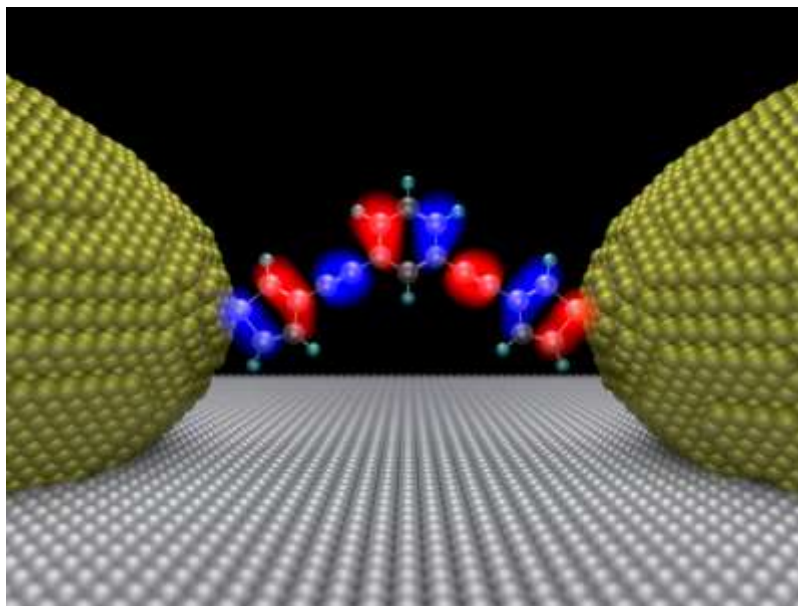


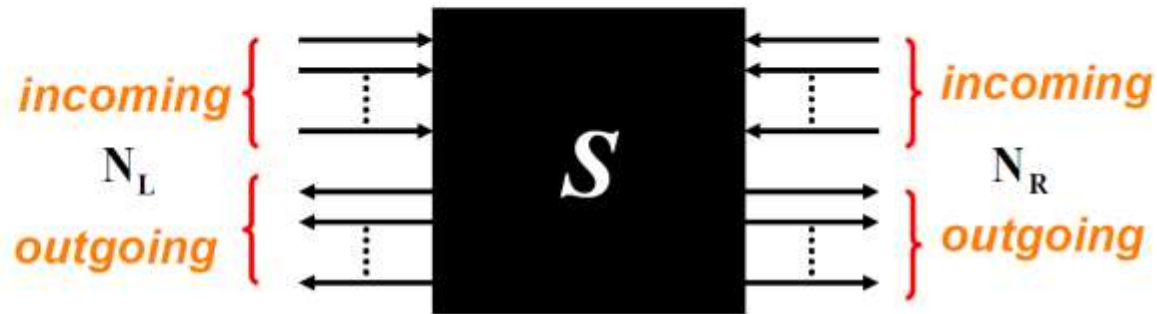
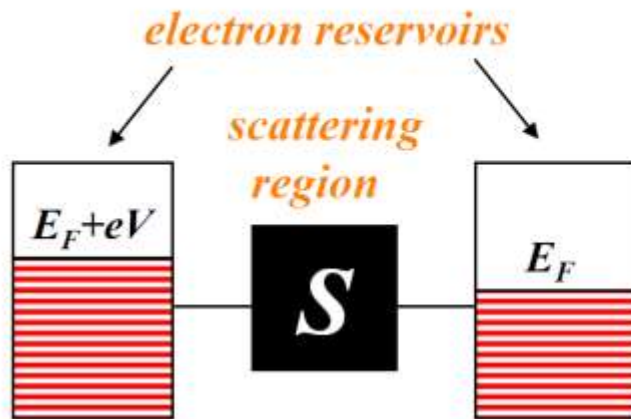
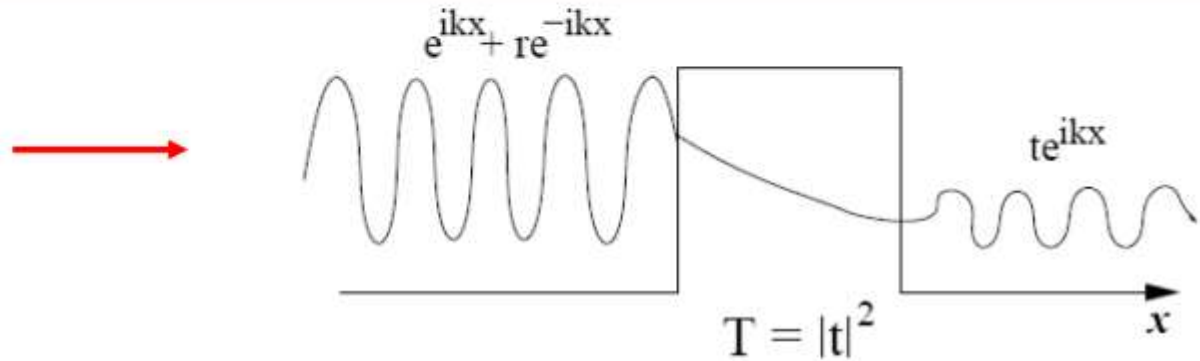
Molecular and carbon-based electronic systems

Lecture 7 - Insights in Density Functional Theory for molecular junctions



Landauer approach to electron transport

Electronic transport as scattering problem



$$G = \frac{2e^2}{h} T(E_F)$$

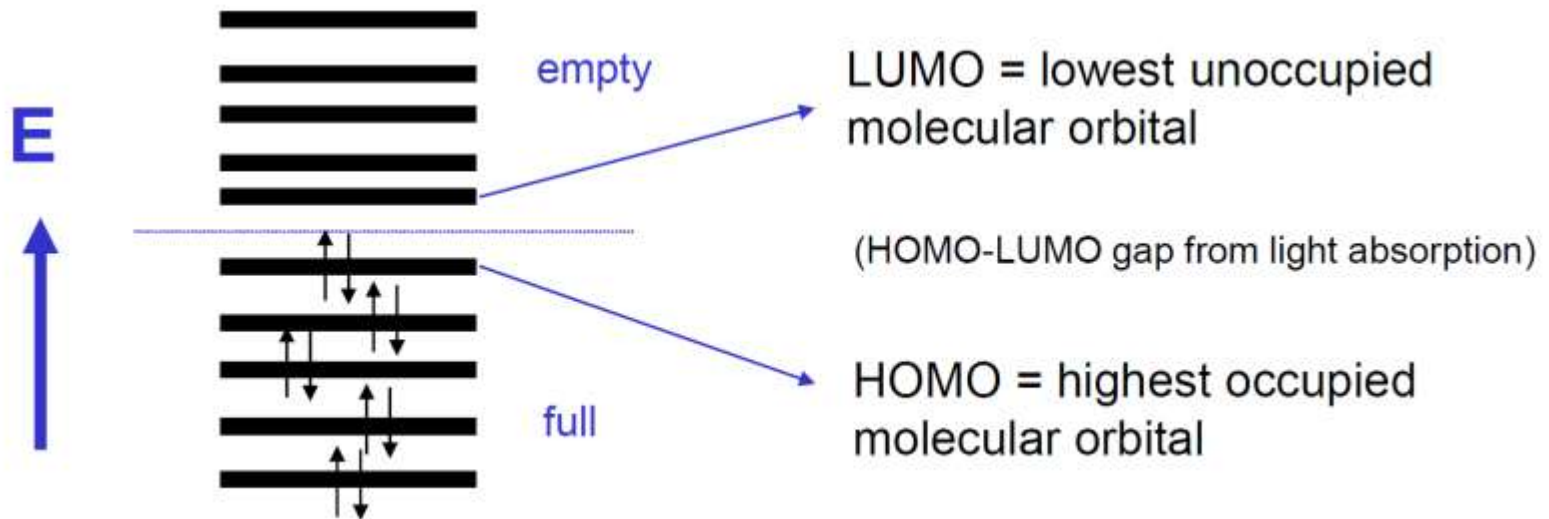
G = conductance

$T(E_F)$ = transmission at the Fermi energy

molecular energy levels

(isolated) molecule

quantum system with spectrum of discrete energy states (molecular orbitals)



'Particle in a box': molecule is small box

box smaller \Leftrightarrow levels more spaced ($\delta E > k_B T$)

simple one-level model

Analytical expression for the current

$$I(E, V) = \frac{2e}{h} \int_{-\infty}^{\infty} T(E, V) [f(E - \mu_2) - f(E - \mu_1)] dE$$

At low T , Fermi functions \approx Heaviside step functions, and

$$I(V) = \frac{2e}{h} \int_{-\frac{eV}{2}}^{\frac{eV}{2}} T(E, V) dE$$

Using the expressions

$$T(E, V) = \frac{4\Gamma_1\Gamma_2}{(E - E_0(V))^2 + (\Gamma_1 + \Gamma_2)^2}$$

$$E_F(V) = \frac{eV}{2} \cdot \frac{\Gamma_1 - \Gamma_2}{\Gamma_1 + \Gamma_2}$$

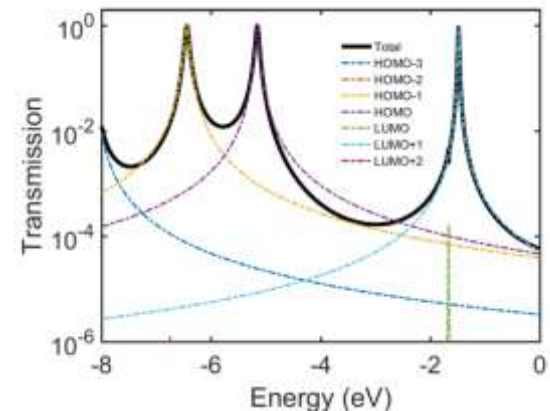
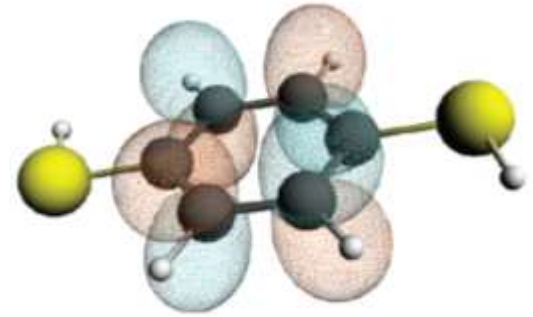
we can write

$$I(V) = \frac{8e}{h} \cdot \frac{\Gamma_L\Gamma_R}{\Gamma_L + \Gamma_R} \left[\arctan \left(\frac{2E_0 + eV \left(\frac{\Gamma_L - \Gamma_R}{\Gamma_L + \Gamma_R} + 1 \right)}{2(\Gamma_L + \Gamma_R)} \right) - \arctan \left(\frac{2E_0 + eV \left(\frac{\Gamma_L - \Gamma_R}{\Gamma_L + \Gamma_R} - 1 \right)}{2(\Gamma_L + \Gamma_R)} \right) \right]$$

3 fit parameters $\Gamma_1, \Gamma_2, E_0 = E_0(V)$

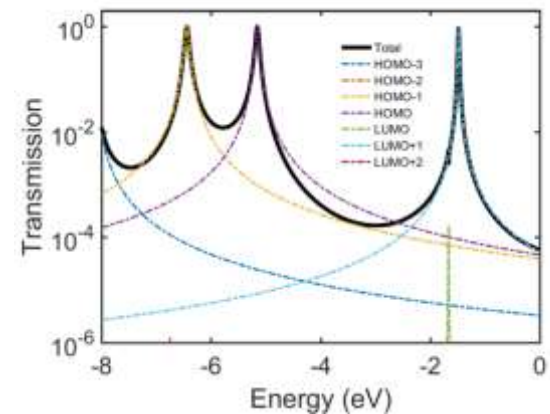
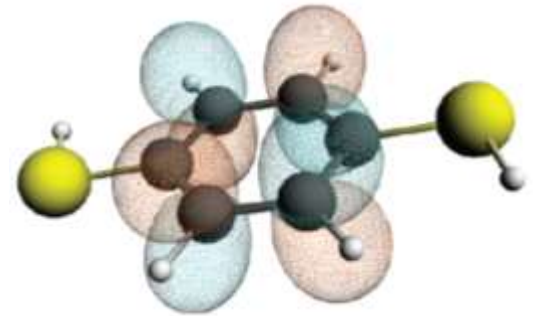
Overview

- Quantum chemistry (QC)
- Density functional theory (DFT)
- Non-equilibrium Green's function (NEGF)



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- Density functional theory (DFT)
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Schrödinger's Equation

- Molecule can be described using the Schrödinger Equation

Time-independent Schrödinger equation (*general*)

$$\hat{H} \Psi = E \Psi$$

Schrödinger's Equation

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Time-independent Schrödinger equation (*general*)

$$\hat{H} \Psi = E \Psi$$

$$\hat{H} = -\frac{\hbar^2}{2m_e} \sum_i \nabla_i^2 + \frac{1}{2} \sum_{i \neq j} \frac{e^2}{|\mathbf{r}_i - \mathbf{r}_j|} - \sum_{i,J} \frac{Z_J e^2}{|\mathbf{r}_i - \mathbf{R}_J|} - \sum_I \frac{\hbar^2}{2M_I} \nabla_I^2 + \frac{1}{2} \sum_{I \neq J} \frac{Z_I Z_J e^2}{|\mathbf{R}_I - \mathbf{R}_J|}$$

Very Complex many body Problem !!
(Because everything interacts)

Born-Oppenheimer approximation

- Electrons are much lighter, and faster
- Decoupling in the wave function

$$\Psi(\vec{r}, \vec{R}) \approx \Psi_e(\vec{r})\Psi_n(\vec{R})$$

- Nuclei described as constant external potential

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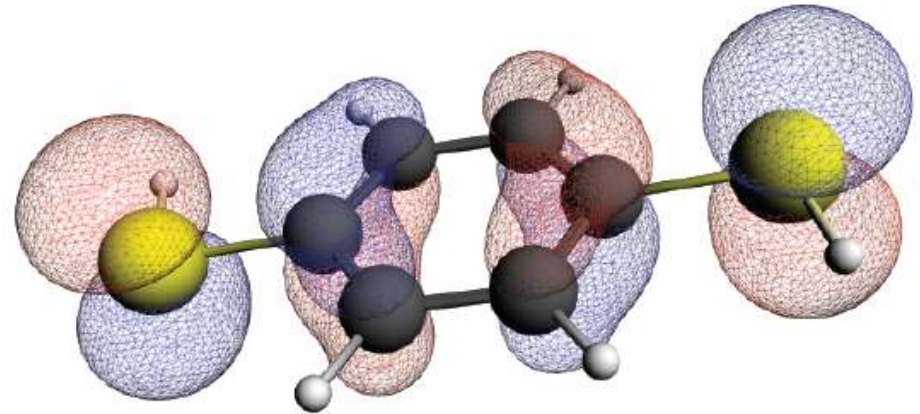
- Nuclei described as constant external potential
- Still numerically intractable

$$\hat{H} = -\frac{\hbar^2}{2m_e} \sum_i \nabla_i^2 + \frac{1}{2} \sum_{i \neq j} \frac{e^2}{|\mathbf{r}_i - \mathbf{r}_j|} - \sum_{i,J} \frac{Z_J e^2}{|\mathbf{r}_i - \mathbf{R}_J|}$$

Quantum chemistry calculations

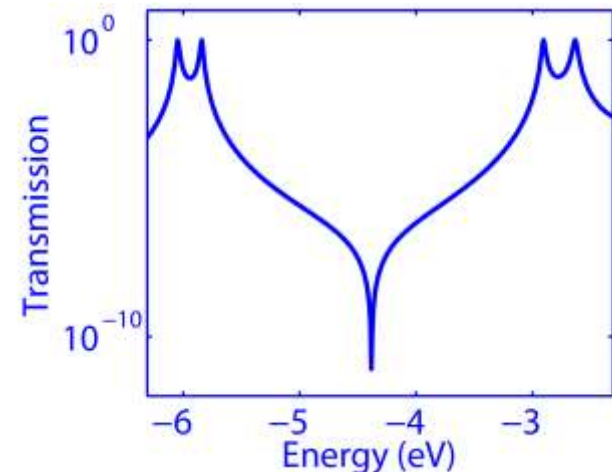
Electronic structure

- Numerical renormalization group
- Configuration interaction
- Hartree Fock
- Density functional theory
- Hückel method



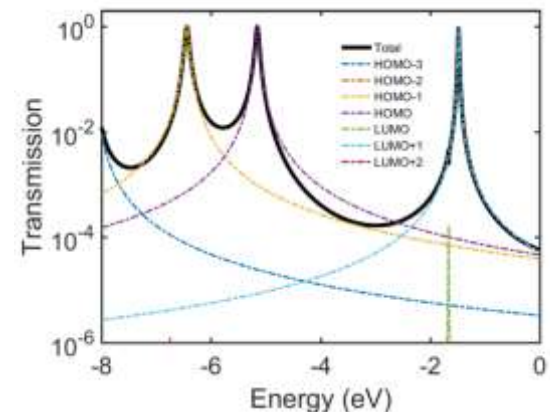
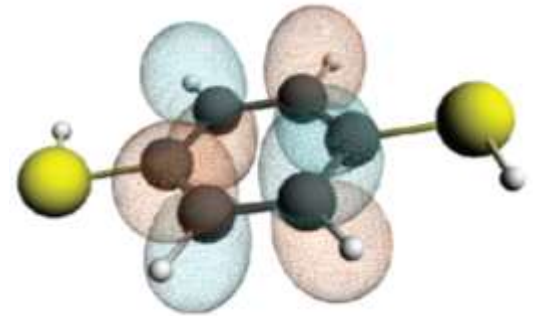
Transport

- NEGF
- GW



Overview

- Quantum chemistry (QC)
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DFT

- Density functional theory (DFT) is an **exact reformulation** of many-body quantum mechanics in terms of the **electron density** rather than the wave function
 - equivalent independent single-particle problem

DFT

- Density functional theory (DFT) is an **exact reformulation** of many-body quantum mechanics in terms of the **electron density** rather than the wave function
→ equivalent independent single-particle problem
- The wave-function is a unique functional of the density (**1^e Hohenberg-Kohn theorem, 1964**)
- All system properties are governed *only* by the ground-state density.
- Nobelprize in 1998!

DFT

- The density $n(r)$ which minimizes $E[n(r)]$ is the ground-state density, and minimization of this functional alone is enough to fully determine the exact ground-state energy and density $n_0(r)$.
(2^e Hohenberg-Kohn theorem, 1964)


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$$E_{\text{KS}} = T_s[n] + \int d^3r V_{\text{ext}}(\mathbf{r})n(\mathbf{r}) + E_H[n] + E_{\text{II}} + E_{\text{xc}}[n]$$

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$$E_{\text{KS}} = T_s[n] + \int d^3r V_{\text{ext}}(\mathbf{r})n(\mathbf{r}) + E_H[n] + E_{\text{II}} + E_{\text{xc}}[n]$$


The diagram shows five arrows pointing from labels below to terms in the equation above:

- An arrow from "Kinetic energy" points to $T_s[n]$.
- An arrow from "Electron-nuclei interaction" points to $\int d^3r V_{\text{ext}}(\mathbf{r})n(\mathbf{r})$.
- An arrow from "Electron-electron interaction" points to $E_H[n]$.
- An arrow from "Ion-Ion interaction" points to E_{II} .
- An arrow from "Exchange + correlations" points to $E_{\text{xc}}[n]$.

$$E_H[n] = \frac{1}{2} \int d^3r \int d^3r' \frac{n(\mathbf{r})n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}$$

Kohn-Sham equations

- But T_s and E_{xc} are unknown. How to find T_s ?

Kohn-Sham equations

- But T_s and E_{xc} are unknown. How to find T_s ?
- Kohn and Sham proposed to model electrons as non-interacting particles that generate the same density as the interacting particles (mean field, 1965)

$$\left(-\frac{\hbar^2}{2m} \nabla^2 + v_{\text{eff}}(\mathbf{r}) \right) \phi_i(\mathbf{r}) = \varepsilon_i \phi_i(\mathbf{r})$$

Exchange-correlation (XC)

- All we know about the functional E_{xc} is that it exists, however, its form is unknown.

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Local density approximation (LDA) : uniform electron gas: $E_{xc}[\rho]$

Generalized gradient approximation (GGA) : $E_{xc}(\rho, \nabla\rho)$

Hybrid (B3LYP) : including some HF for exchange

Dispersion/long-range corrected functionals

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Dispersion/long-range corrected functionals

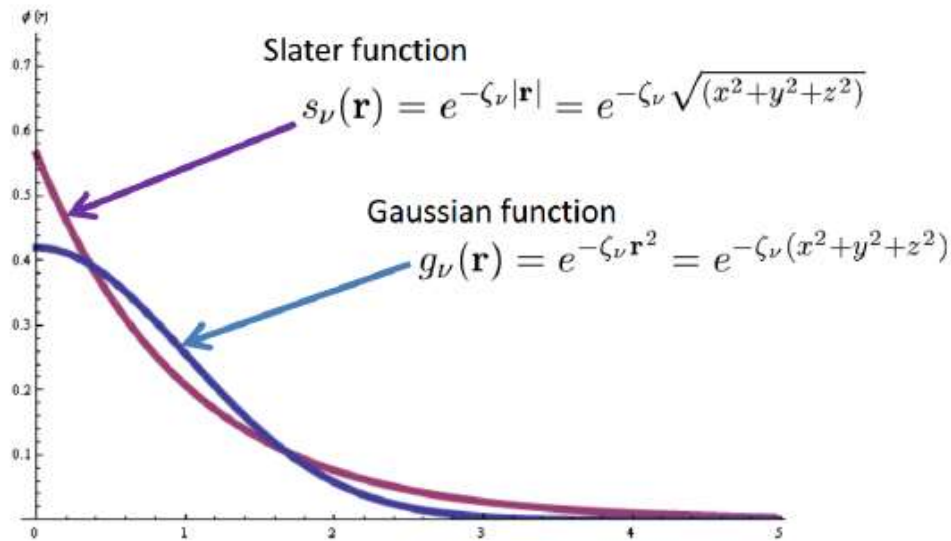
→ As the XC functional is not exact, electron exchange and correlations are not taken properly into account.

Basis set used to describe orbitals

- A **basis set** is a set of basis functions which are combined in linear combinations to create molecular orbital wave-functions.

Basis set used to describe orbitals

- A **basis set** is a set of basis functions which are combined in linear combinations to create molecular orbital wave-functions.
- Basis set can be any type of set of functions: atomic orbitals, plane waves, ...



Convergence (SCF)

Guess wavefunction

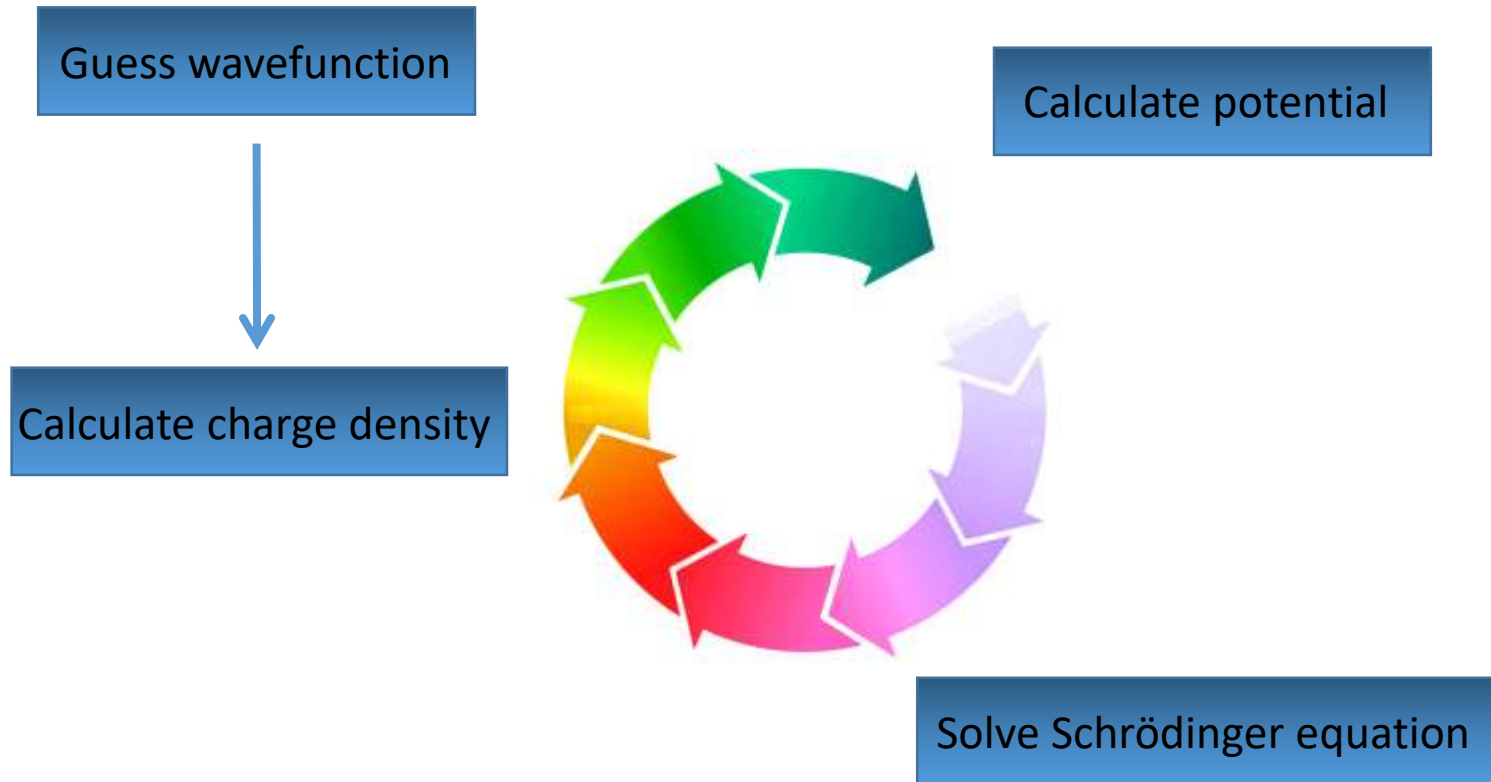
Convergence (SCF)

Guess wavefunction

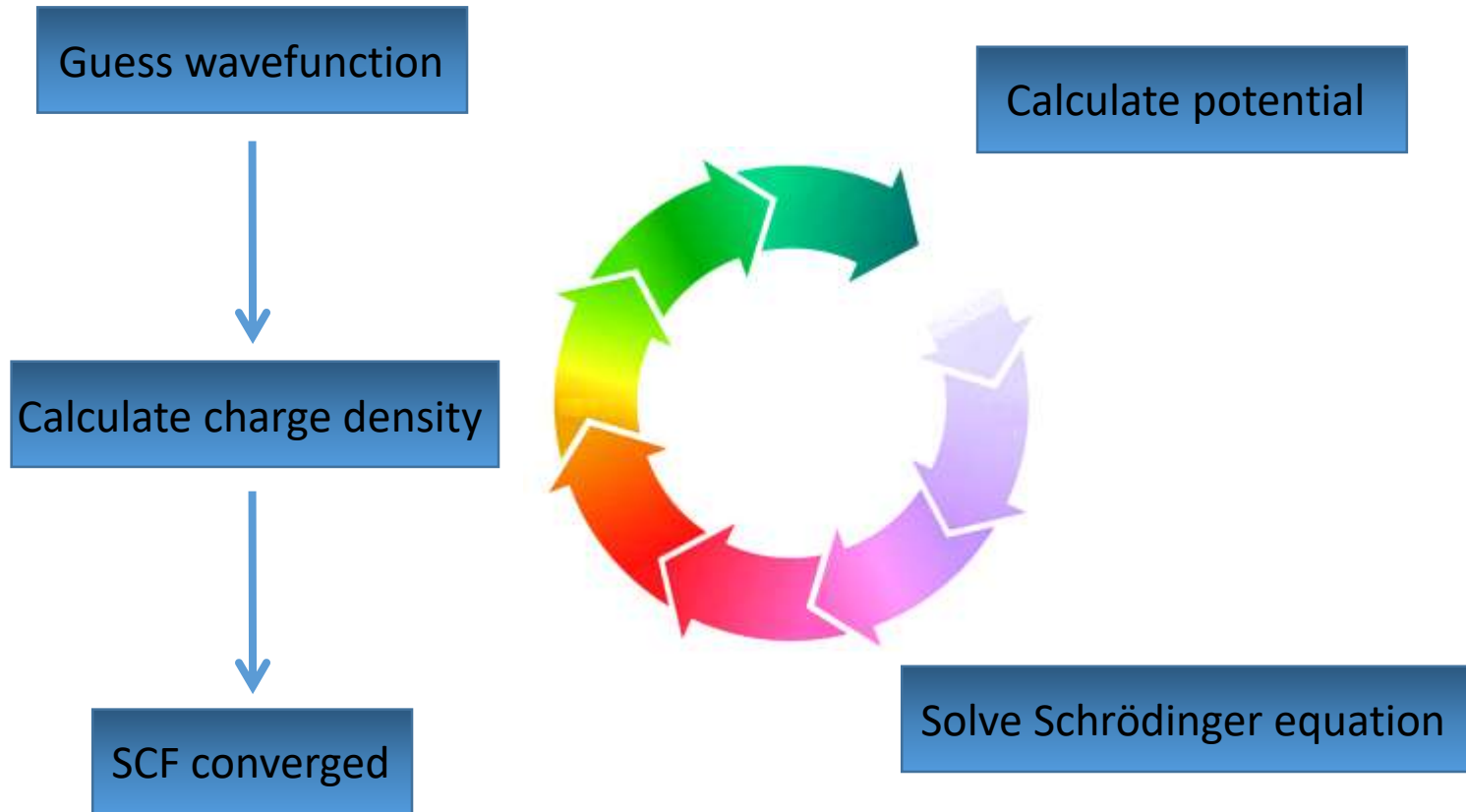


Calculate charge density

Convergence (SCF)

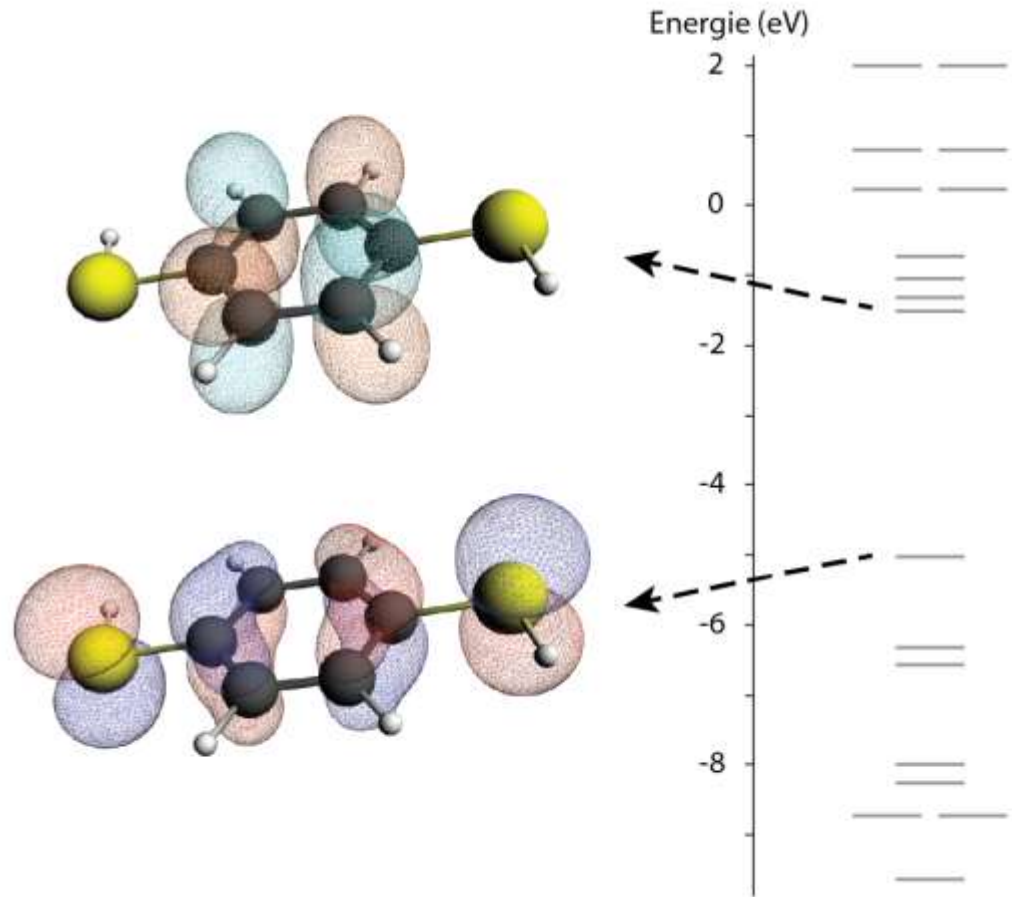


Convergence (SCF)



DFT Results

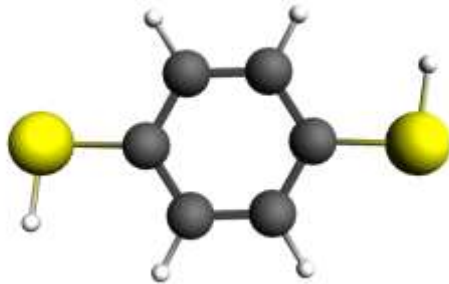
- DFT provides the energy of the molecular orbitals
- DFT provides the wave-functions of the molecular orbitals



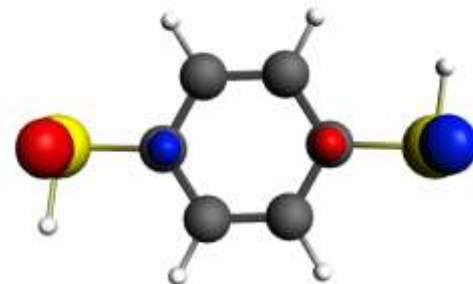
Molecular orbitals

- # electrons per cubic ångström
- Positive and negative part of the wave function

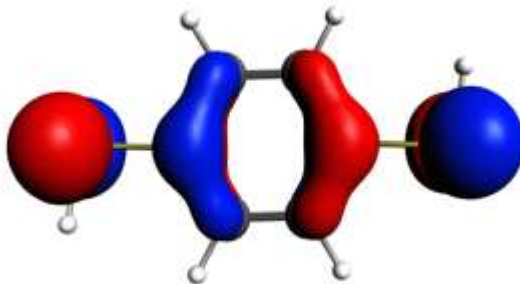
0.5



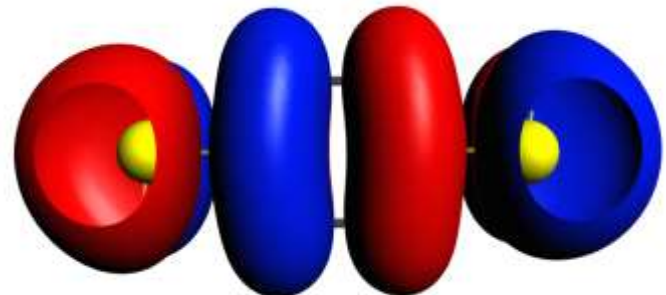
0.1



0.05

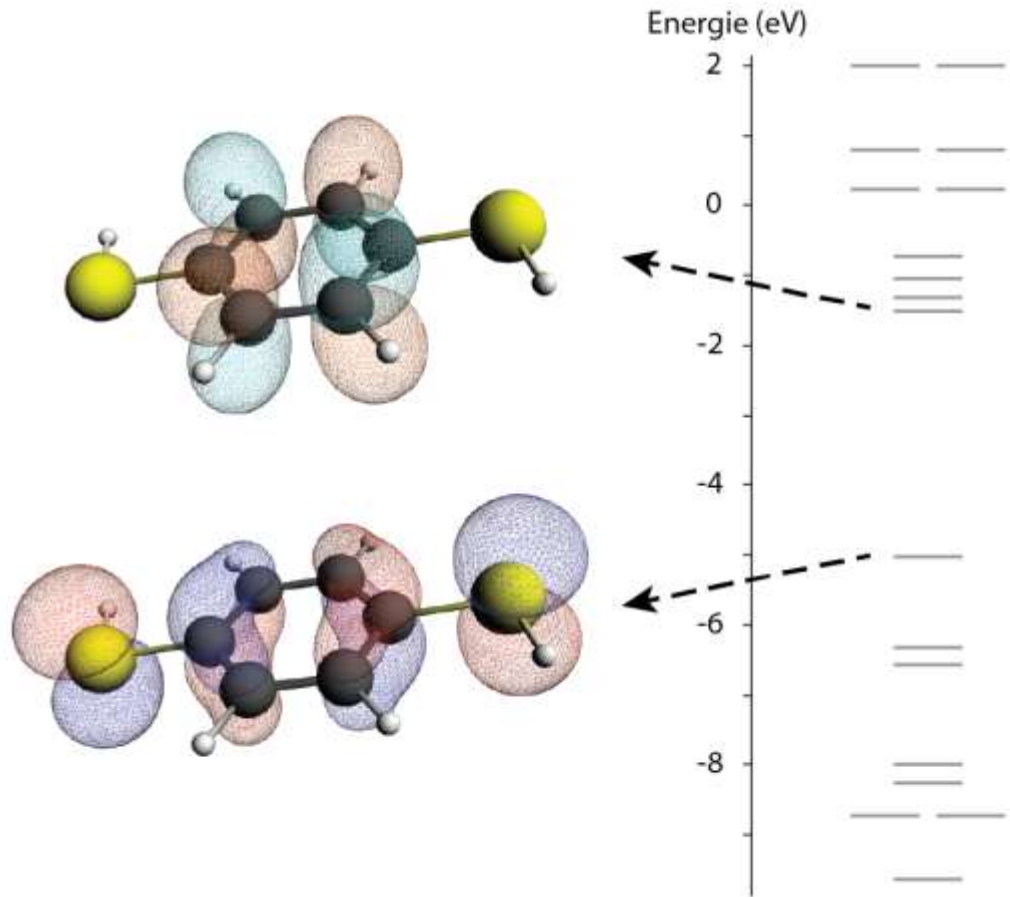


0.01



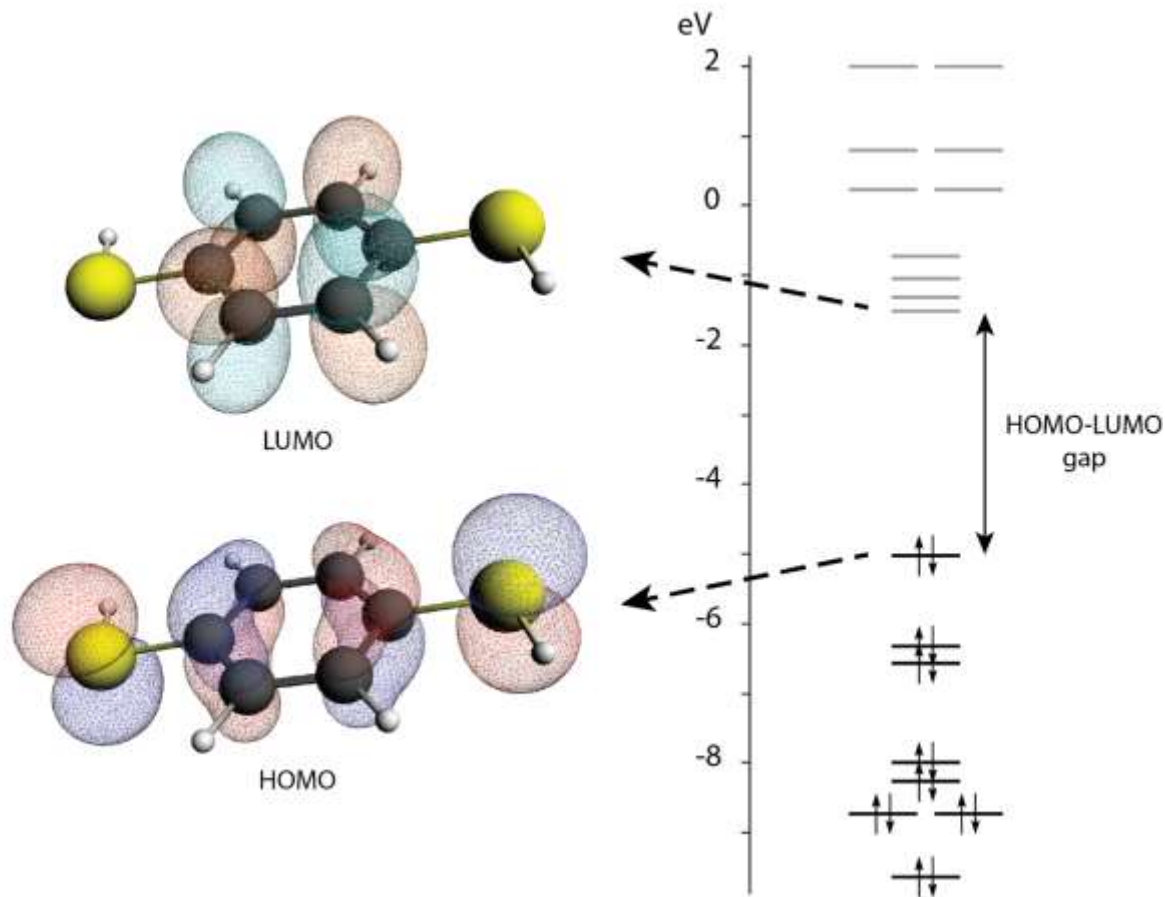
DFT Results

- DFT provides the energy of the molecular orbitals
- DFT provides the wave-functions of the molecular orbitals
- Molecular Orbitals are linear combinations of basis-set functions



DFT Results

- Orbitals are populated from the bottom (Aufbau principle)
- Each orbital can be occupied by at most two electrons (Pauli principle)
- HOMO/LUMO

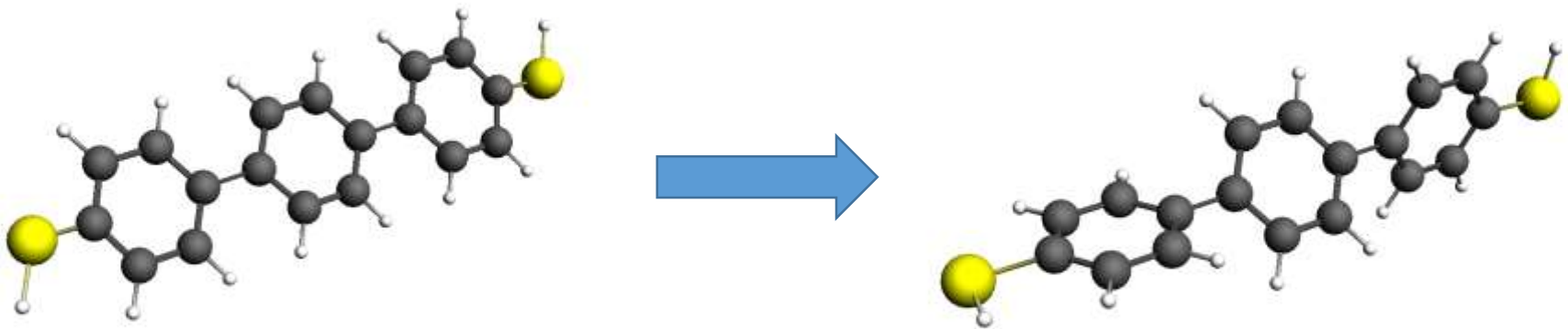


Summary DFT

- Density functional theory is an **exact reformulation** of many-body quantum mechanics in terms of the density rather than the wave function
- DFT provides electron density and hence the wave function
- XC functional unknown : electron exchange and correlation effects are approximated
- The result is as good as the functional/basis-set you are using

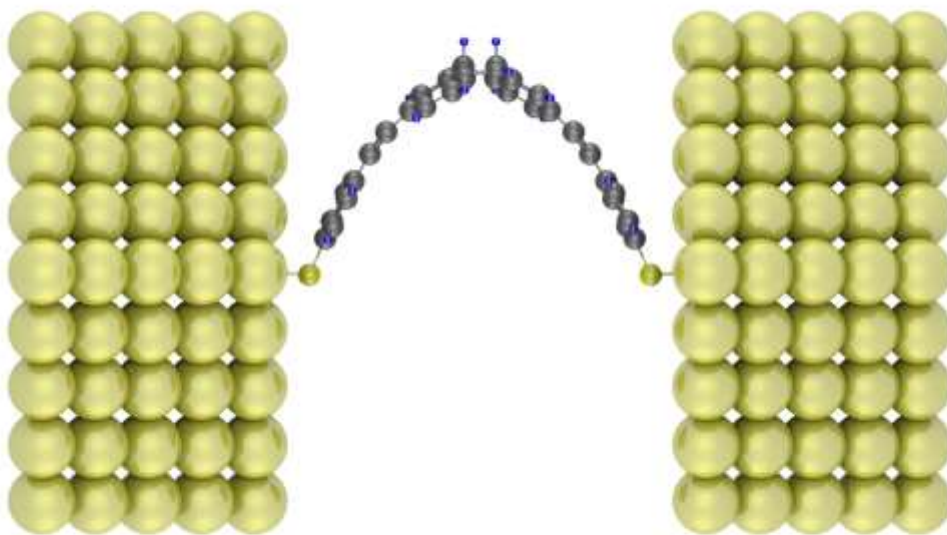
Geometry optimization

- Look for lowest energy configuration
- Based on energy gradient



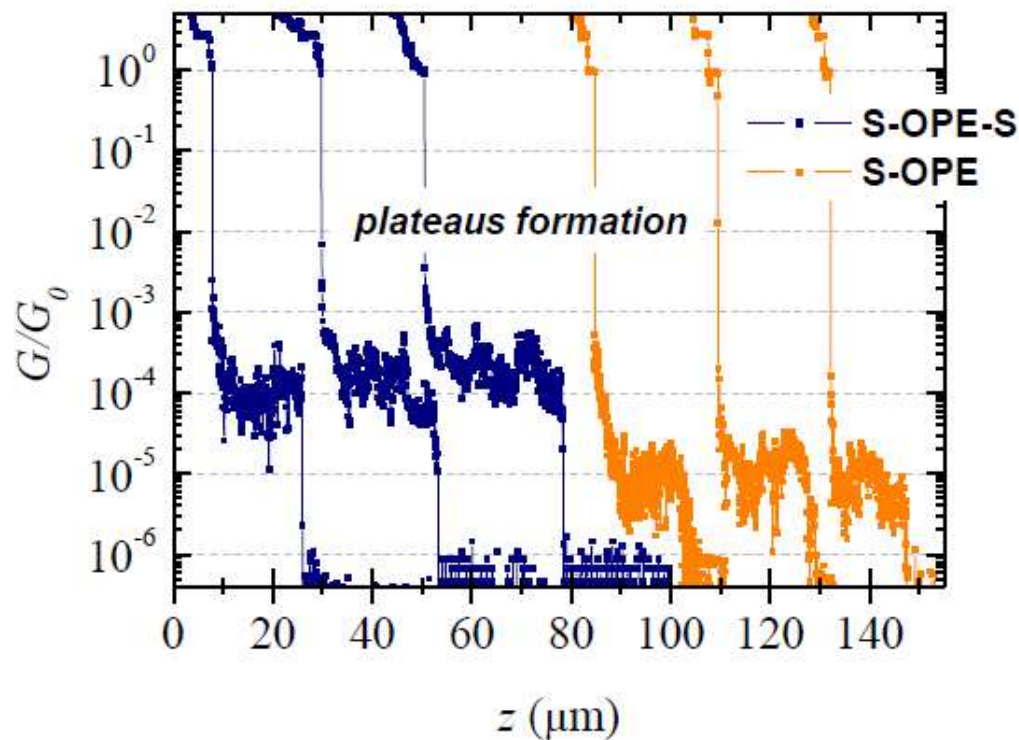
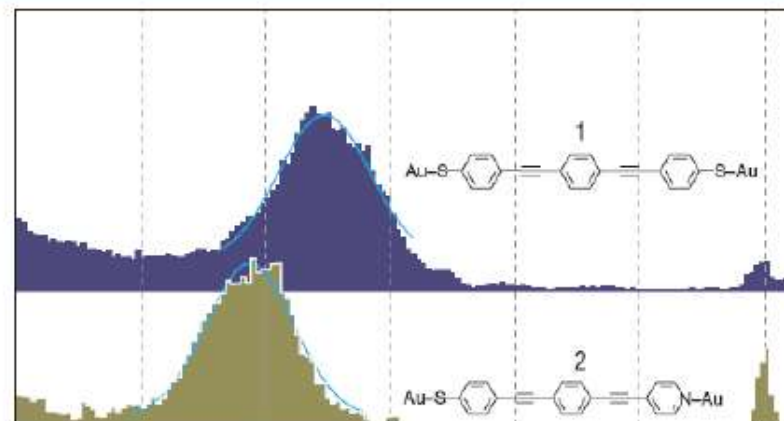
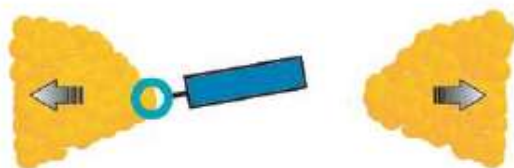
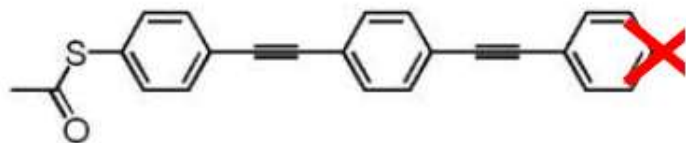
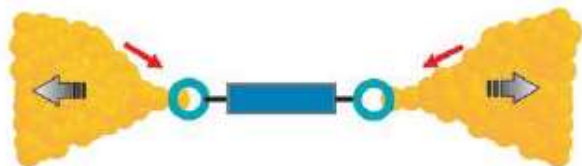
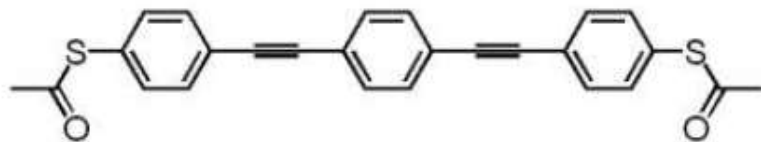
Geometry constraints

- Fix specific atoms or blocks of atoms
- Fix specific bond length, angles etc...

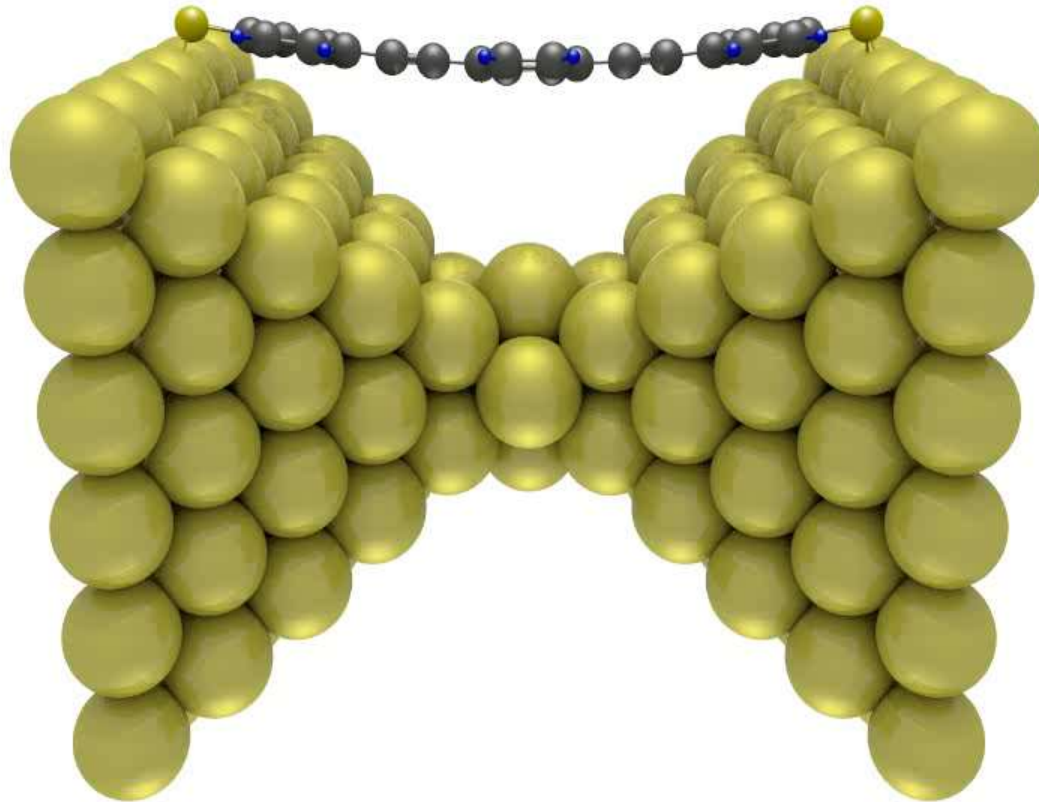


importance of intermolecular interactions

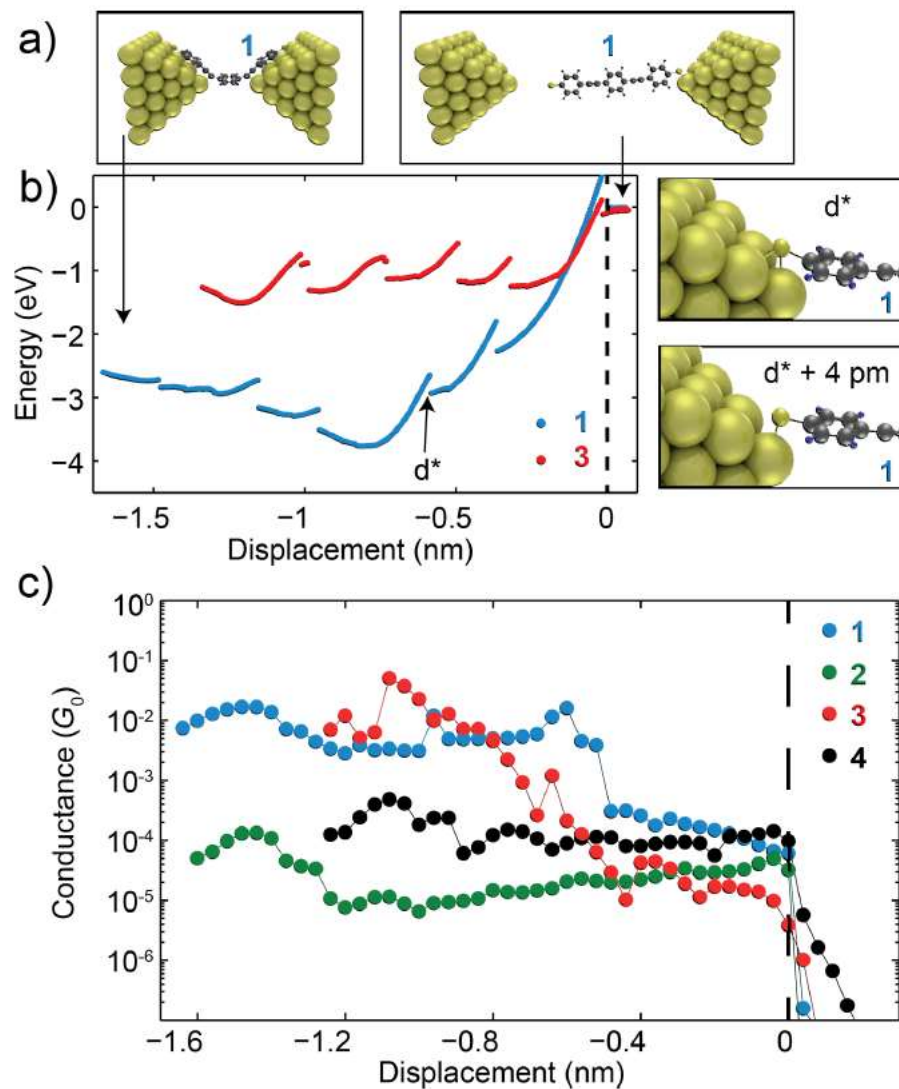
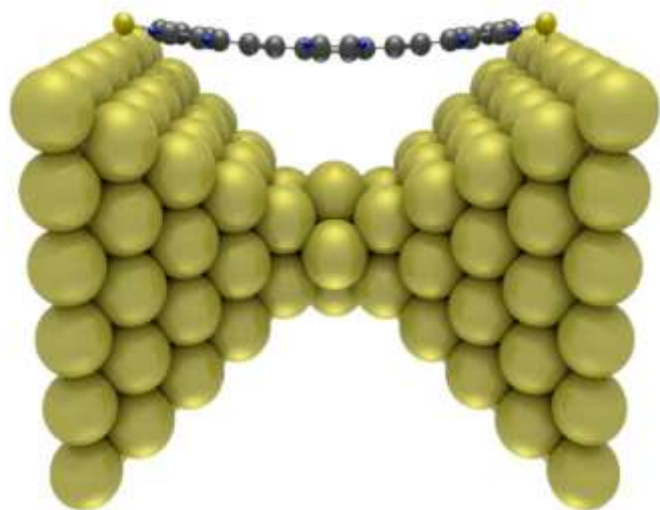
OPE: oligo (phenylene ethynylene)



Geometry constraints

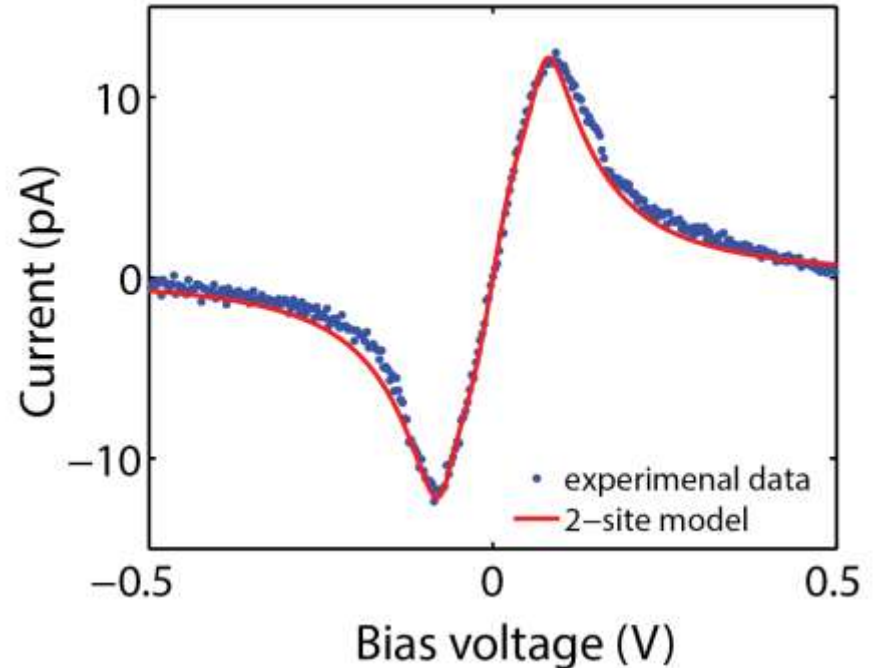
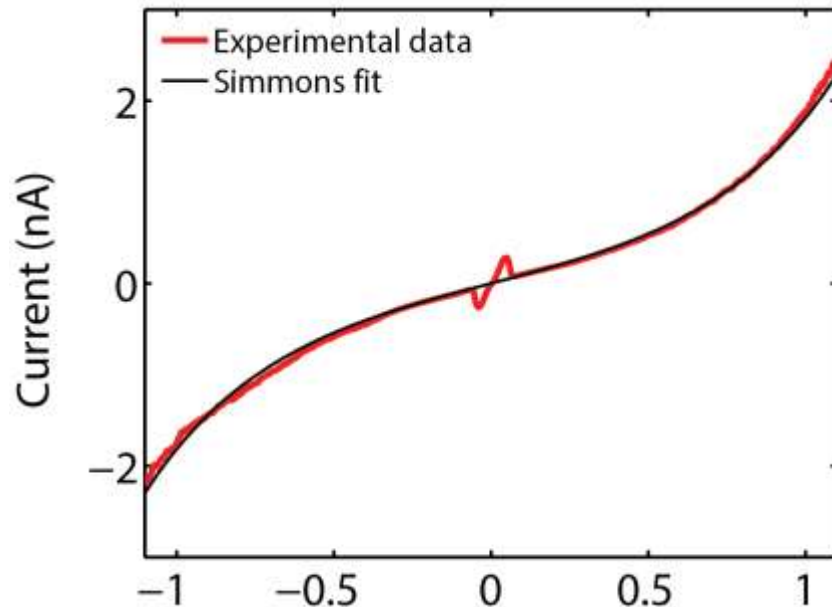


Geometry constraints



Electric fields

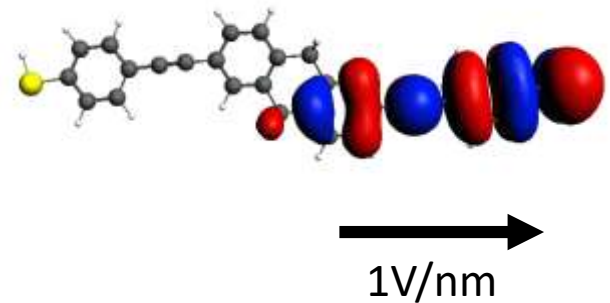
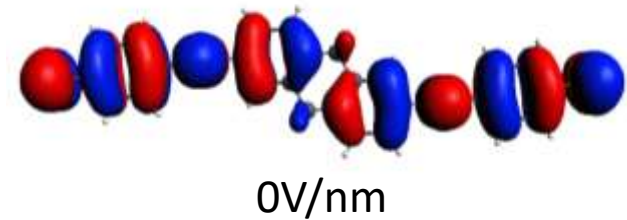
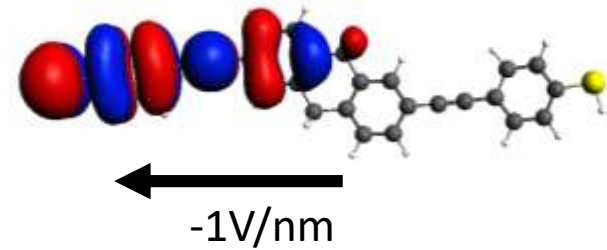
Perrin *et al.* Nat. Nano 2014



- High bias is dominated by single barrier tunneling
- Experimental data can be fitted with 2-site model

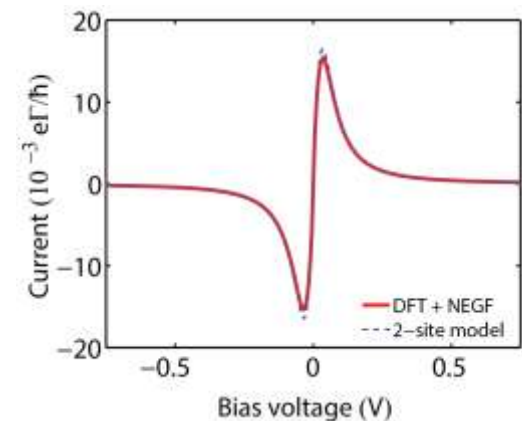
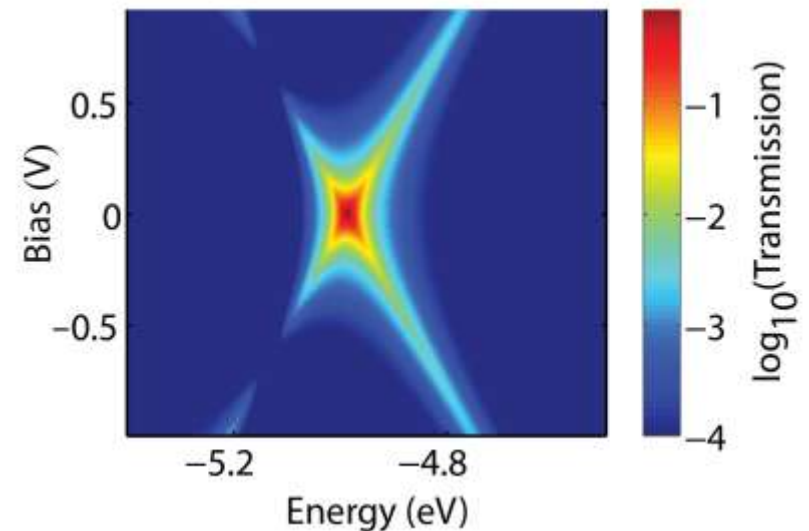
Electric fields

- An electric field can be added



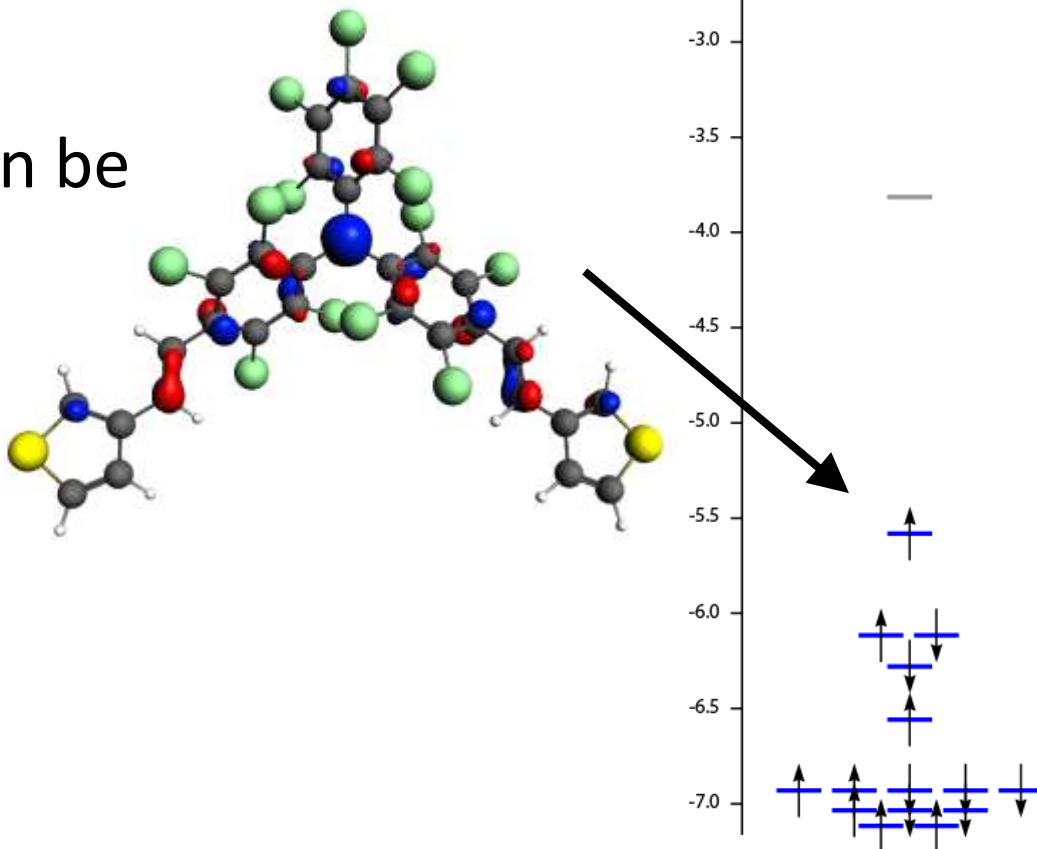
Electric fields

- An electric field can be added
- Bias dependence transmission
- Useful to calculate IVs

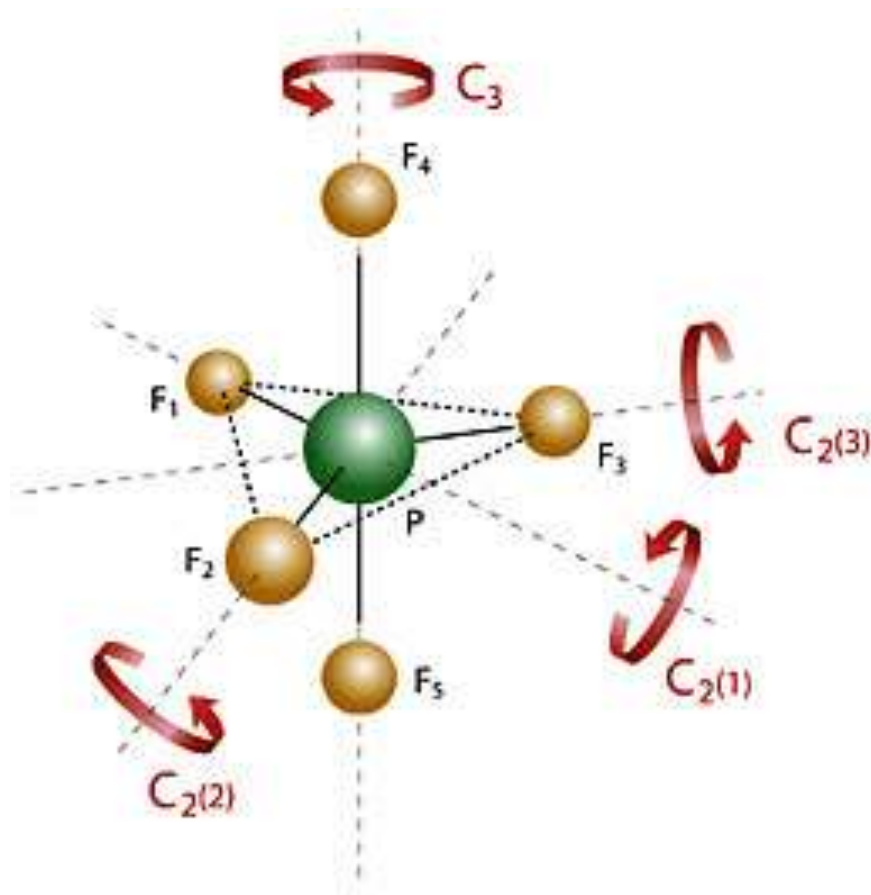


Spin resolved calculations

- Spin up and down are treated separately
- Spin unbalance can be added
- Add charge to molecule



SCF tricks/speed-up - Symmetry



- Enforce symmetry in geometry optimization (reflection, rotation,...)
- Easier to converge
- Speeds up the calculations

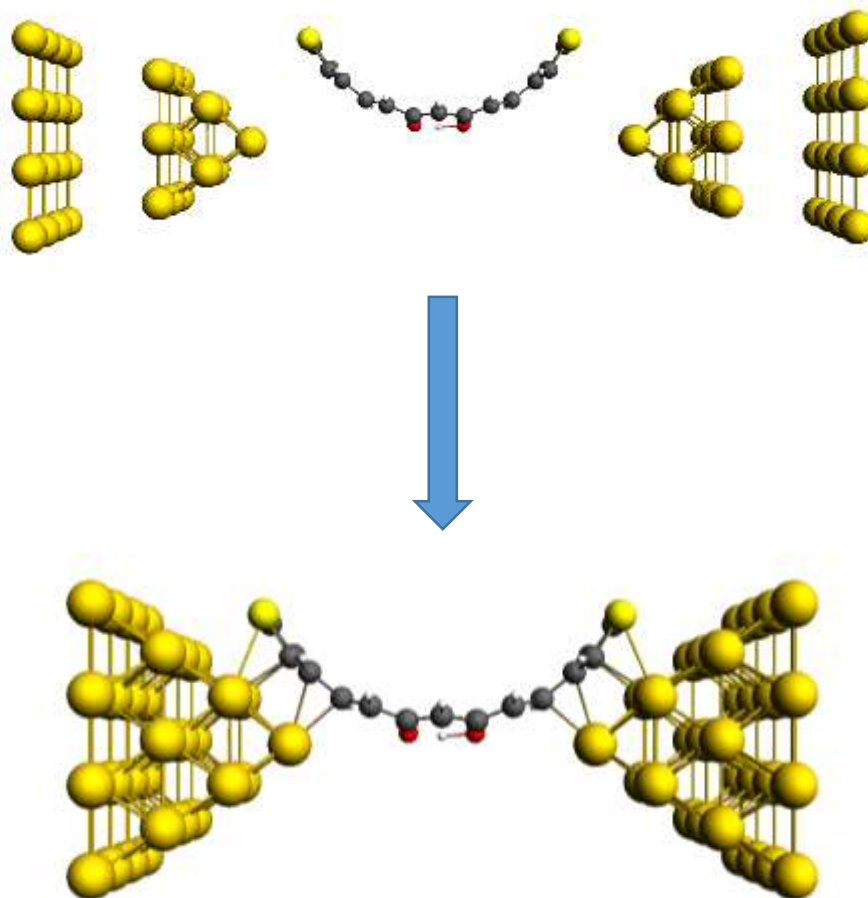
SCF tricks/speed-up – Fragments

- Fragments are first converged separately
- Provides better guess for initial density



SCF tricks/speed-up – Fragments

- Fragments are first converged separately
- Provides better guess for initial density
- Allows for convergence which otherwise would be difficult/impossible
- Speeds up convergence



SCF tricks/speed-up – Frozen core

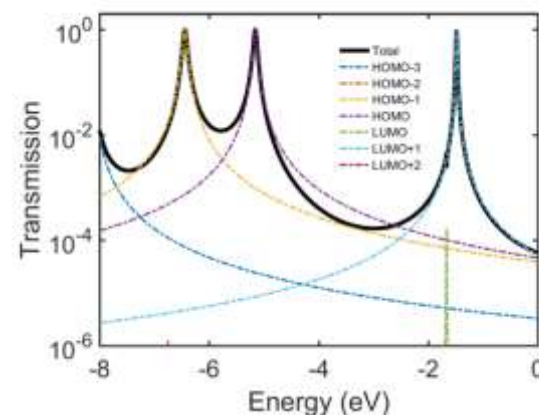
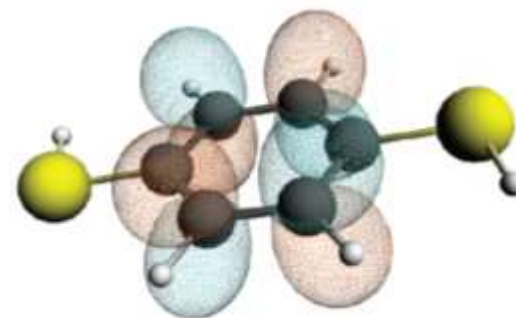
- Deep-core atomic orbitals change very little upon bond formation
- DFT can ‘freeze’ the core electrons of atoms

SCF tricks/speed-up – Frozen core

- Deep-core atomic orbitals change very little upon bond formation
- DFT can ‘freeze’ the core electrons of atoms
- Reduces the size of the variational basis set and speeds up calculations
- Core electrons taken into account when calculating energy

Overview

- Quantum chemistry (QC)
- Density functional theory (DFT)
- Non-equilibrium Green's function (NEGF)



NEGF

For charge transport, time dependent Schrödinger equation needs to be solved

$$i\hbar \frac{\partial}{\partial t} \Psi(\mathbf{r}, t) = \hat{H} \Psi(\mathbf{r}, t)$$

NEGF

For charge transport, time dependent Schrödinger equation needs to be solved

$$i\hbar \frac{\partial}{\partial t} \Psi(\mathbf{r}, t) = \hat{H} \Psi(\mathbf{r}, t)$$

After a lot of math, the transmission is given by

$$T(\epsilon) = \text{Tr} \{ \Gamma_L(\epsilon) \mathbf{G}^r(\epsilon) \Gamma_R(\epsilon) \mathbf{G}^a(\epsilon) \}$$

\mathbf{G}^r is the retarded Green's function

$$\mathbf{G}^r(\epsilon) = [\epsilon \mathbf{S} - \mathbf{H} - \Sigma_L^r(\epsilon) - \Sigma_R^r(\epsilon)]^{-1}$$

NEGF

$$\mathbf{G}^r(\epsilon) = [\epsilon \mathbf{S} - \mathbf{H} - \Sigma_L^r(\epsilon) - \Sigma_R^r(\epsilon)]^{-1}$$

$\epsilon \rightarrow$ Energy incoming electron

$\mathbf{S} \rightarrow$ Overlap matrix (DFT)

Overlap of basis-set functions (atomic orbitals).

If orthonormal basis, \mathbf{S} is identity matrix

$\mathbf{H} \rightarrow$ Fock matrix (DFT)

Eigenvalues: Orbital energies

Eigenvectors: Orbital shape, linear combination atomic orbitals

$\Sigma \rightarrow$ Self energy matrix (DFT/arbitrary)

$$\Sigma_{L,R}^r(\epsilon) = \Lambda_{L,R}(\epsilon) - \frac{i}{2}\Gamma_{L,R}(\epsilon)$$

Wide-band limit

Wide-band limit (WBL) assumes self-energy is energy independent

$$\Sigma_{L,R}^r = \Lambda_{L,R} - \frac{i}{2}\Gamma_{L,R}$$

and neglects the real part

$$\Sigma_{L,R}^r = -\frac{i}{2}\Gamma_{L,R}$$

Wide-band limit

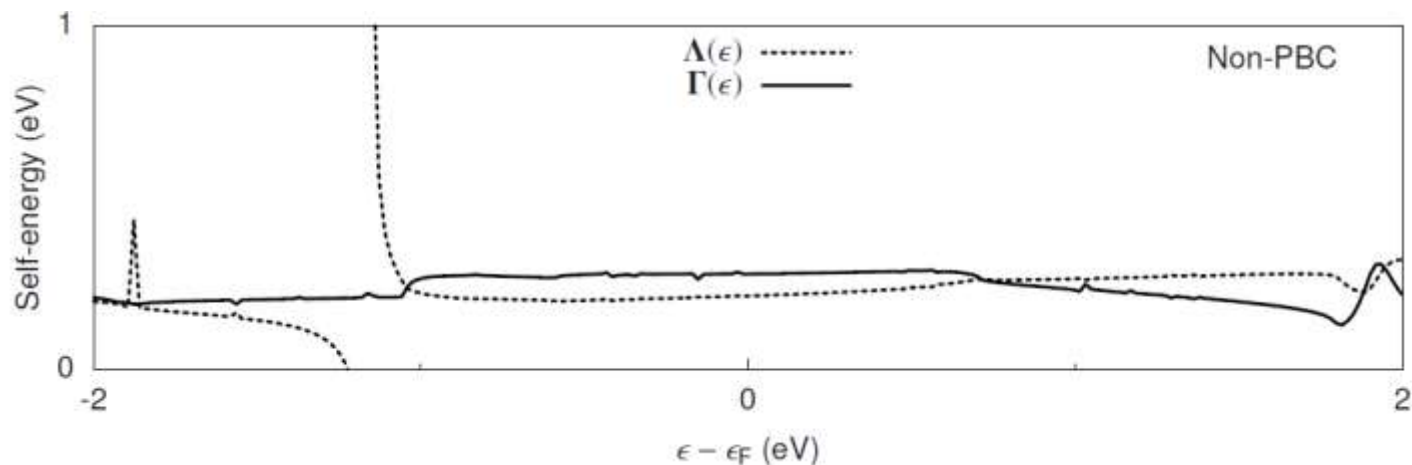
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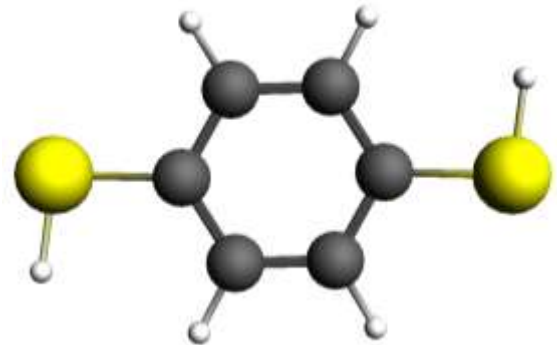
$$\Sigma_{L,R}^r = -\frac{i}{2}\Gamma_{L,R}$$

For gold this is a not-so-crazy assumption



Gas phase transmission

1. Take fock matrix from DFT (H)
2. Take overlap matrix from DFT (S)
3. Chose were to inject charge and construct $\Gamma_{R,L}$ accordingly.
4. Compute transmission

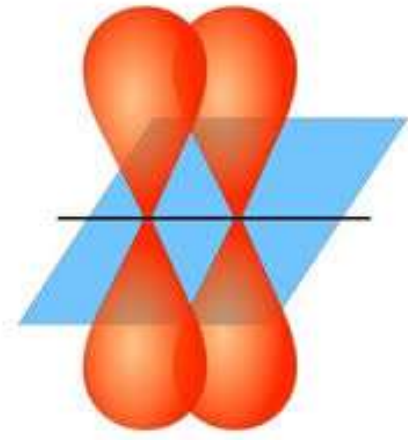


$$T(\epsilon) = \text{Tr} \{ \Gamma_L(\epsilon) G^r(\epsilon) \Gamma_R(\epsilon) G^a(\epsilon) \}$$

Neglects all electrodes/molecule interactions!

Gas phase transmission

1. Take fock matrix from DFT (H)
2. Take overlap matrix from DFT (S)
3. Chose were to inject charge and construct $\Gamma_{R,L}$ accordingly.
4. Compute transmission

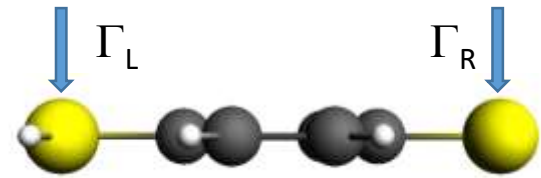


$$T(\epsilon) = \text{Tr} \{ \Gamma_L(\epsilon) G^r(\epsilon) \Gamma_R(\epsilon) G^a(\epsilon) \}$$

Neglects all electrodes/molecule interactions!

Gas phase transmission

1. Take fock matrix from DFT (H)
2. Take overlap matrix from DFT (S)
3. Chose were to inject charge and construct $\Gamma_{R,L}$ accordingly.
4. Compute transmission



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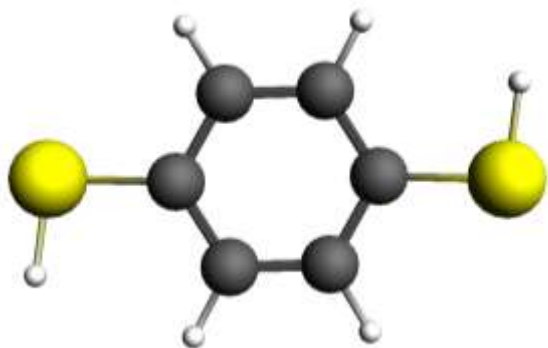
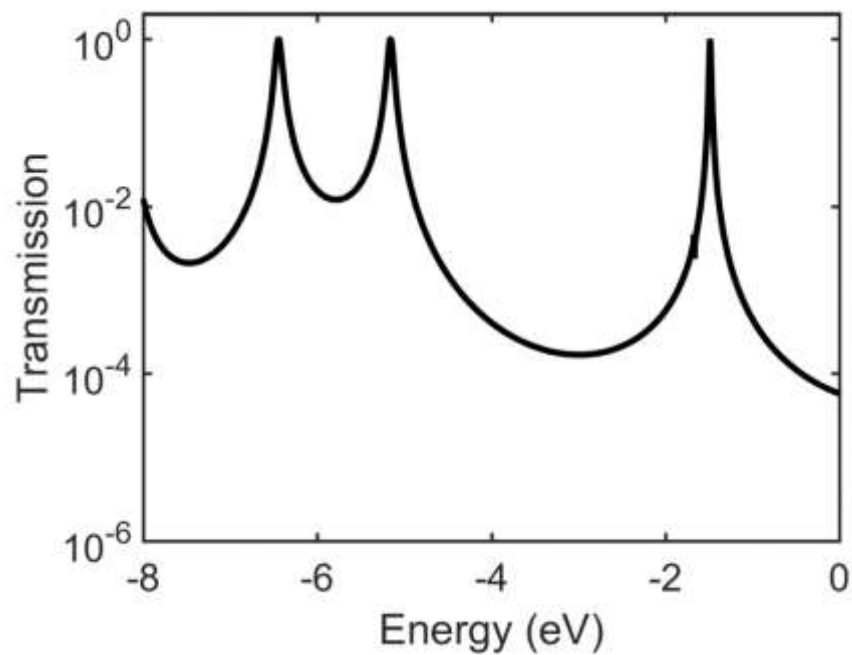
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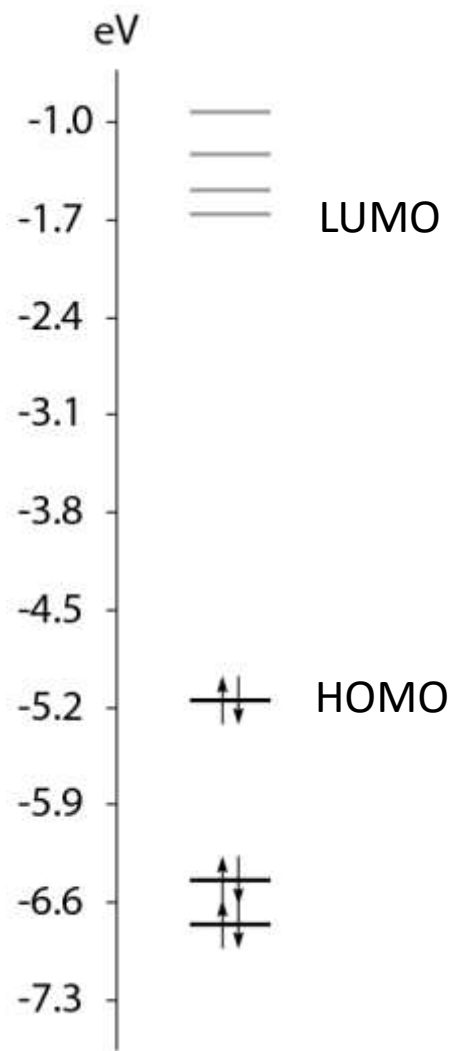
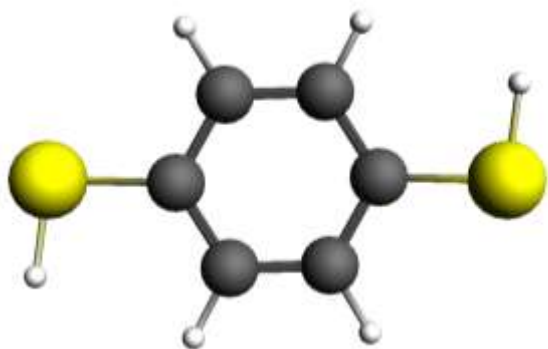
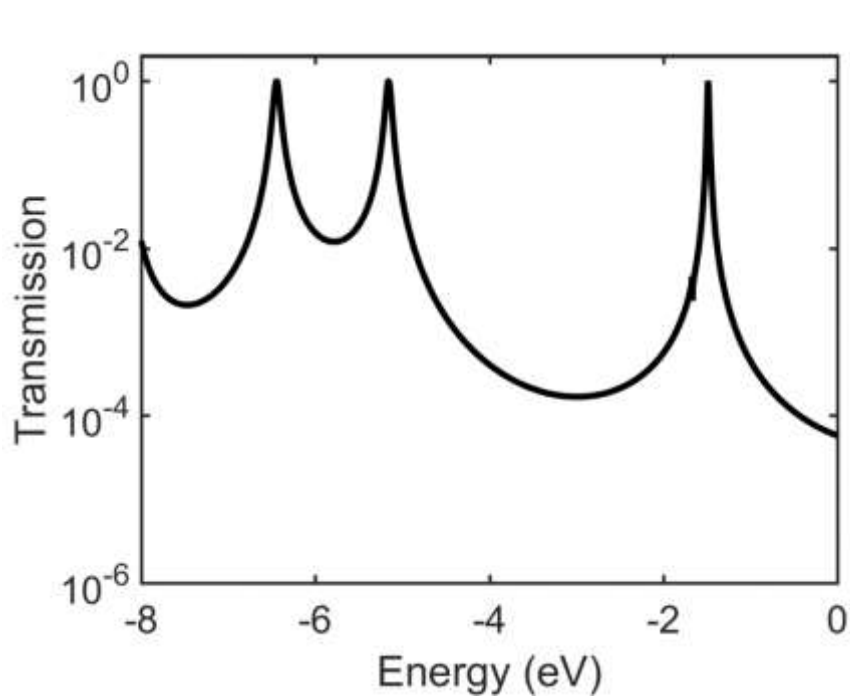
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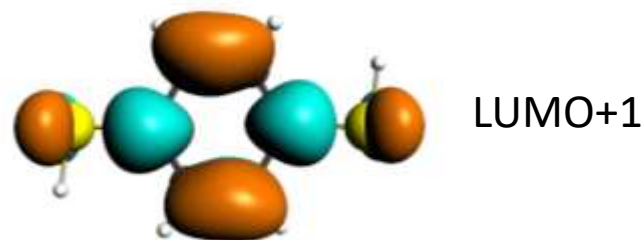
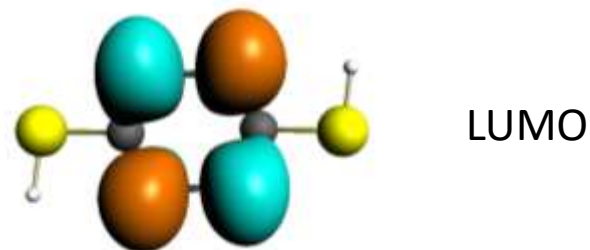
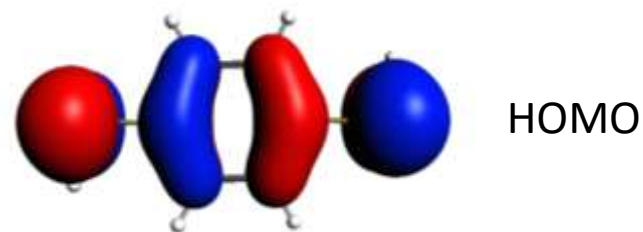
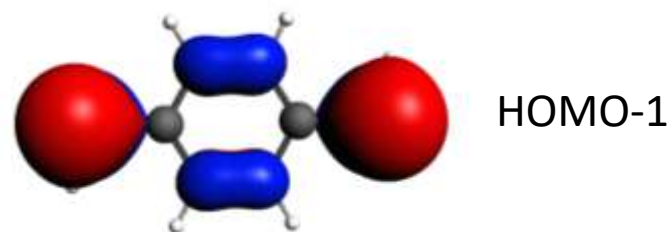
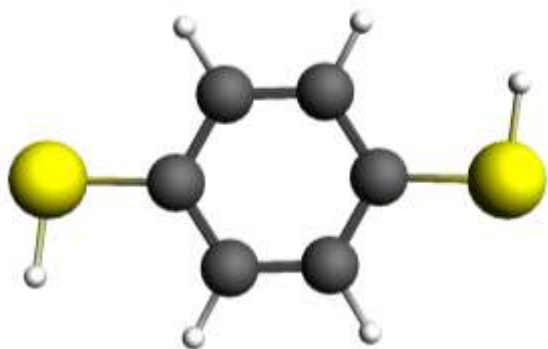
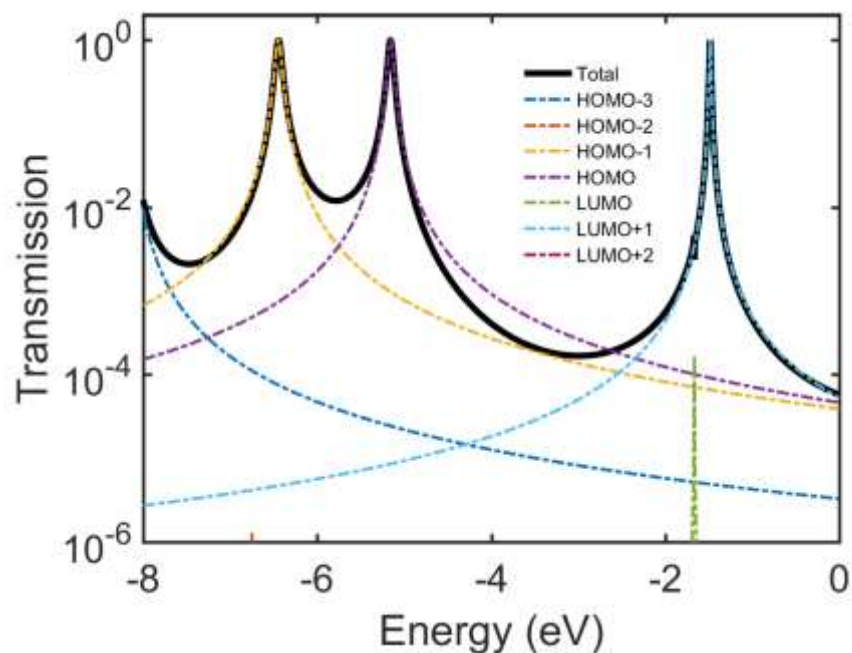
Gas phase transmission



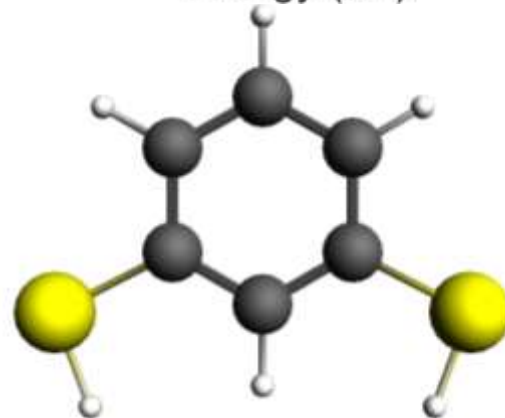
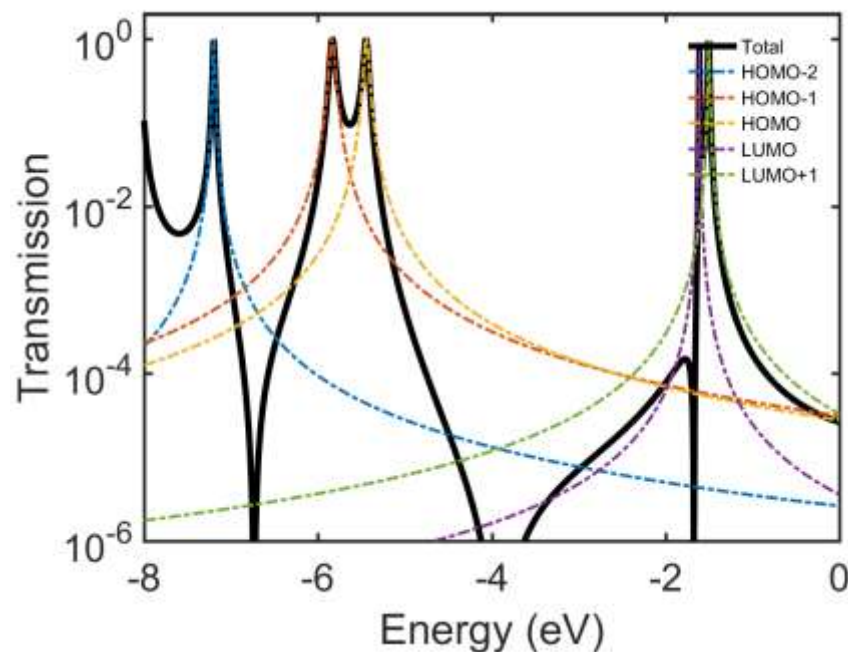
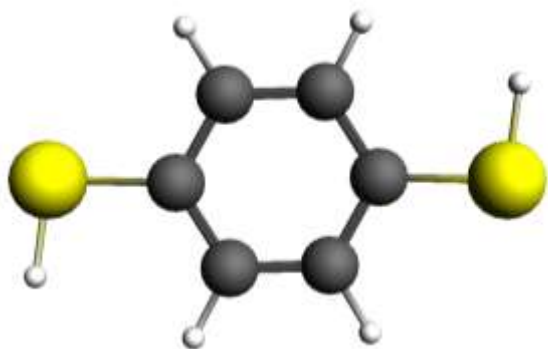
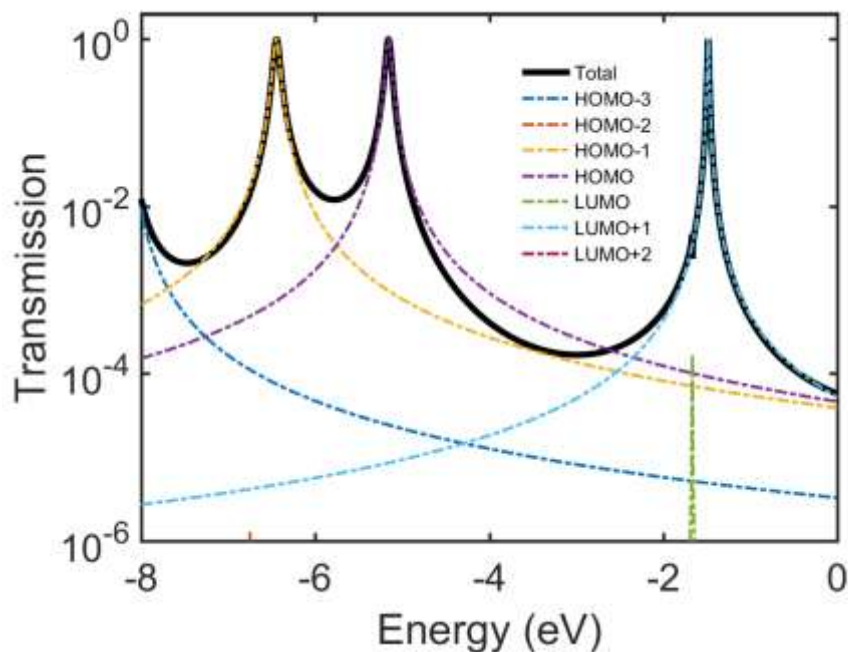
Gas phase transmission



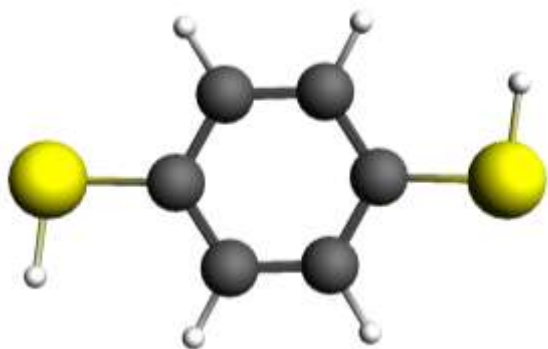
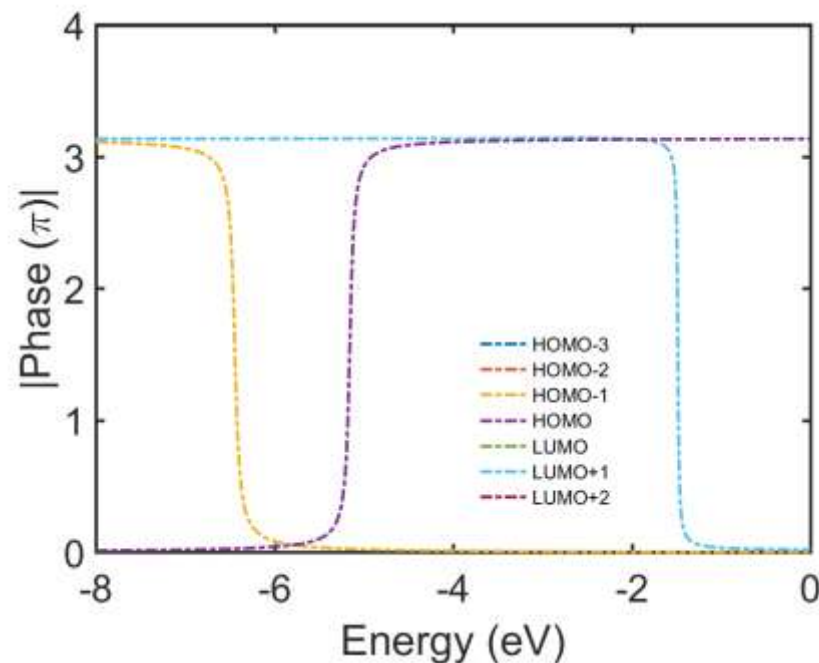
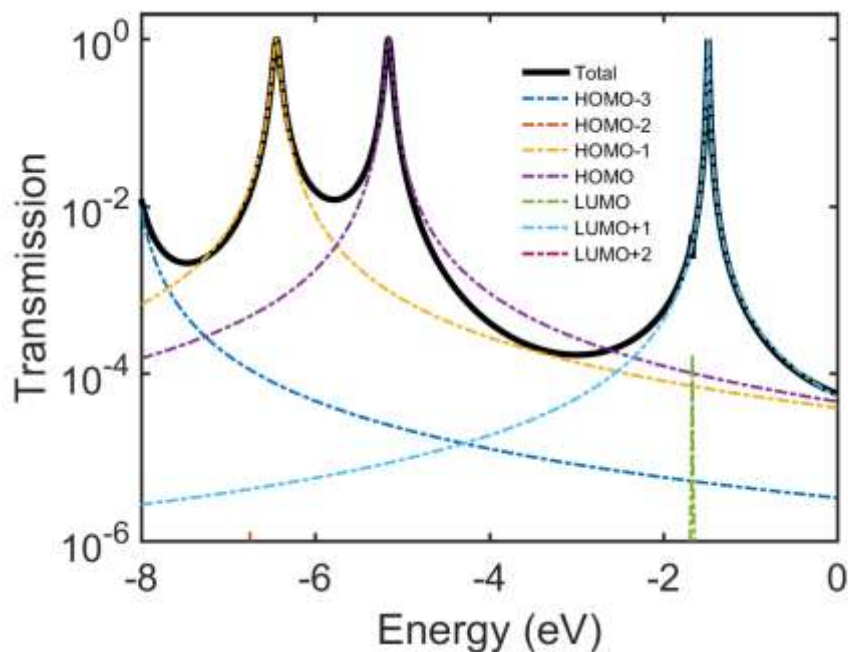
Orbital decomposition - amplitude



Orbital decomposition - amplitude



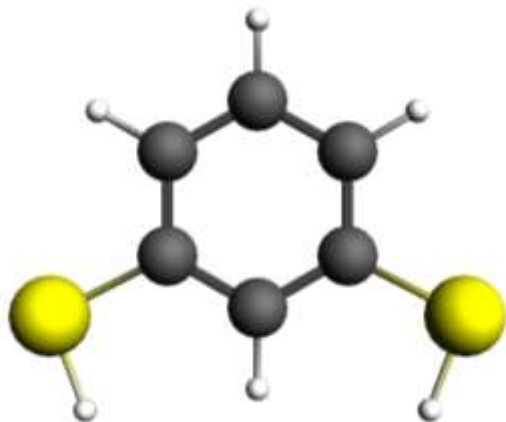
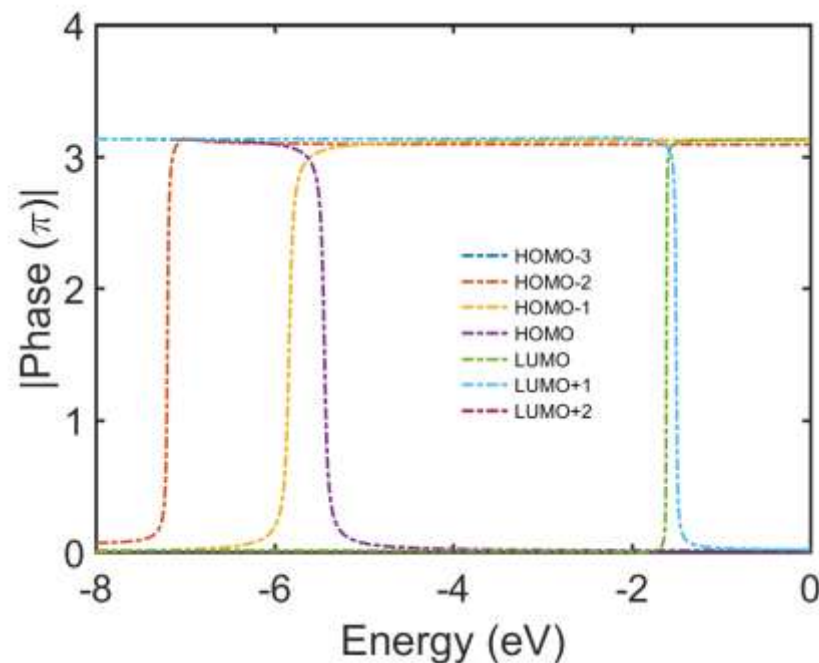
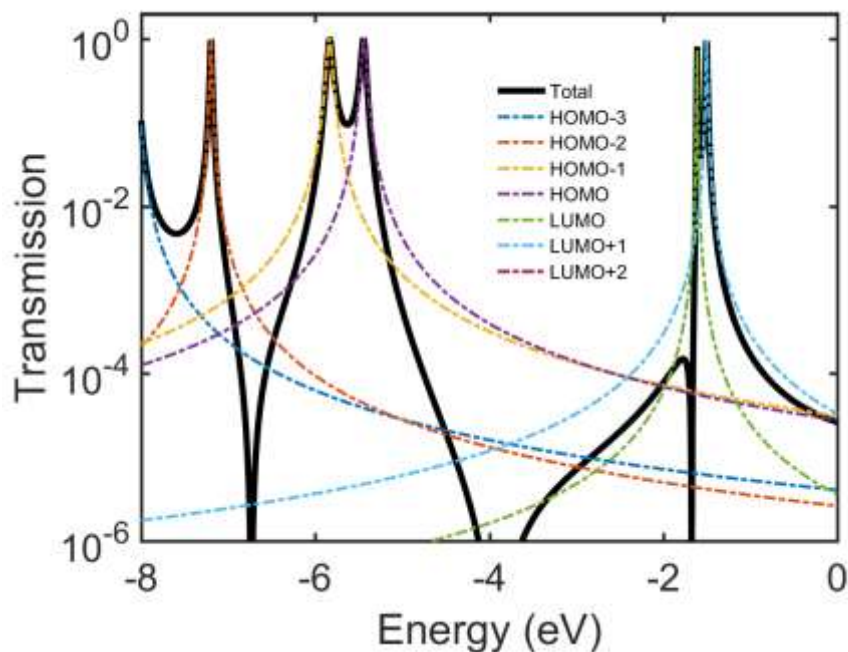
Orbital decomposition - phase



Phase is angle between real and complex part of the transmission

π -phase shift at energy of corresponding orbital

Orbital decomposition - phase

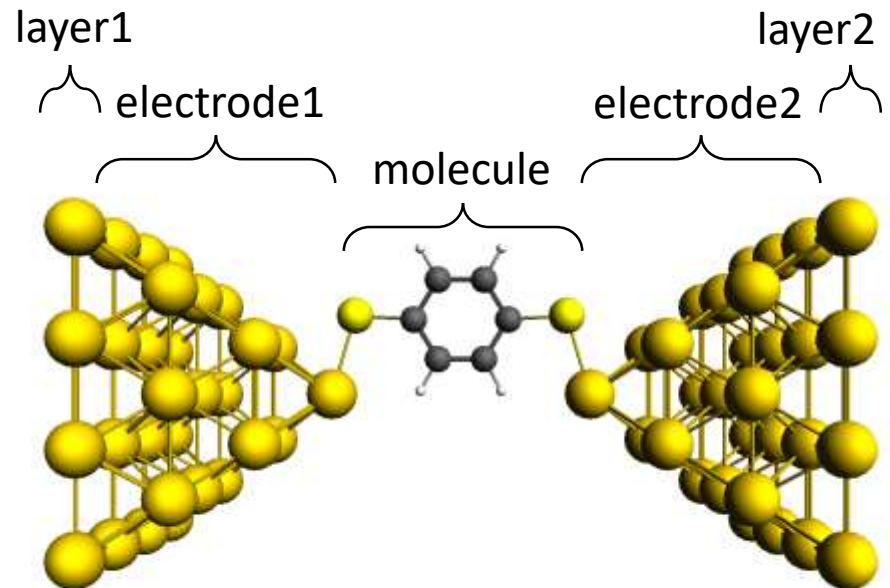
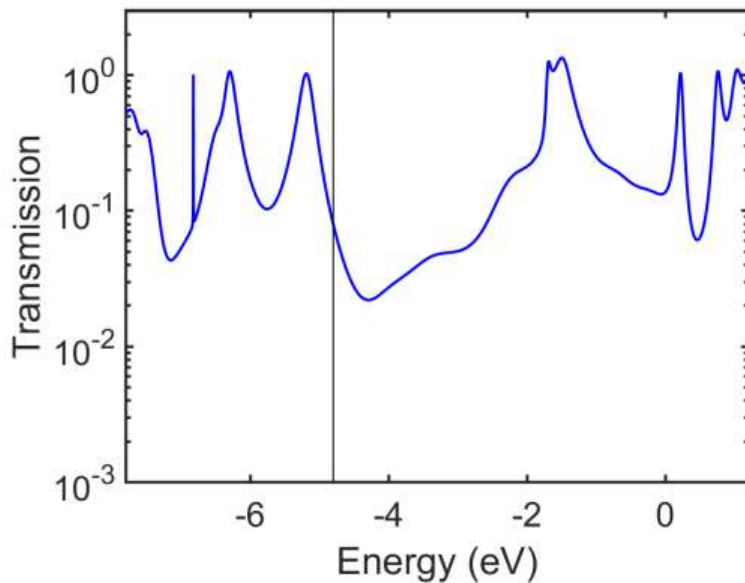


Phase is angle between real and complex part of the transmission

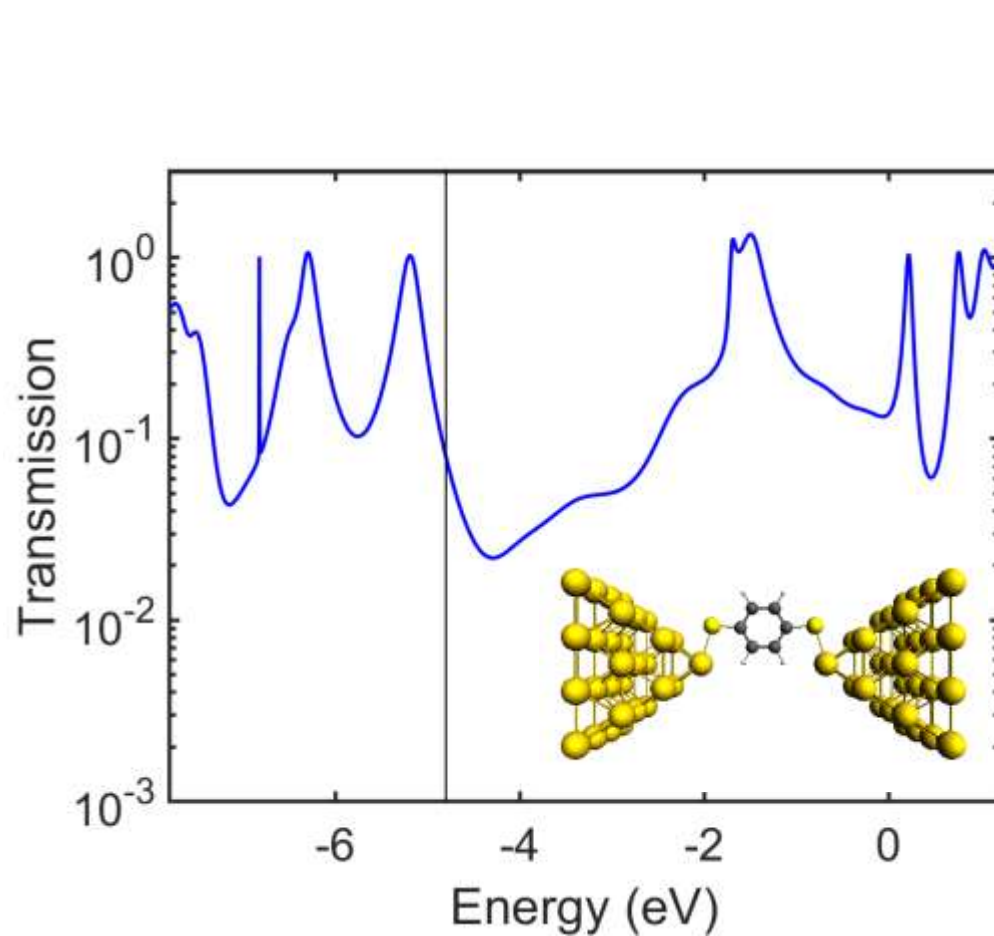
Destructive interference occurs when amplitude is the same, but π -phase shift

Transport with gold electrodes

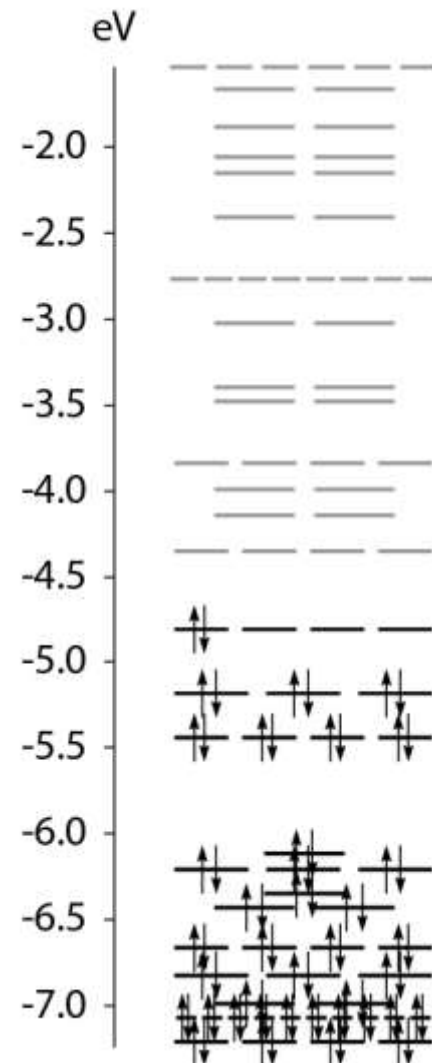
- Couple to atomic orbitals of gold of outermost layer instead of S atoms.



Transport with gold electrodes

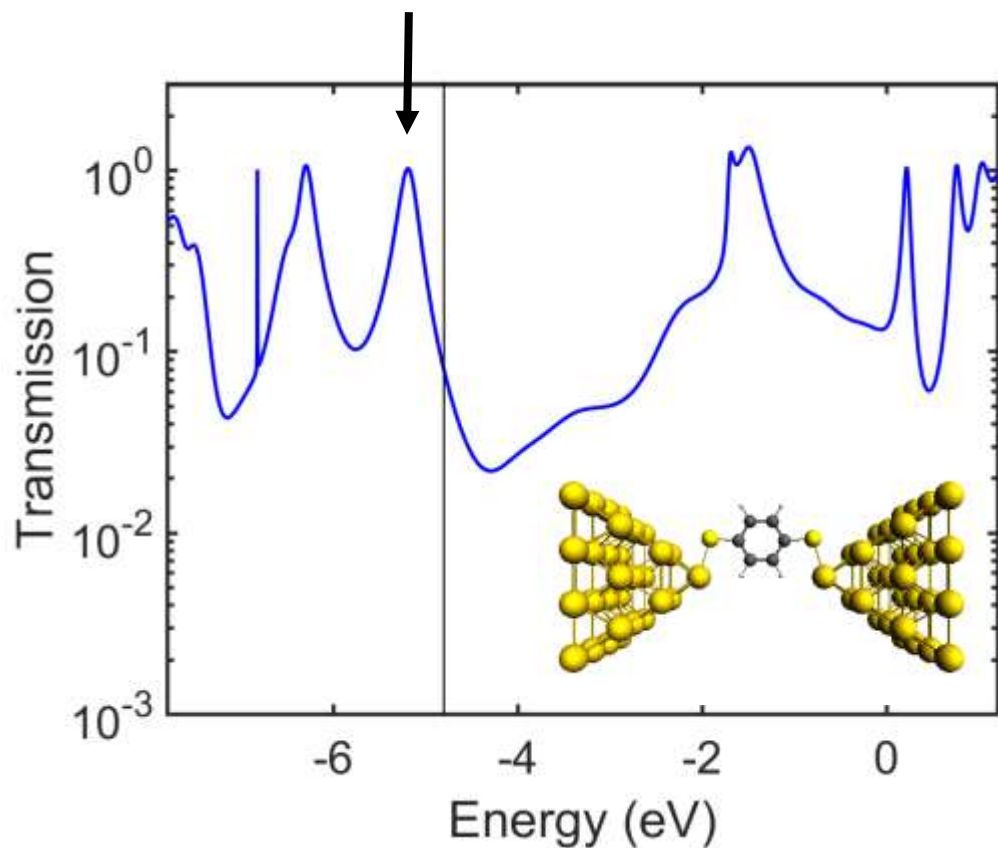


With so many orbitals, how to identify relevant ones?



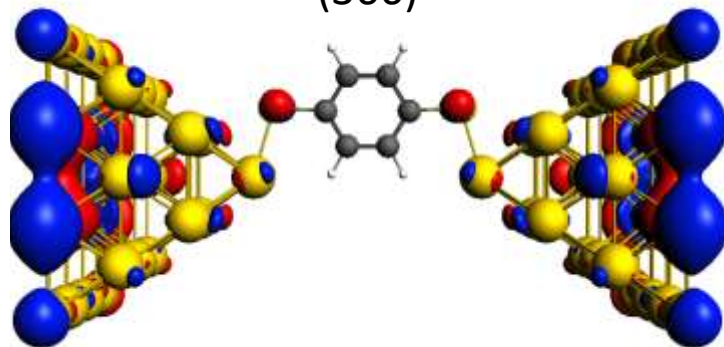
Transport with gold electrodes

'true' HOMO

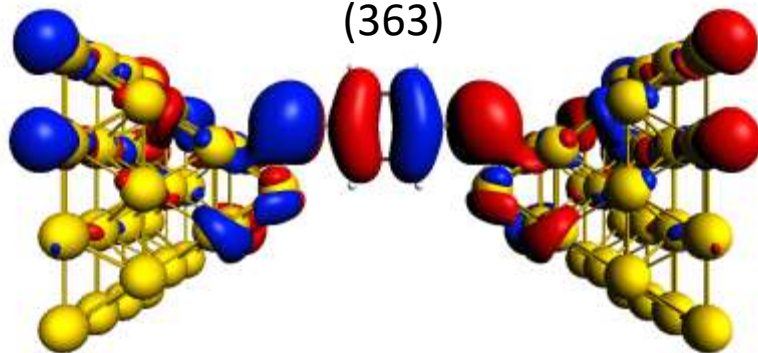


Which 'gas phase' orbital is responsible for transport?

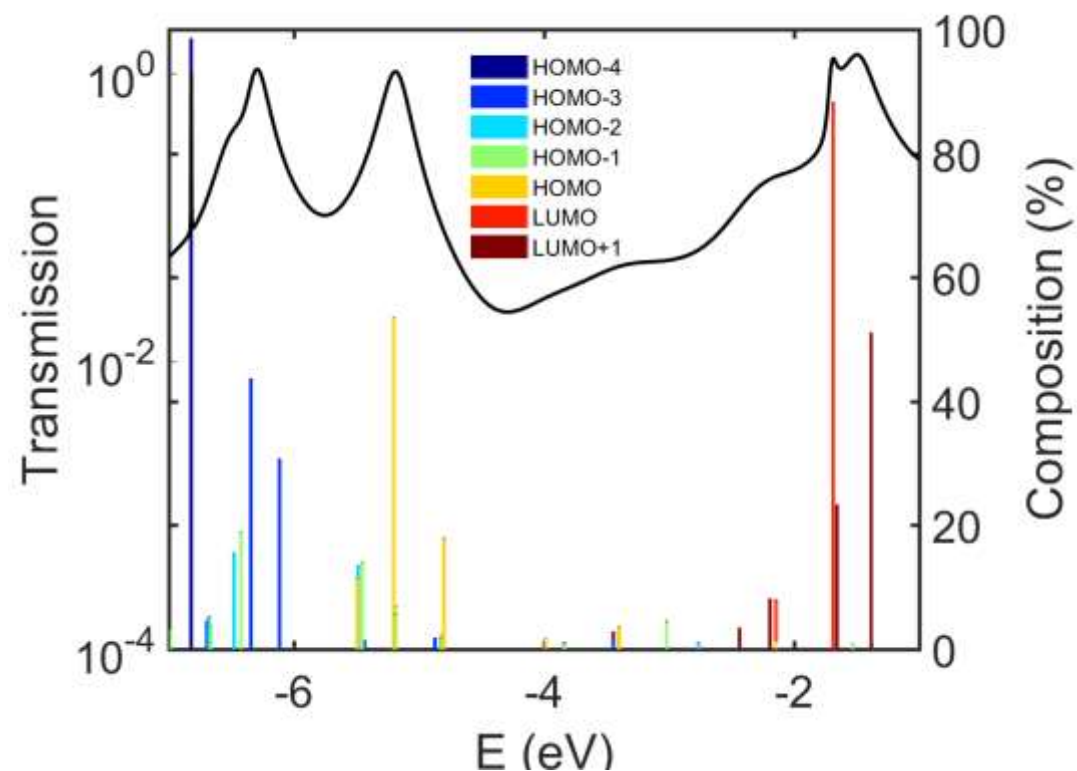
junction HOMO
(366)



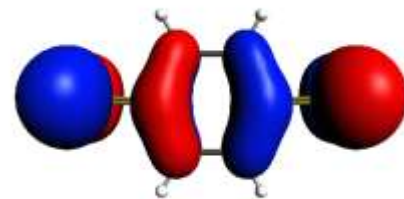
'true' HOMO
(363)



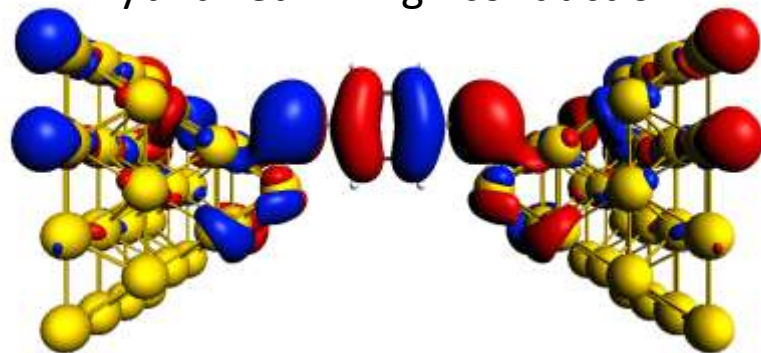
Orbital projections



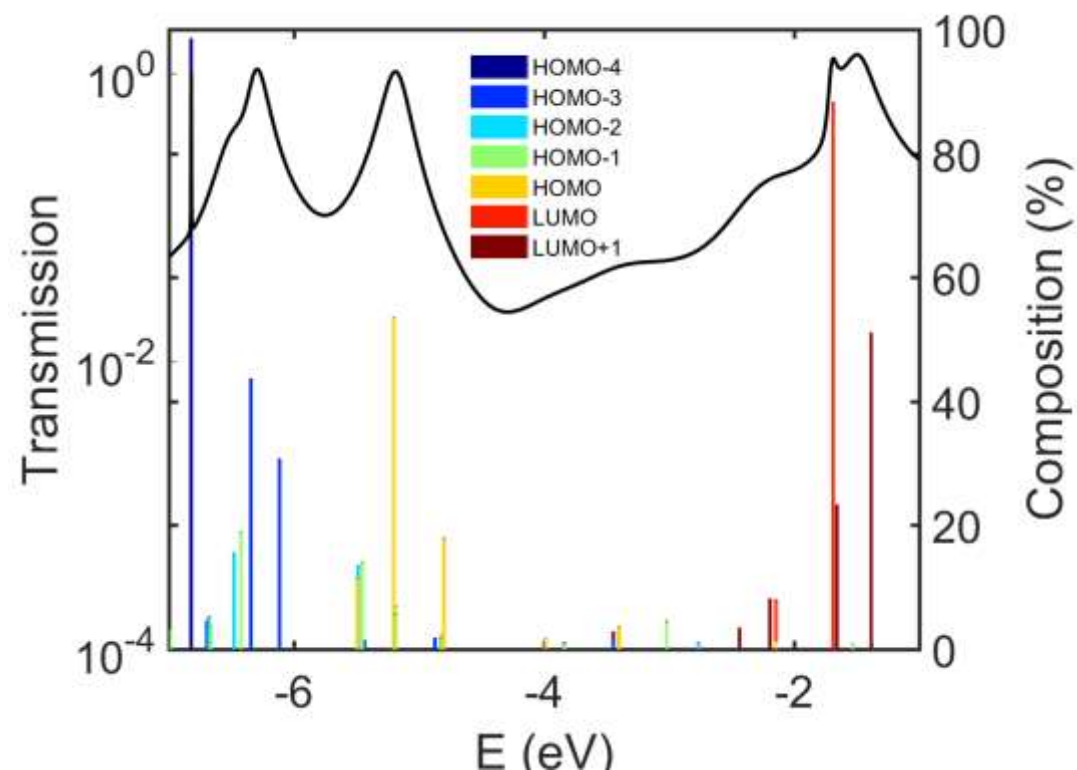
Gas phase HOMO



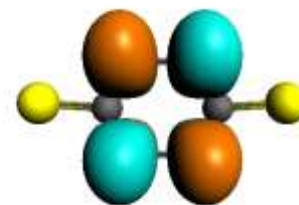
hybridized \rightarrow high conduction



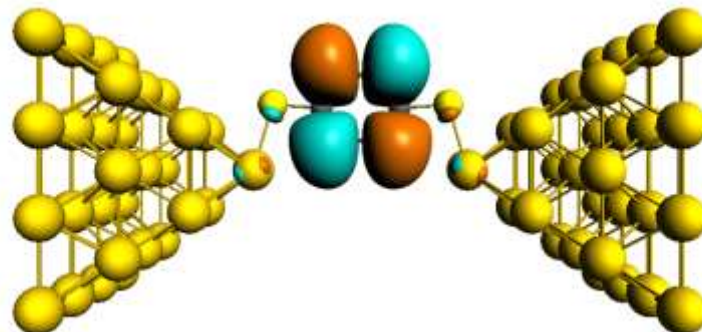
Orbital projections



Gas phase LUMO

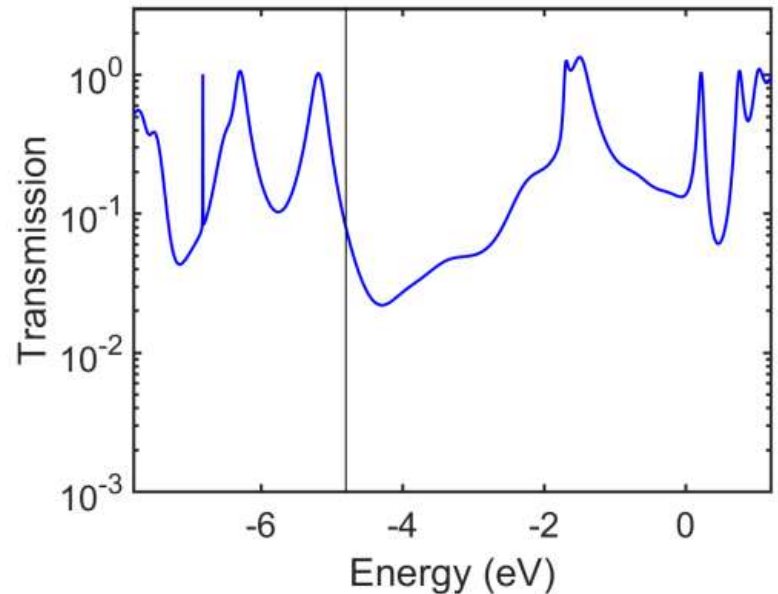


not hybridized \rightarrow poor conduction



How does this compare to experiments?

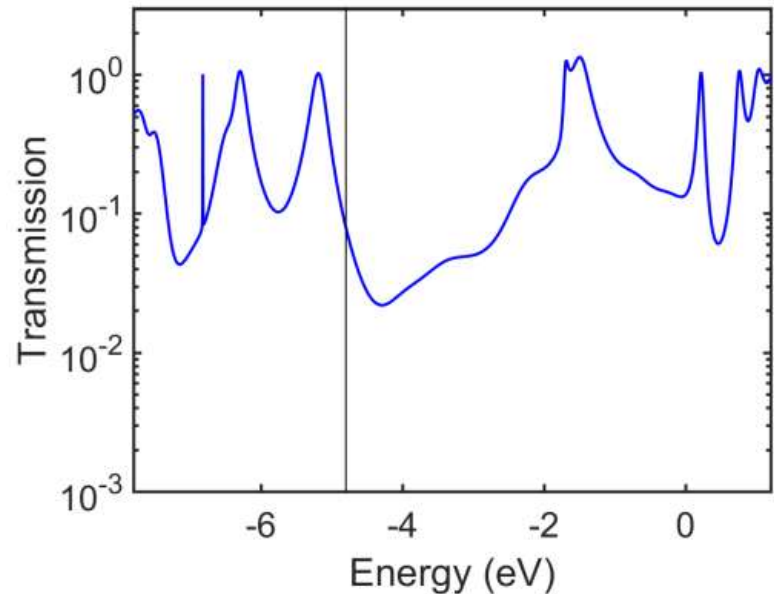
**Bandgap underestimated,
conductance overestimated**



How does this compare to experiments?

**Bandgap underestimated,
conductance overestimated**

1. DFT does not properly account for addition/removal of charge
2. Formation of image-charges in the electrodes upon addition/removal of charge not taken into account

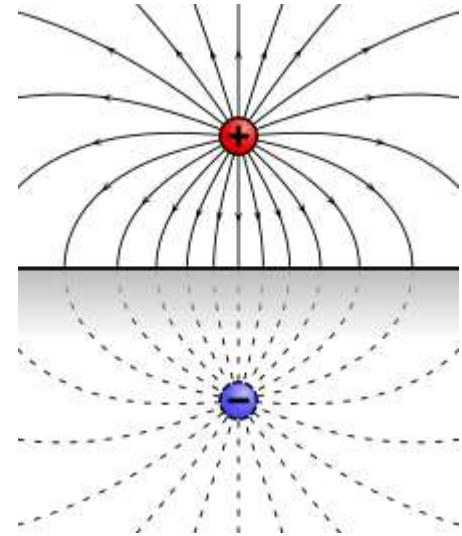


DFT + Σ

- Correct HOMO and LUMO by performing a **calculation for $\pm 1e$**
 - increases bandgap a lot

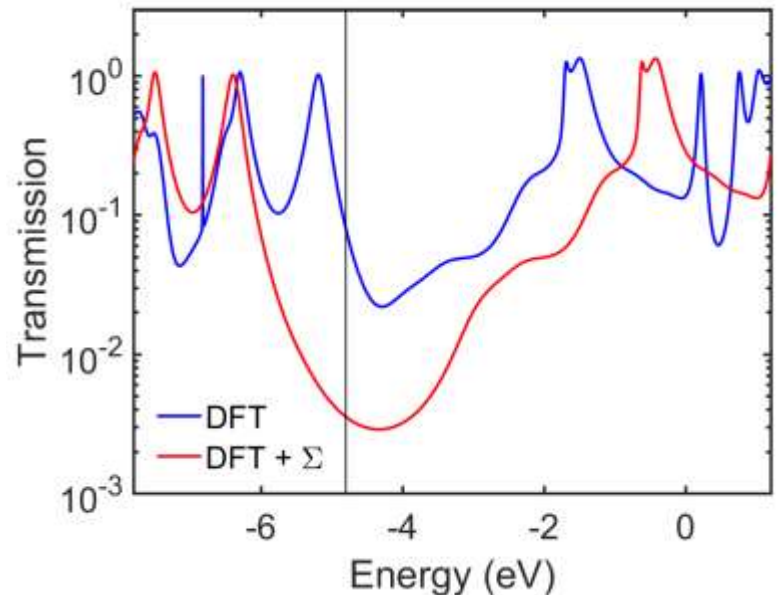
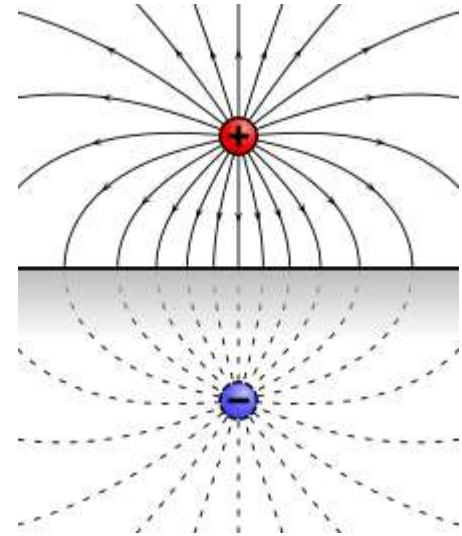
DFT + Σ

- Correct HOMO and LUMO by performing a **calculation for $\pm 1e$**
→ increases bandgap a lot
- Correct for **image-charge formation** in electrodes (classical electrostatics)
→ reduces bandgap a bit

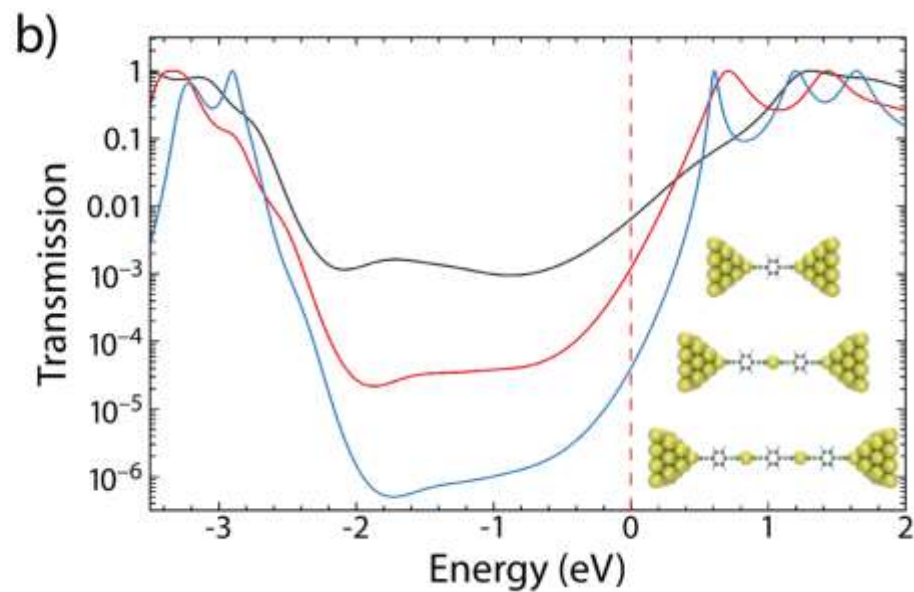
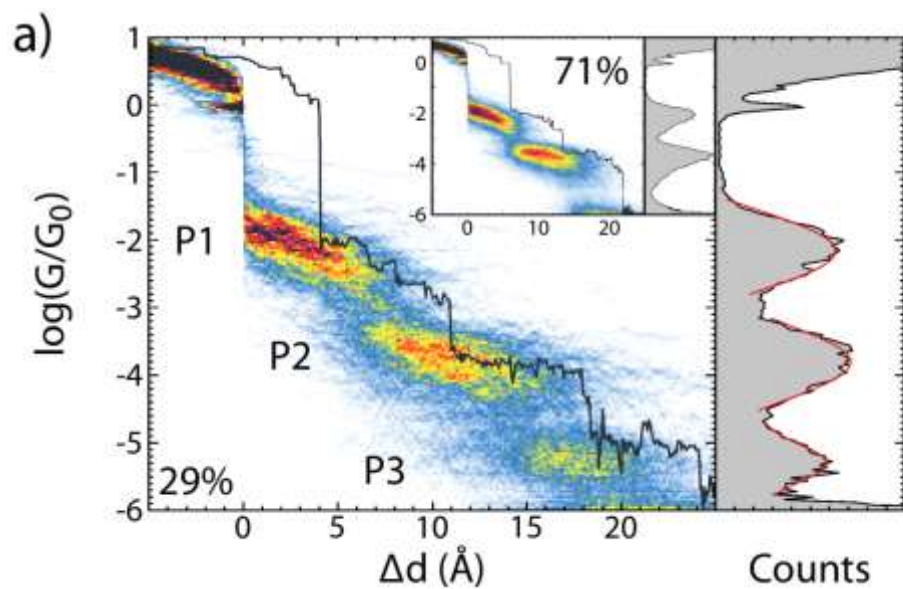


DFT + Σ

- Correct HOMO and LUMO by performing a **calculation for $\pm 1e$**
→ increases bandgap a lot
- Correct for **image-charge formation** in electrodes (classical electrostatics)
→ reduces bandgap a bit
- Shift for (un)occupied levels implemented as **scissor operator**
- Significantly improves agreement with experimental conductance values



Some examples...



Summary

- DFT provides electron density and hence the wave function
- NEGF gives you the electron transmission function through the system
- DFT+NEGF is very useful to understand experimental data
- Results should be interpreted with care

