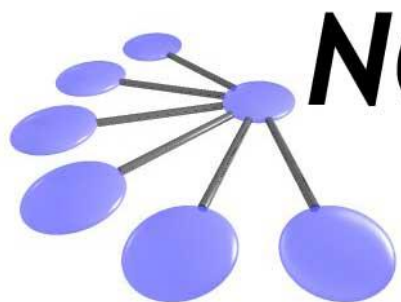
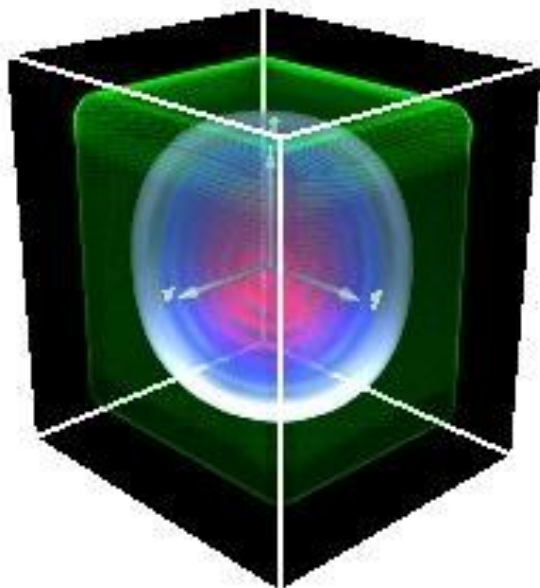


Network for Computational Nanotechnology (NCN)

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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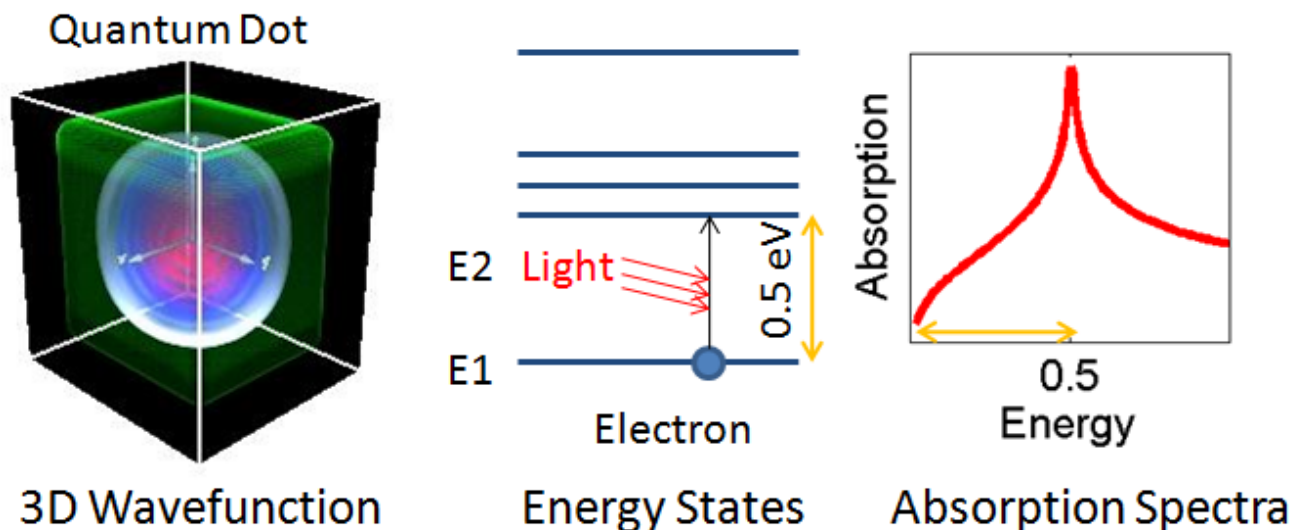
*<http://www.nanohub.org/tools/qdot/>.

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- References 19

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: 7

Surface passivation: "yes"

Device Structure

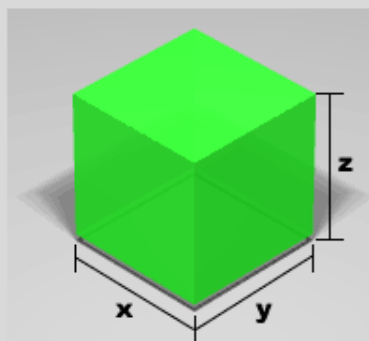
Light Source

Geometry: Cuboid

X dimensions: 10nm

Y dimensions: 10nm

Z dimensions: 10nm



Effective Mass: 0.067

Discretization: 0.565nm

Energy gap: 1.43eV

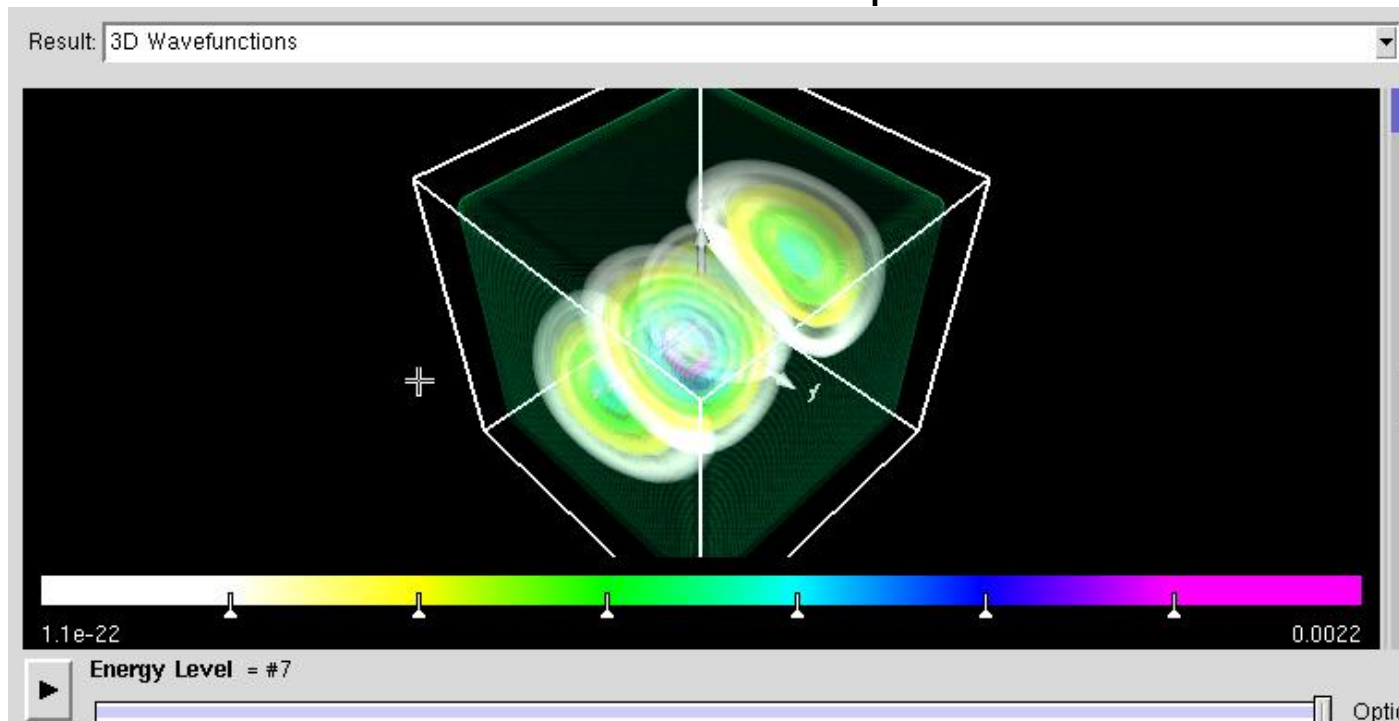
Simulate >

Input Interface

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

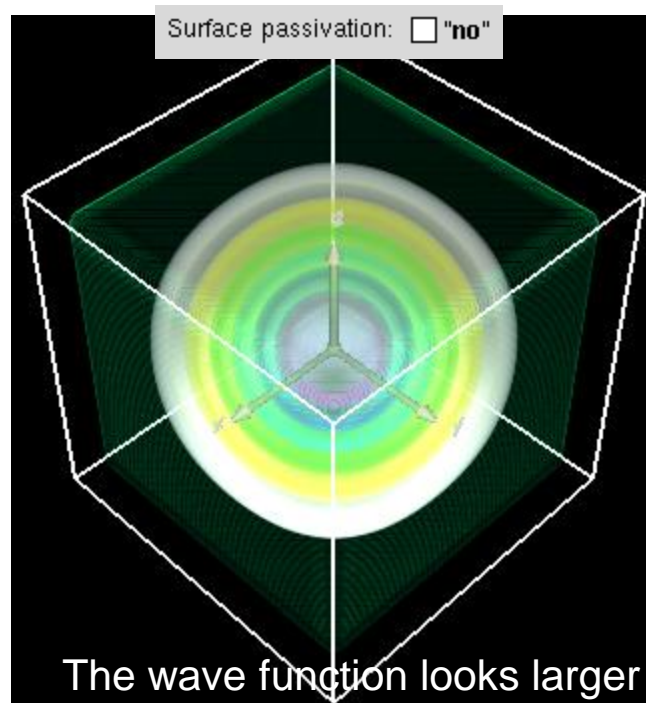
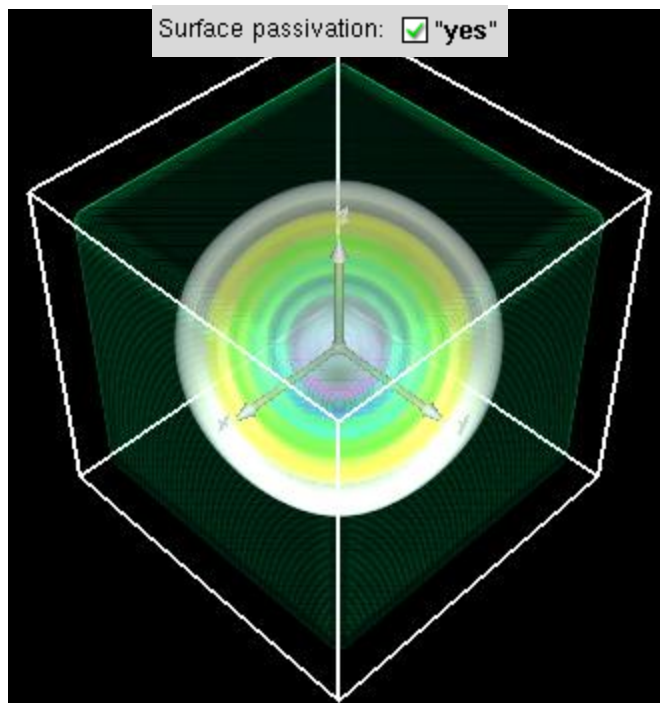
The Number of States

- 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:



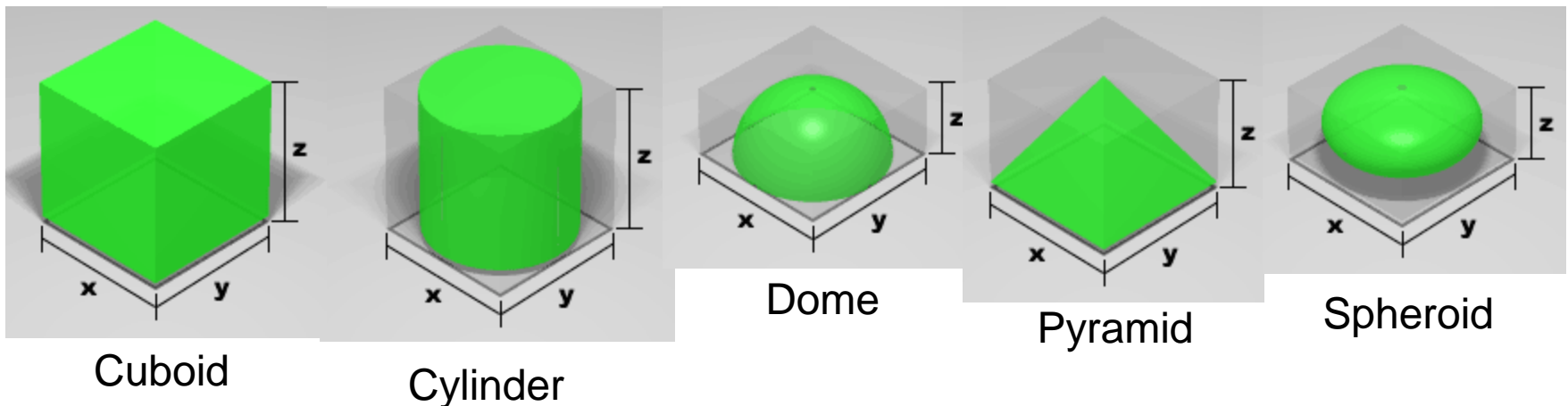
Device Structure: Geometry

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

Geometry:	Cuboid	
	Cuboid	
	Cylinder	
	Dome	
	Pyramid	
	Spheroid	

Click to expand

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



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)

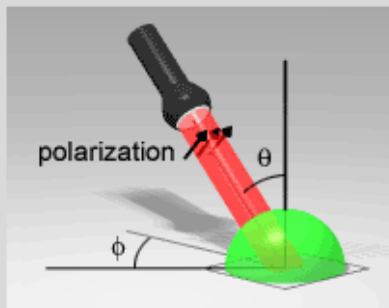
Energy gap: 1.43eV

- 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 θ or phi Φ 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

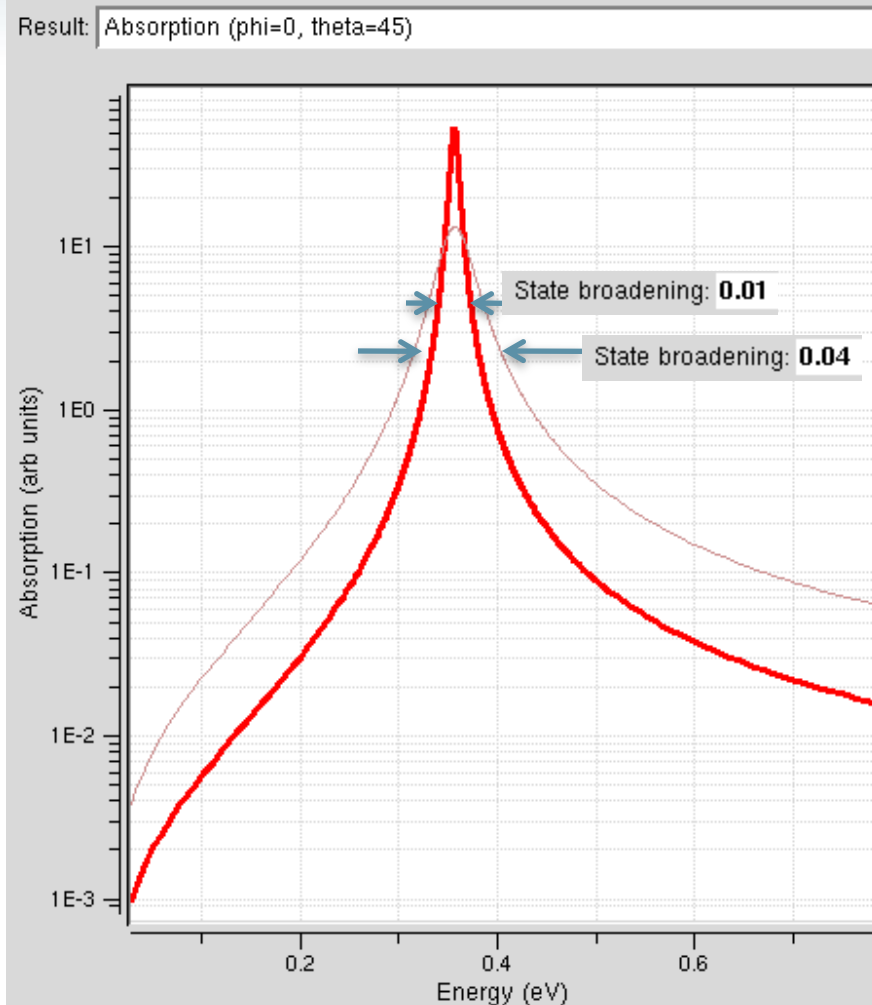
Optical Parameters

Electron Fermi level: 0eV

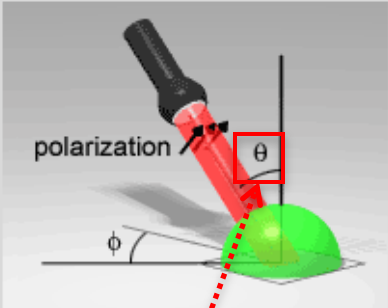
Temperature: 300K

State broadening: 0.01

- 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**

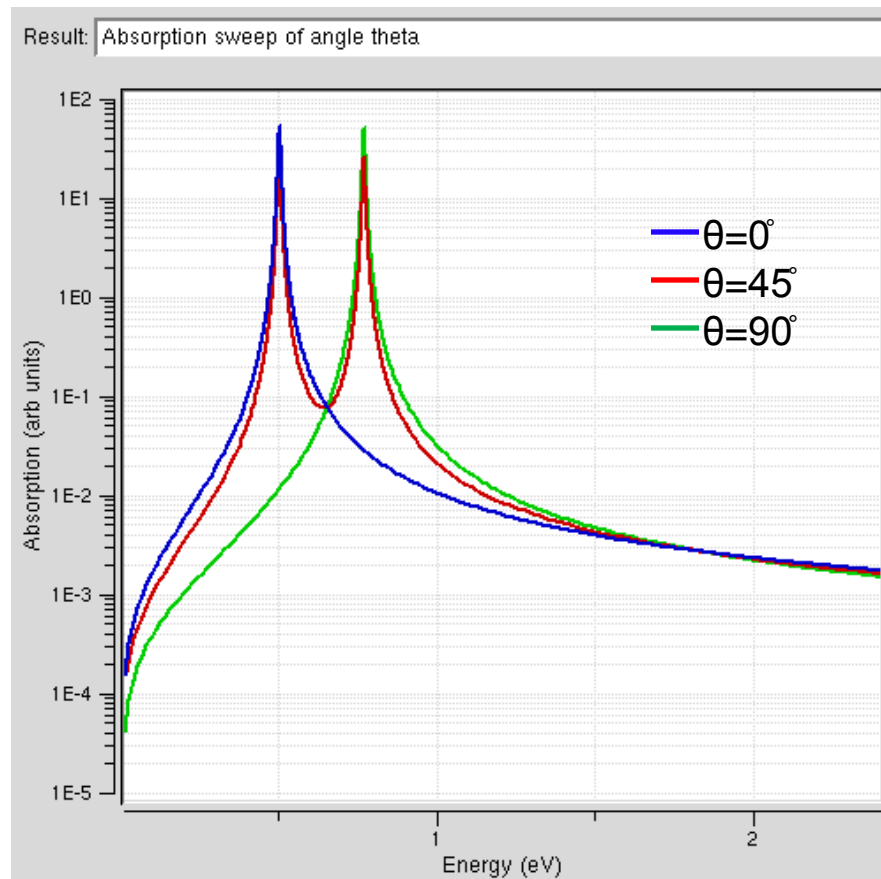
Sweep

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

Minimum: **0**

Maximum: **90**

Number of points: **3**



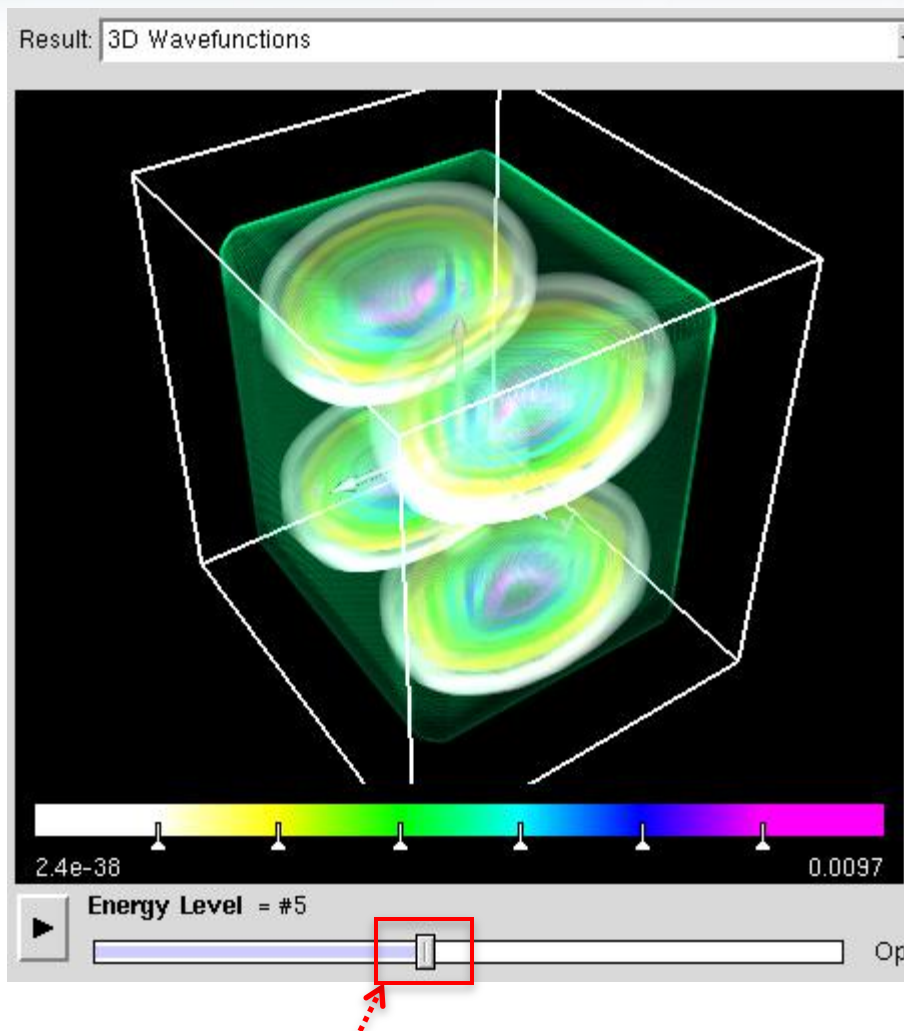
Output Interface

- 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 θ, ϕ , Fermi level, or temperature is swept.
- Integrated absorption: the integrated (the area under the graph of) absorption for each sweeping variable.

Optical Properties

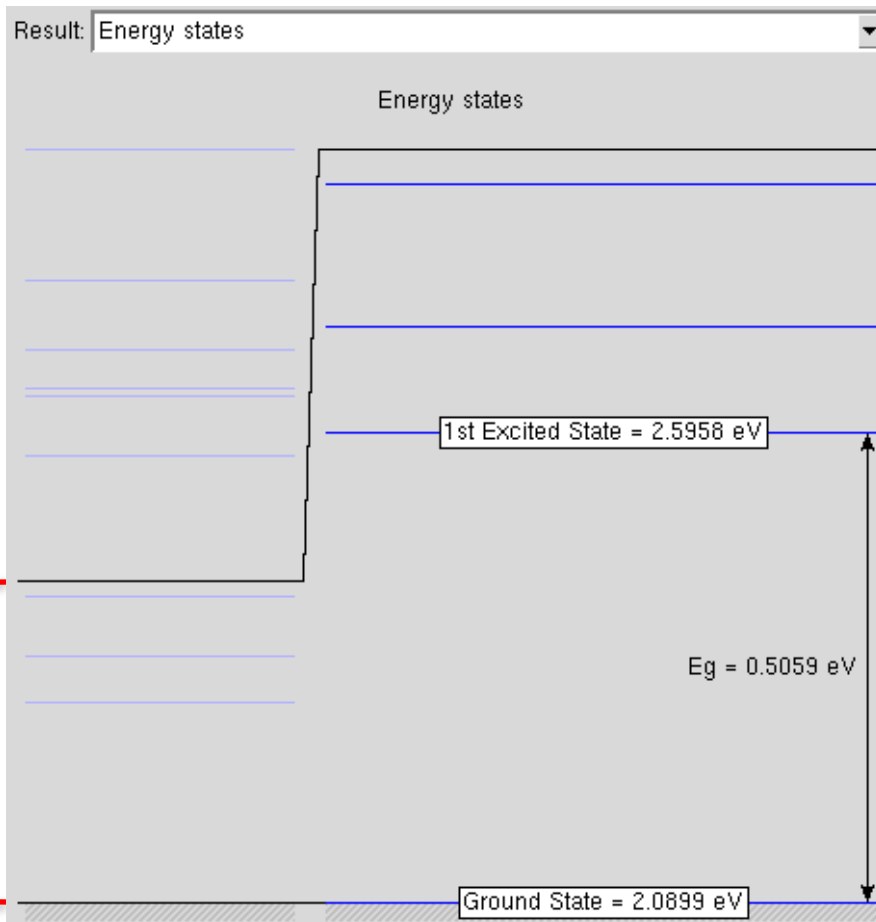
Refer to the introductory tutorial for more examples of the optical properties <https://www.nanohub.org/resources/4194>

3D wavefunctions



Use this tab to explore different energy states

Energy States



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

Magnified view of the selected portion

Total energy range

Energy range selected

Optical Properties: Transition Strength

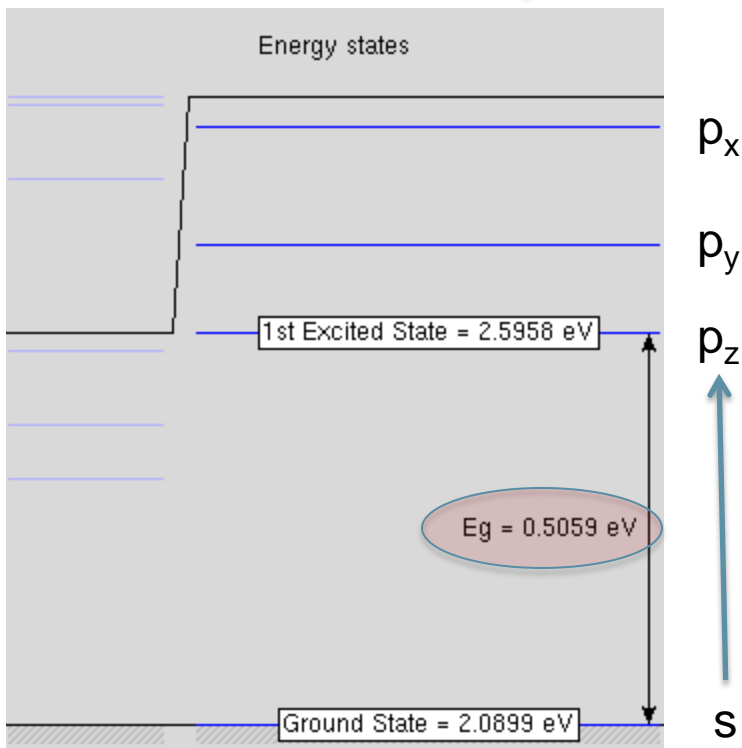
Geometry: Cuboid

X dimensions: 5nm

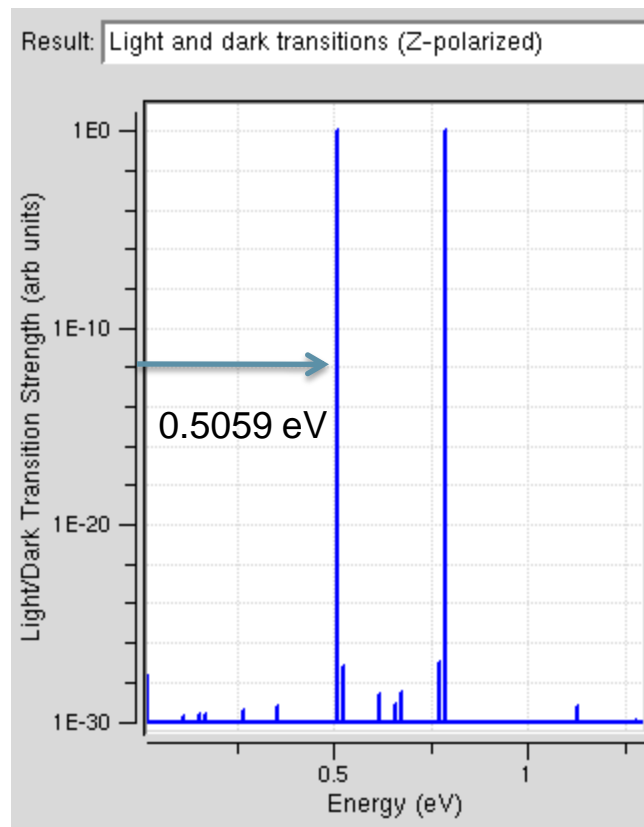
Y dimensions: 5.5nm

Z dimensions: 6nm

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

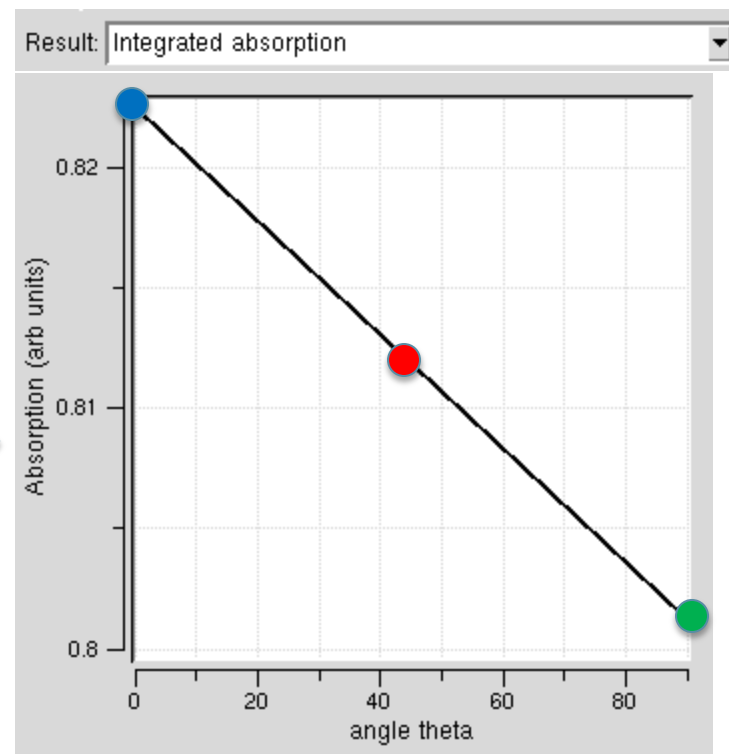
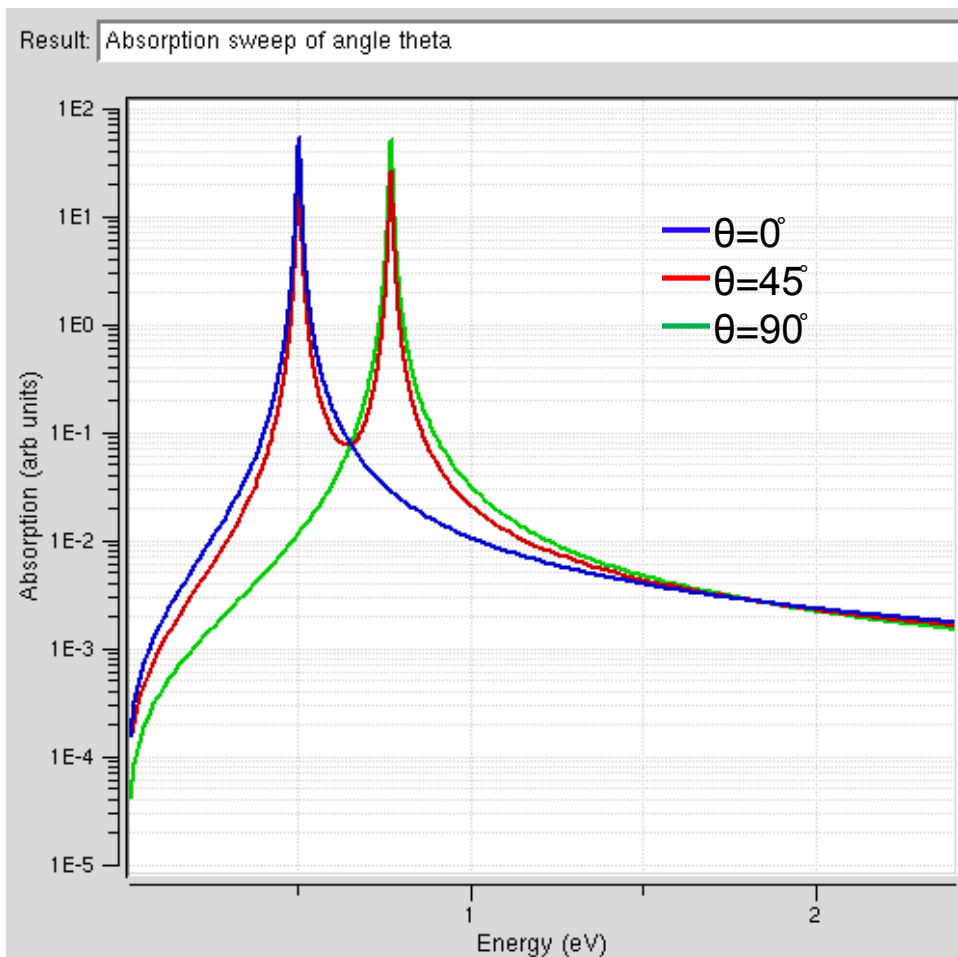


s to p_z orbital transition can be observed by Z-polarized light



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.^[2]
- 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

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