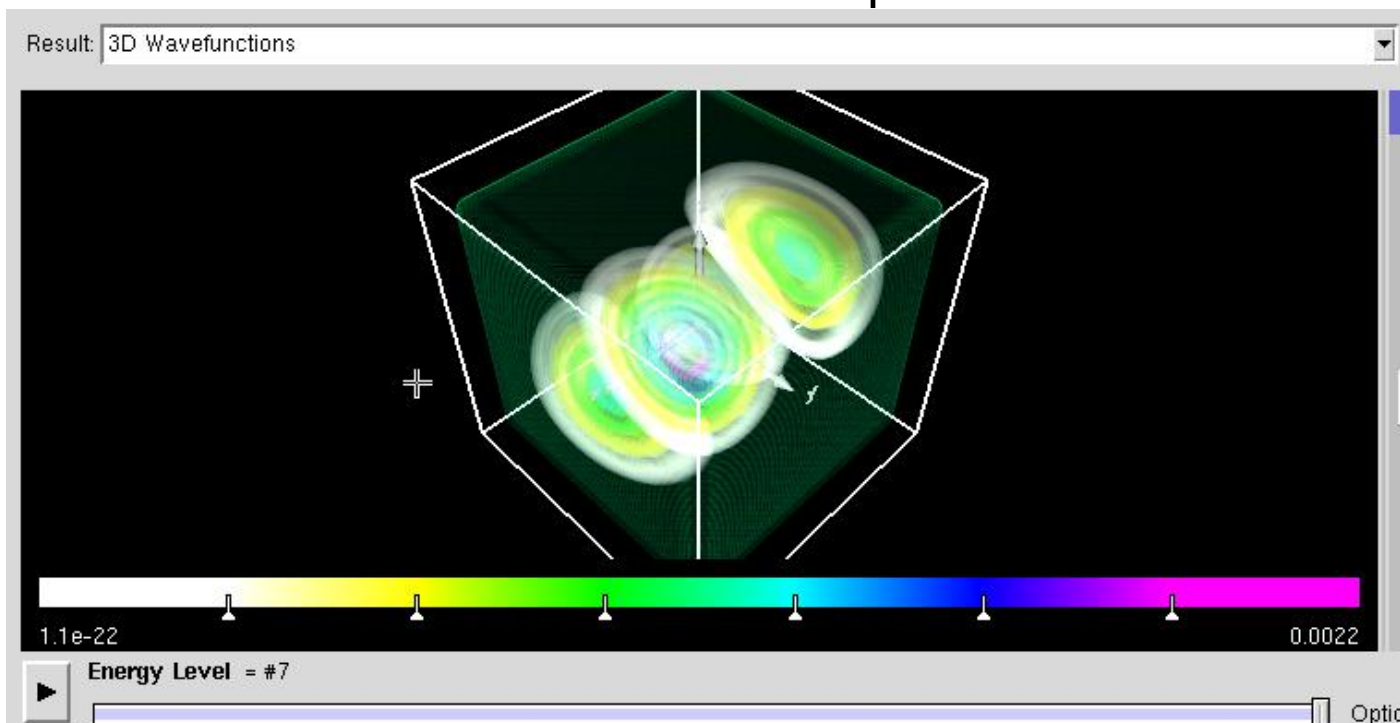
**Simulate >**

# Input Interface

- Number of states
- Device Structure
  - » Geometry
  - » Effective mass/Discretization/Energy gap
- Light Source
  - » Light polarization
  - » Optical parameters
  - » Sweep

## The Number of States

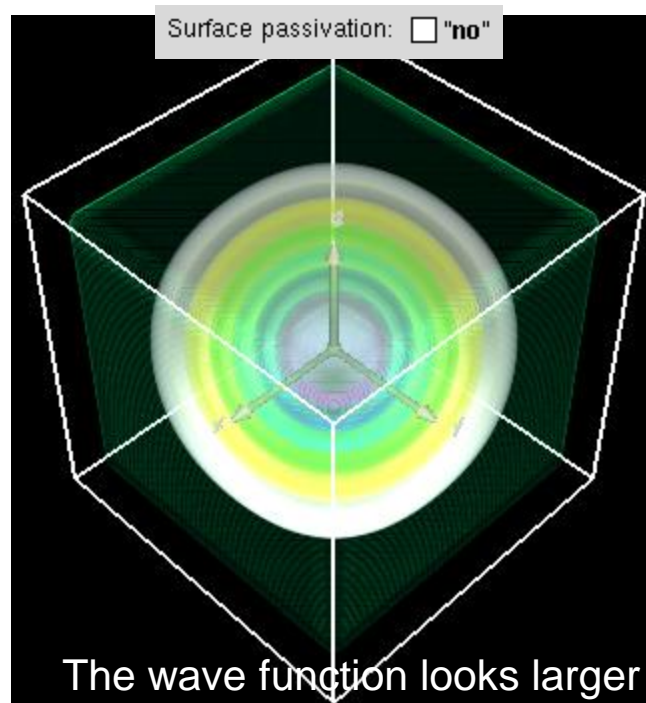
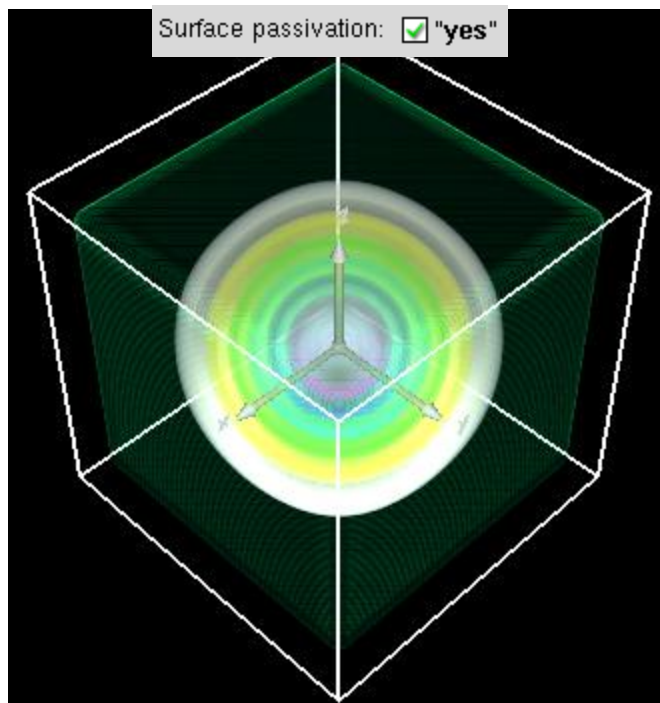
- First, choose the number of states: the default value is **Number of States: 7**
- How many states do you want to see in the output?
- Do not choose an unnecessarily large number: it increases run time.
- The output below shows that up to 7 energy levels are viewable, if number of states is chosen to be 7 in the input.






# Surface Passivation

- Surface passivation option passivates the surface so that the electron feels an infinite potential barrier at the surface of quantum dot.
  - » Surface passivation forces the electron wave function at the surface of the quantum dot to go to zero.
  - » If “no” is chosen, then the wavefunction is allowed to leak out of the quantum dot. The result is illustrated in the following figures:



# Device Structure: Geometry

- The geometry can be set by choosing x, y, and z dimensions for each of the configurations shown below.

Geometry:  

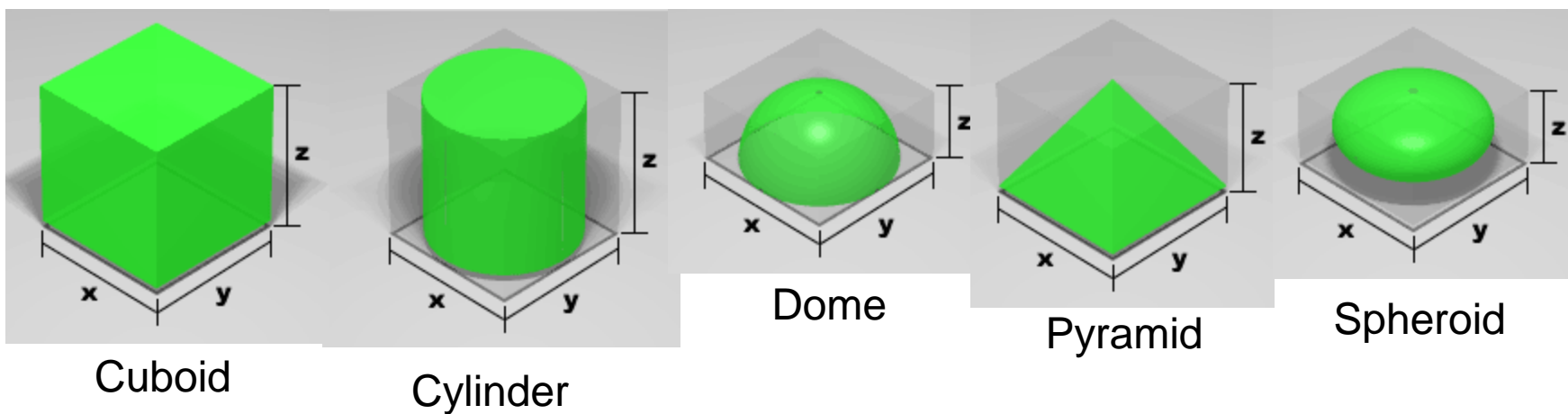
- Cuboid
- Cylinder
- Dome
- Pyramid
- Spheroid

X dimensions:

Y dimensions:

Z dimensions:

Click to expand





# Other Device Structure Parameters

Click to expand

Effective Mass: 0.067

- InAs
- GaAs**
- Ge
- Si (heavy mass)
- Si (light mass)
- Free Space

• Effective mass

- » Ratio to the free electron mass ( $m_0$ )
- » e.g., 0.067 means  $m = 0.067 \times m_0$

Discretization: 0.565nm

- InAs
- GaAs**
- Ge
- Si

• Discretization

- » The discrete mesh spacing in the quantum dot domain (simple cuboid for all shapes of quantum dot)

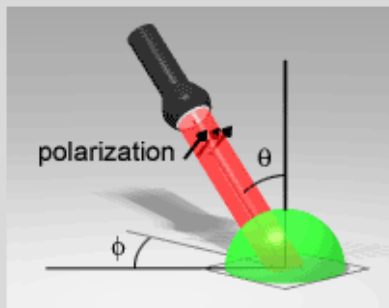
Energy gap: 1.43eV

- InAs
- GaAs**
- Ge
- Si

• Energy gap

- » The energy gap between the valance and the conduction band edge

# Light Source: Polarization/Optical parameters



## Light Polarization

Angle theta: **45deg**

Angle phi: **0deg**

## Optical Parameters

Electron Fermi level: **0eV**

Click to expand

Temperature:  **300K**

**300K (room temperature)**

**77K (liquid nitrogen)**

**4.2K (liquid helium)**

State broadening: **0.01**

## Sweep

Sweep parameter: Angle theta in units of 'degree'

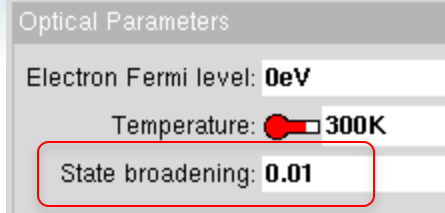
Minimum: **0**

Maximum: **90**

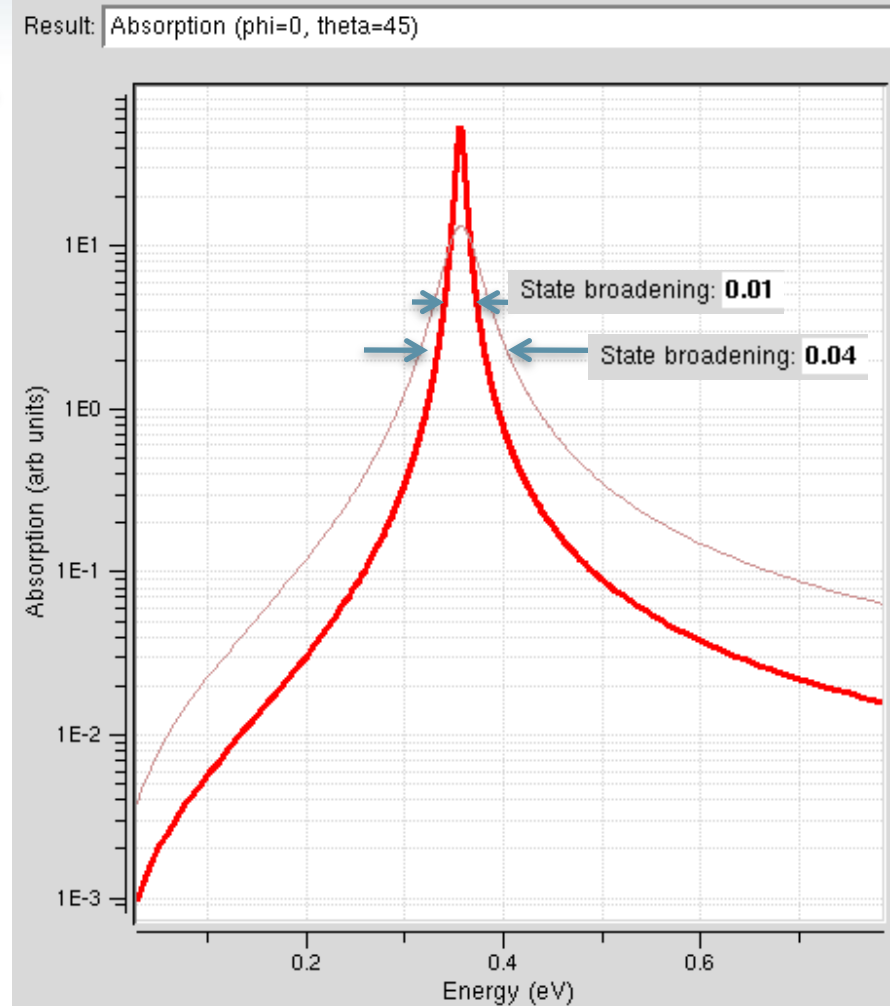
Number of points: **3**

- The light source is shined on the quantum dot to see the optical properties.
- Users can choose the angles theta  $\theta$  or phi  $\Phi$  as shown in the figure to the top left.
- Fermi level: relative to the lowest energy level
- Temperature: ambient temperature
- Detailed description: <https://www.nanohub.org/resources/4194>

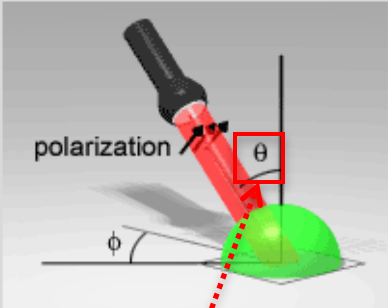
# Light Source: State Broadening



- State broadening determines
  - » the broadening width of the energy states in the quantum dot
  - » the width of the Lorentzian shape of optical absorption



# Light Source: Sweep



**Light Polarization**

Angle theta: **45deg**

Angle phi: **0deg**

**Optical Parameters**

Electron Fermi level: **0eV**

Temperature: **300K**

State broadening: **0.01**

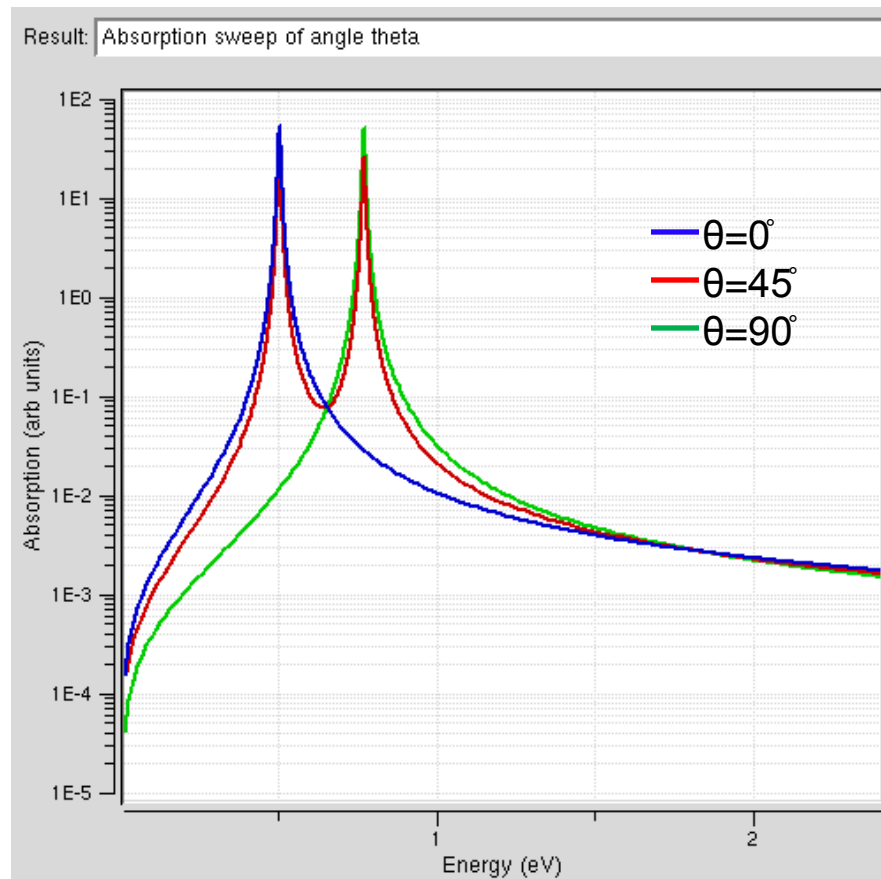
**Sweep**

Sweep parameter: **Angle theta** in units of 'degree'

Minimum: **0**

Maximum: **90**

Number of points: **3**



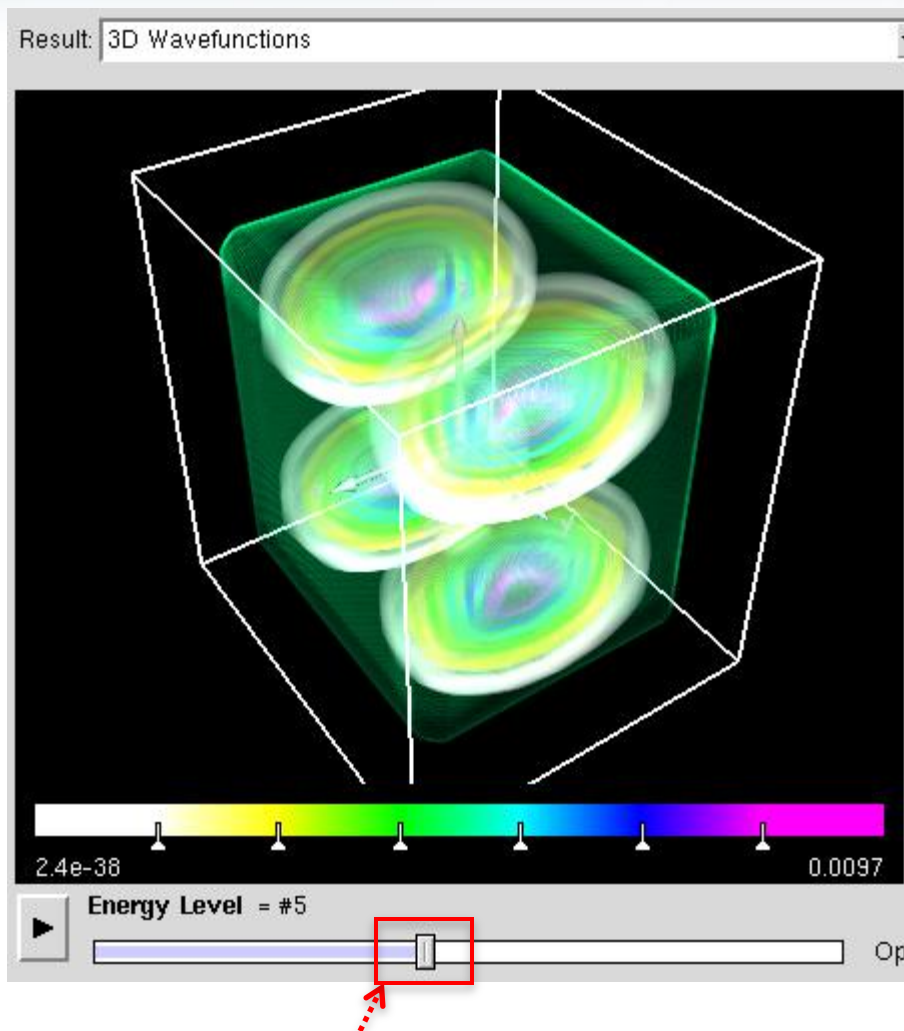
## Output Interface

- 3D wavefunctions : 3D plot of the electron wavefunction in a quantum dot
- Energy states : the energy levels of the electron in a quantum dot
- Light and dark transitions: the transition strength of electrons when light shines to a quantum dot
  - » X-polarized: when X-polarized light is shined
  - » Y-polarized: when Y-polarized light is shined
  - » Z-polarized: when Z-polarized light is shined
- Absorption: the absorption strength
- Absorption sweep: the absorption strength plot when the angles  $\theta, \phi$ , Fermi level, or temperature is swept.
- Integrated absorption: the integrated (the area under the graph of) absorption for each sweeping variable.

### Optical Properties

Refer to the introductory tutorial for more examples of the optical properties <https://www.nanohub.org/resources/4194>

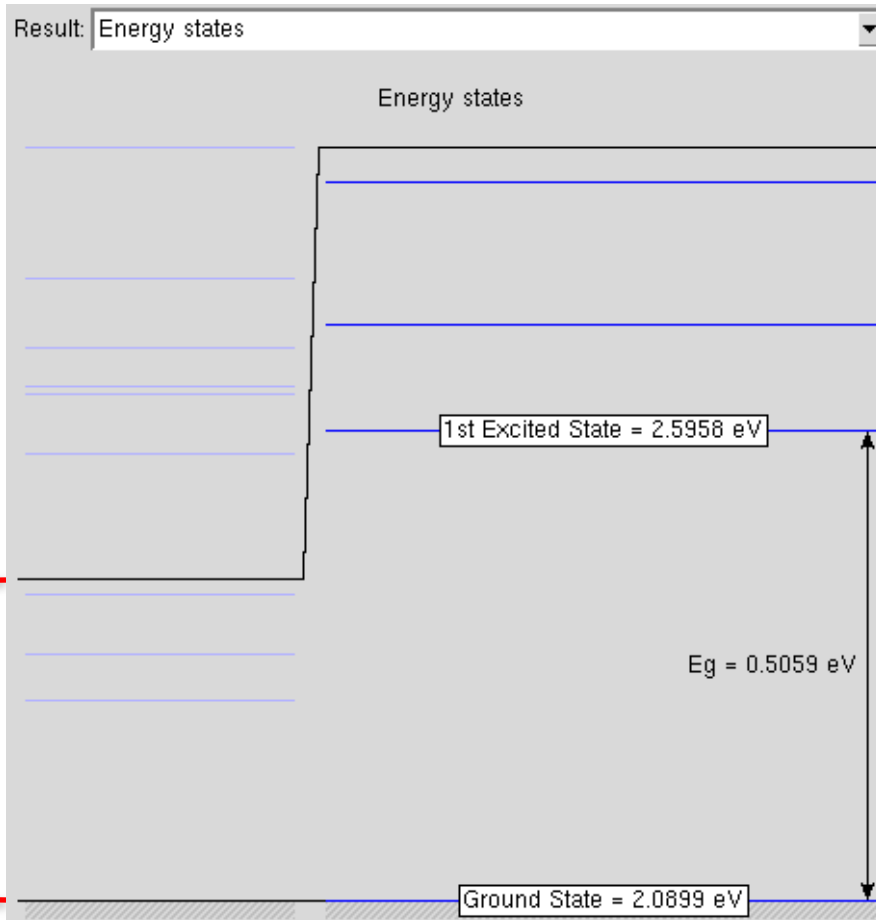
# 3D wavefunctions



Use this tab to explore different energy states



# Energy States



Total energy range

Energy range selected

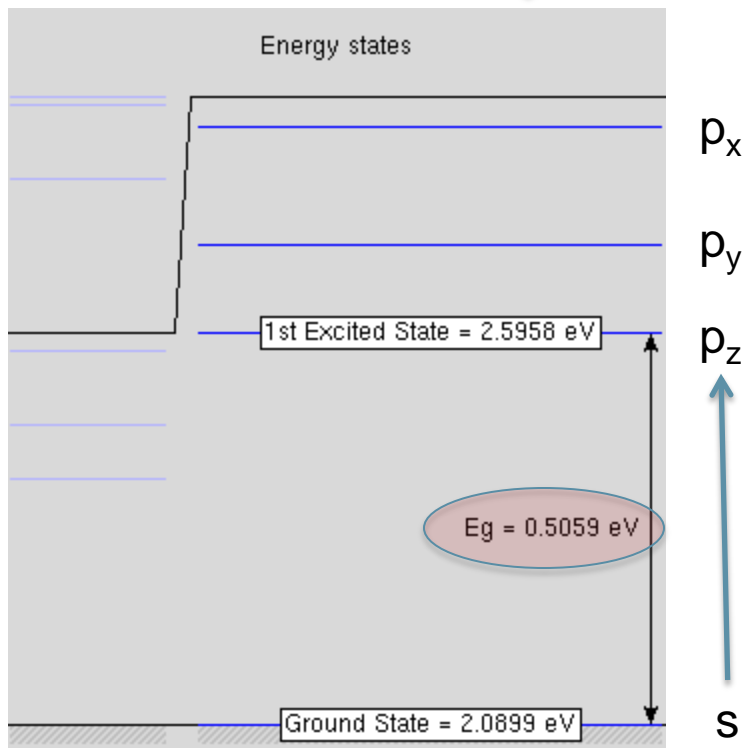
States Order	Notations in Qdot Lab
1 <sup>st</sup> state	Ground state
2 <sup>nd</sup> state	1 <sup>st</sup> excited state
3 <sup>rd</sup> state	2 <sup>nd</sup> excited state
...	...

Magnified view of the selected portion

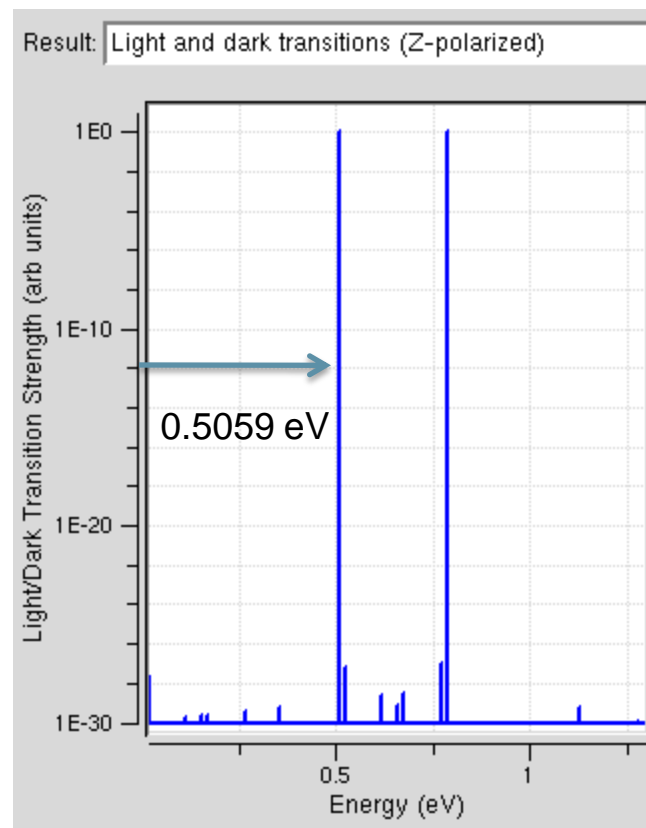
# Optical Properties: Transition Strength

Geometry: Cuboid  
X dimensions: 5nm  
Y dimensions: 5.5nm  
Z dimensions: 6nm

From the geometry, expect that the  $p_z$ -type orbital has the lowest energy. (Inverse order to the real space dimensions.)

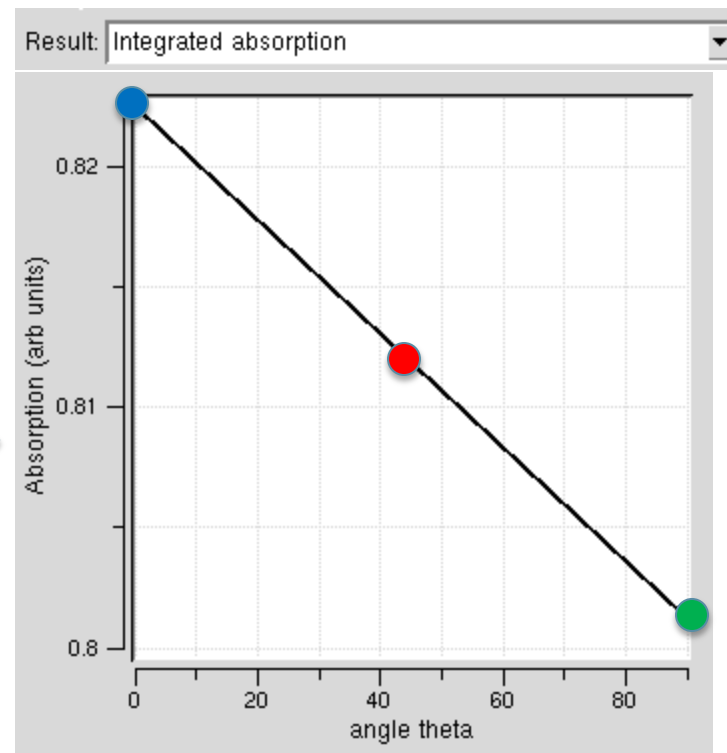
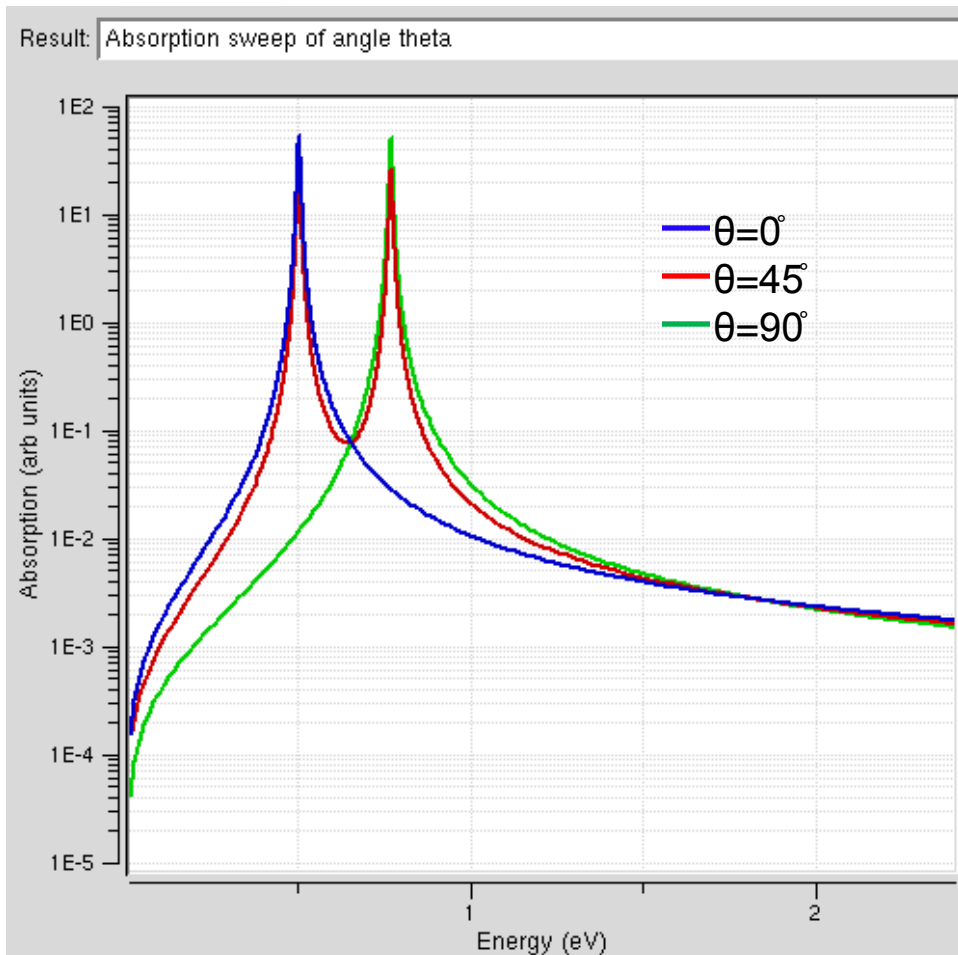


s to  $p_z$  orbital transition can be observed by Z-polarized light



The first large transition comes exactly at the transition energy from s to  $p_z$  type orbital.

# Optical Properties : Absorption



Each point is calculated by integrating each absorption graph (note that the color of each point matches the color of the line in the left figure)

## Simulation Engine behind the tool: NEMO 5

- Right now, the quantum dot lab's engine is NEMO 5.
- NEMO 5 is a **N**ano **E**lectronic **M**Odeling tool.<sup>[2]</sup>
- Quantum dot lab tool mainly uses the following parts of NEMO5
  - » Structure construction
  - » Schrodinger solver
  - » Optical properties solver

[2] [https://engineering.purdue.edu/gekcogrp/research-group/SebastianSteiger/quad\\_NEMO5.pdf](https://engineering.purdue.edu/gekcogrp/research-group/SebastianSteiger/quad_NEMO5.pdf)

## References

- [1] Gerhard Klimeck, Introduction to Quantum Dot Lab:  
<https://www.nanohub.org/resources/4194>
- [2] Sebastian Steiger, NEMO 5 quad chart:  
[https://engineering.purdue.edu/gekcogrp/research-group/SebastianSteiger/quad\\_NEMO5.pdf](https://engineering.purdue.edu/gekcogrp/research-group/SebastianSteiger/quad_NEMO5.pdf)