

# *1<sup>st</sup> Tutorial:*

## The Basics of DFT

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MAX-PLANCK-GESSELLSCHAFT

Hands-on Summer School: Electronic Structure Theory  
for Materials and (Bio)molecules

# The ultimate goal!

$$H\Psi = E\Psi$$

Second order differential equation for a  $3N_e$ -variable function  $\Psi$

⇒ **Complex problem**

Unsolved issues at the simplest level of approximations (multiple solutions, ...)

# Goals of this tutorial

- Familiarise with practical aspects of electronic structure theory in general and density functional theory (DFT) in particular
- Hartree-Fock (HF) method and Kohn-Sham DFT (non-periodic)
- Numerical solution of the approximate equations (tool: FHI-aims)
- Exploring potential energy surfaces (total energies at fixed nuclei, local minima, vibrational spectra)
- Electronic structure analysis (visualisation tools, electron density, Kohn-Sham orbitals and spectrum)

# Solving the Kohn-Sham equations

**Hohenberg-Kohn Theorem**  $\Psi(\mathbf{r}_1 \dots \mathbf{r}_{N_e}) \Leftrightarrow n(\mathbf{r})$

## Kohn-Sham scheme

$$\left( -\frac{1}{2} \nabla^2 + \int d^3 r' \frac{n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} + v_{xc} + v \right) \psi_i = \epsilon_i \psi_i$$
$$\Rightarrow n = \sum_i f_i |\psi_i|^2$$

**KS Orbitals**  $\{\psi_i\}$   $\langle \psi_i, \psi_j \rangle = \delta_{ij}$

**XC Potential**  $v_{xc}$  unknown, but many approximations exist

**LDA, PBE, ...**  $\rightarrow$  control.in

**External potential** contains ionic contributions  $\rightarrow$  geometry.in

# Basis set

**Expand in a finite basis  $\{\phi_i\}$ :**  $\psi_j = \sum_{i=1}^N c_{ij}\phi_i$

## Finite Basis

### Numeric atom centered

Gaussians

Plane waves + pseudoisation

Slater type

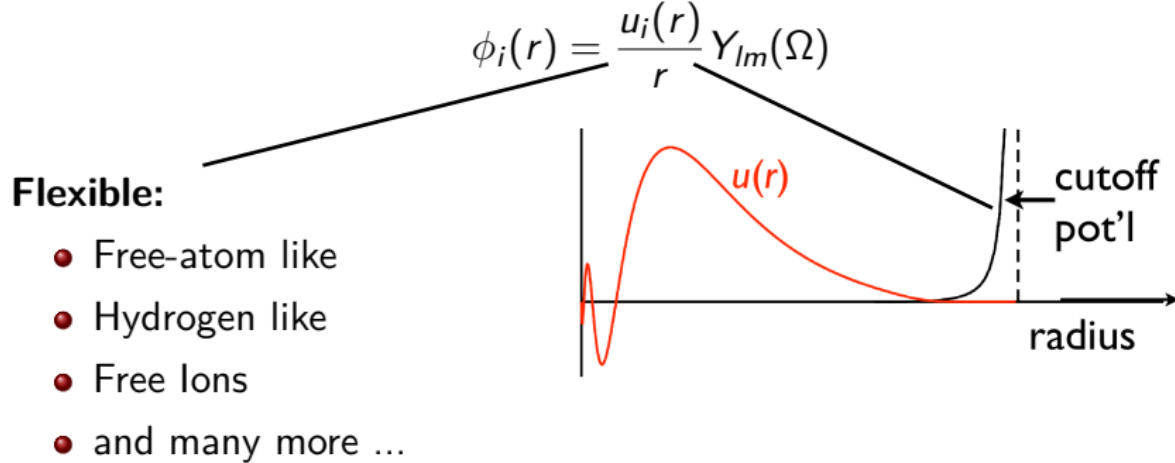
Grid based

... many more

# Basis set

**Expand in a finite basis  $\{\phi_i\}$ :**  $\psi_j = \sum_{i=1}^N c_{ij} \phi_i$

## Numeric atom centered (FHI-aims)



Courtesy V. Blum

## Basis set

**Expand in a finite basis  $\{\phi_i\}$ :**  $\psi_j = \sum_{i=1}^N c_{ij} \phi_i$



**Generalized matrix eigenvalue equation in  $c_{ij}$**

$$\hat{h}^{KS} \psi = E \psi \quad \Rightarrow \quad \sum_j h_{ij}(c) c_{jl} = \epsilon_l \sum_j s_{ij} c_{jl}$$

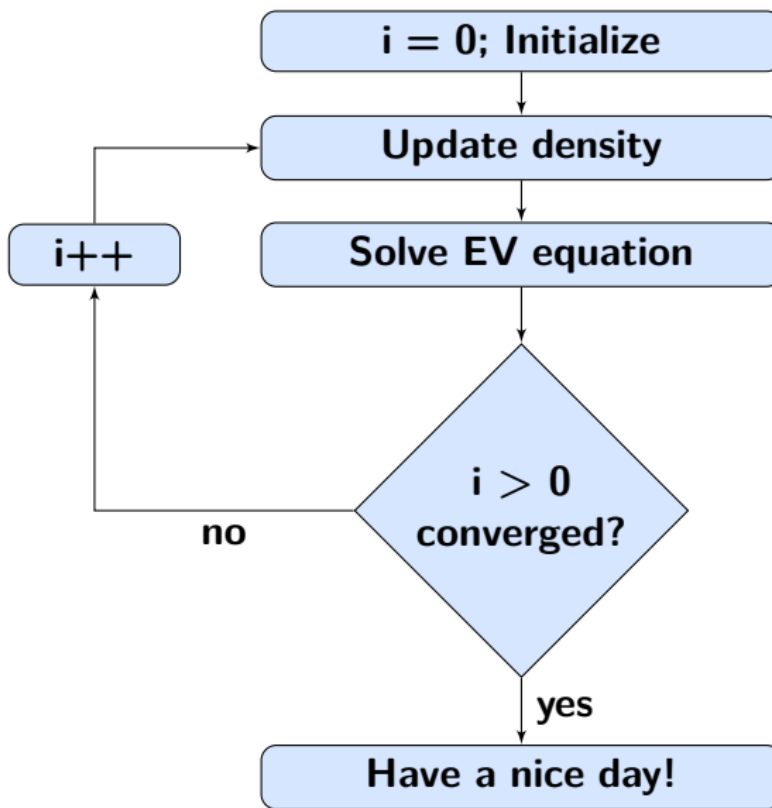
Overlap matrix  $s_{ij} = \langle \phi_i, \phi_j \rangle$

Hamilton matrix  $h_{ij} = \langle \phi_i, \hat{h}^{KS} \phi_j \rangle$



**Self-consistent solution**

# Finding the self-consistent solution



Mixing (Pulay)

(Sca)Lapack  
ELPA

Criteria

- Energy
- Charge density
- Sum of eigenvalues
- Force

# Problem I: The hydrogen atom

## Tasks:

- Input files needed to run FHI-aims.
- Test the convergence of the total energy with basis size.
- Compare the total energy of the hydrogen atom computed with different methods implemented in FHI-aims. Do all methods converge to the same result?

Basic electronic structure with FHI-aims

# FHI-aims: input files

`geometry.in`

`control.in`

## Basic electronic structure with FHI-aims

## FHI-aims: input files

**geometry.in**

# Atomic structure

# x y z

atom 0.0 0.0 0.0 H

atom 1.0 0.0 0.0 H

initial\_moment 1.0

# That's a comment

**control.in****Units:**

Positions in Å

Energies in eV

**Manual, chap. 2.1**

## Basic electronic structure with FHI-aims

## FHI-aims: input files

**geometry.in****# Atomic structure**

# x y z

atom 0.0 0.0 0.0 H

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# That's a comment

**control.in****# Physical model settings**

xc pw-lda

charge 0.

spin collinear

**Units:****Positions in Å****Energies in eV****Manual, chap. 2.1**

## Basic electronic structure with FHI-aims

## FHI-aims: input files

**geometry.in****# Atomic structure**

```
#      x     y     z
atom 0.0 0.0 0.0 H
atom 1.0 0.0 0.0 H
initial_moment 1.0
# That's a comment
```

**control.in****# Physical model settings**

```
xc pw-lda
charge 0.
spin collinear
```

**# SCF convergence settings**

```
sc_accuracy_eev 1E-2
sc_accuracy_etot 1E-5
sc_accuracy_rho 1E-4
sc_iter_limit 100
```

**Units:**

Positions in Å

Energies in eV

Manual, chap. 2.1

## Basic electronic structure with FHI-aims

## FHI-aims: input files

**geometry.in**

## # Atomic structure

# x y z

atom 0.0 0.0 0.0 H

atom 1.0 0.0 0.0 H

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**control.in**

## # Physical model settings

xc pw-lda

charge 0.

spin collinear

## # SCF convergence settings

sc\_accuracy\_eev 1E-2

sc\_accuracy\_etot 1E-5

sc\_accuracy\_rho 1E-4

sc\_iter\_limit 100

## # Species specifics

...

**Units:**

Positions in Å

Energies in eV

Manual, chap. 2.1

Basic electronic structure with FHI-aims

# species\_default

```
$HANDSON/aimsfiles/species_defaults
```

**Predefined species**

**Copy-paste into control.in**

**Manual, chap. 2.2**

- light
- tight
- really tight

# species\_default

```
$HANDSON/aimsfiles/species_defaults
```

**Predefined species**

**Copy-paste into control.in**

**Manual, chap. 2.2**

- **light**

**Increased accuracy:**

- **tight**

Basis  
Hartree potential

- **really tight**

Basis cutoff potential  
Integration grids

# species\_default

```
$HANDSON/aimsfiles/species_defaults
```

Predefined species

Copy-paste into control.in

Manual, chap. 2.2

- light
- tight
- really tight



- Fast, many production tasks**  
Fast pre-relaxation
- Used to verify important results**  
Converged settings
- Heavily converged numerical settings**  
Explicit convergence tests

# species\_default

```
$HANDSON/aimsfiles/species_defaults
```

Predefined species

Manual, chap. 2.2

Copy-paste into control.in

- light
- tight
- really tight



- Fast, many production tasks**  
Fast pre-relaxation
- Used to verify important results**  
Converged settings
- Heavily converged numerical settings**  
Explicit convergence tests

**Additionally converge basis ("tiers")!**

Basic electronic structure with FHI-aims

# FHI-aims output

1

Invoking FHI-aims ...

Introduction

# FHI-aims output

1

Invoking FHI-aims ...

---

2

Reading file control.in.

---

Summary of control.in file

## Basic electronic structure with FHI-aims

# FHI-aims output

1

Invoking FHI-aims ...

2

Reading file control.in.

3

Reading geometry description geometry.in.

Summary of geometry.in file

# FHI-aims output

1

Invoking FHI-aims ...

2

Reading file control.in.

3

Reading geometry description geometry.in.

4

Preparing all fixed parts of the calculation.

Geometry independent preparations  
Basis set generation

## Basic electronic structure with FHI-aims

# FHI-aims output

5

```
-----  
Begin self-consistency loop: Initialization.  
Date : 20130610, Time : 162002.389  
-----
```

Geometry dependent preparations  
Integration grid  
Initialization of charge density

## Basic electronic structure with FHI-aims

## FHI-aims output

5

```
-----  
Begin self-consistency loop: Initialization.  
Date : 20130610, Time : 162002.389  
-----
```

6

```
-----  
Begin self-consistency iteration # 1  
Date : 20130610, Time : 162002.445  
-----
```

First SCF cycle

## Basic electronic structure with FHI-aims

## FHI-aims output

5

```
-----  
Begin self-consistency loop: Initialization.  
Date : 20130610, Time : 162002.389  
-----
```

6

```
-----  
Begin self-consistency iteration # 1  
Date : 20130610, Time : 162002.445  
-----
```

First SCF cycle

THIS  
TUTORIAL

## » Energy

Total energy	:	-13.01991124 eV
Total energy, T → 0	:	-13.01991124 eV
Electronic free energy	:	-13.01991124 eV

Periodic metals only

## Basic electronic structure with FHI-aims

# FHI-aims output

5

```
-----  
Begin self-consistency loop: Initialization.  
Date : 20130610, Time : 162002.389  
-----
```

6

```
-----  
Begin self-consistency iteration # 1  
Date : 20130610, Time : 162002.445  
-----
```

First SCF cycle

## » Self-consistency convergence accuracy

Change of charge density : 0.6753E-02
Change of sum of eigenvalues : 0.4376E+00 eV
Change of total energy : 0.1143E-01 eV

## Basic electronic structure with FHI-aims

# FHI-aims output

5

```
-----  
Begin self-consistency loop: Initialization.  
Date : 20130610, Time : 162002.389  
-----
```

6

```
-----  
Begin self-consistency iteration # 6  
Date : 20130610, Time : 162002.560  
-----
```

Sixth SCF cycle

## » Self-consistency convergence accuracy

- | Change of charge density : 0.3163E-05
- | Change of sum of eigenvalues : -.9415E-05 eV
- | Change of total energy : 0.2388E-10 eV

Basic electronic structure with FHI-aims

# FHI-aims output

7

Self-consistency cycle converged.

# FHI-aims output

7

Self-consistency cycle converged.

## » Energy and forces

| Total energy uncorrected : -0.130198526094581E+02 eV  
~~| Total energy corrected : -0.130198526094581E+02 eV~~  
| Electronic free energy : -0.130198526094581E+02 eV

## » SCF info

| Number of self-consistency cycles : 6

## » Timings

## Basic electronic structure with FHI-aims

# FHI-aims output

7

Self-consistency cycle converged.

## » Energy and forces

| Total energy uncorrected : -0.130198526094581E+02 eV  
~~| Total energy corrected : -0.130198526094581E+02 eV~~  
| Electronic free energy : -0.130198526094581E+02 eV

## » SCF info

| Number of self-consistency cycles : 6

## » Timings

8

Have a nice day.

# FHI-aims output

7

Self-consistency cycle converged.

Postprocessing

Structure optimization

- » Get next relaxation step
- » Redo SCF for new geometry

8

Have a nice day.

## Problem II: Hydrofluoric acid (HF)

One of the first papers which systematically investigated the performance of DFT was published by John A. Pople and coworker in 1993.

### Tasks:

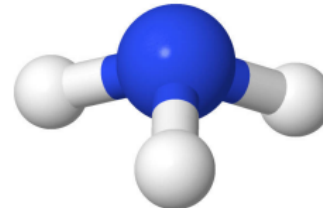
- Find the equilibrium bond distance of HF.
- Compare the HF bond length for different methods.
- Calculate the atomization energy ( $\Delta H_{at}$ ).
- Compute the dipole moment for different methods and bond lengths.

Local structure optimisation

# Problem IV to VIII: Hydronium cation ( $\text{H}_3\text{O}^+$ )

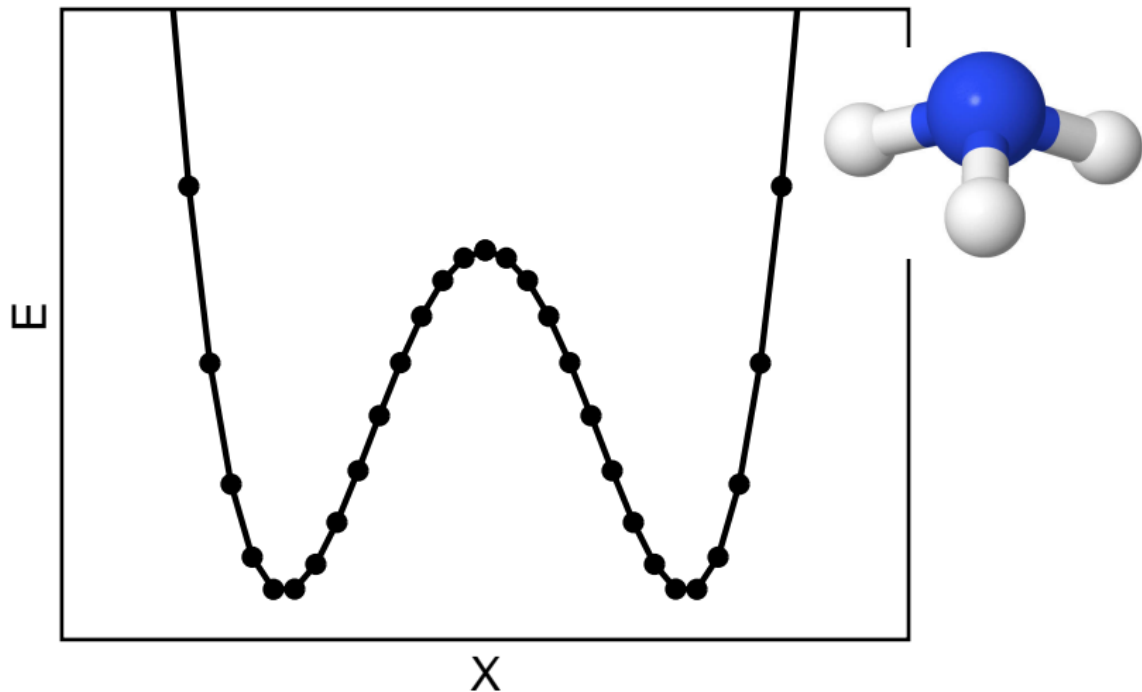
## Tasks:

- Relax structure with two different starting points.
- Make a vibrational analysis.



Local structure optimisation

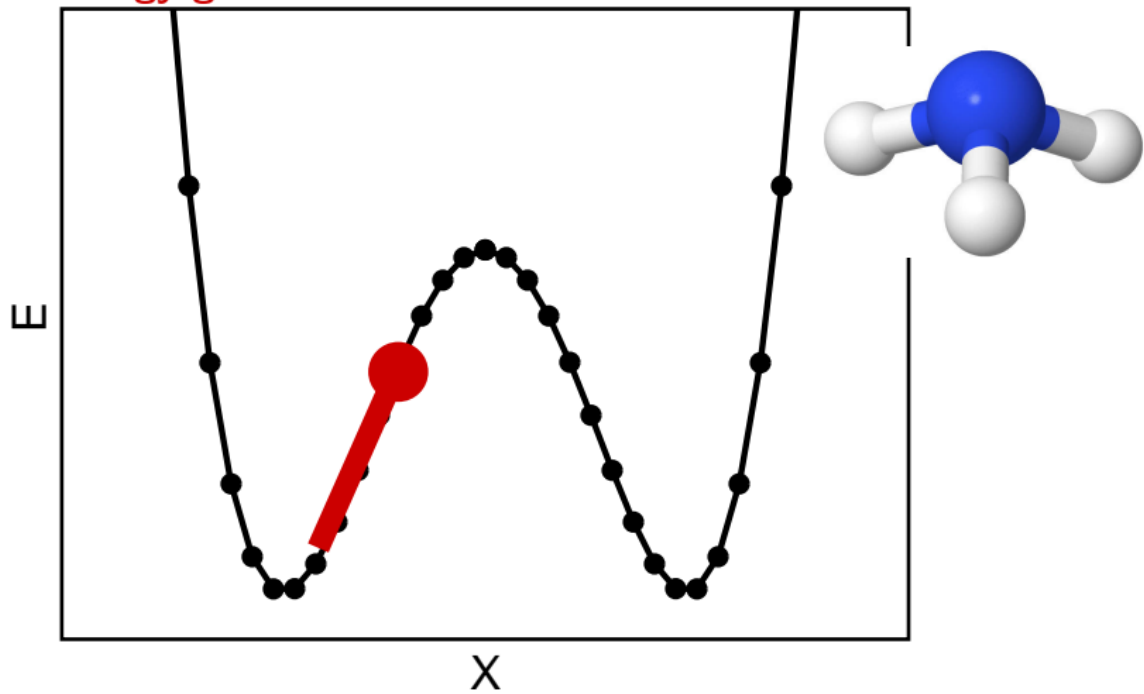
# Forces



Local structure optimisation

## Forces

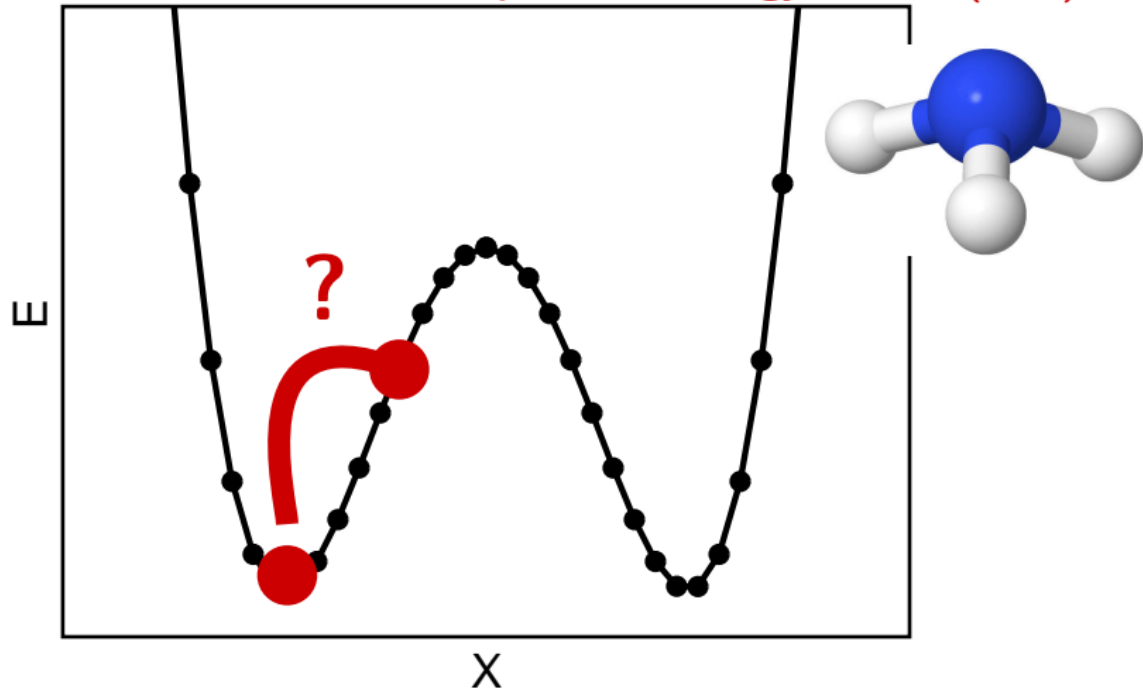
## Energy gradient



Local structure optimisation

# Forces

**Structure optimization:**  
Find local minimum on potential energy surface (PES)



Local structure optimisation

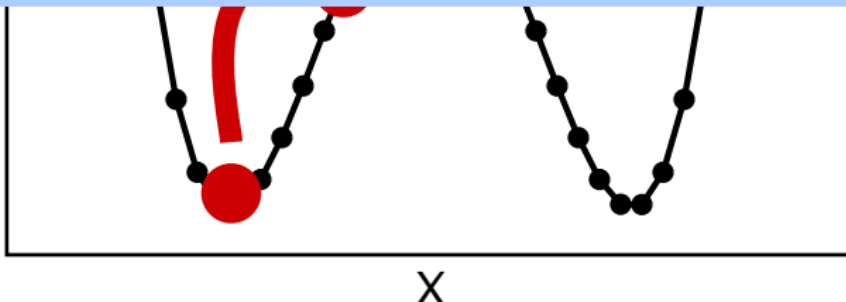
# Forces

**Structure optimization:**  
**Find local minimum on potential energy surface (PES)**



Many methods !

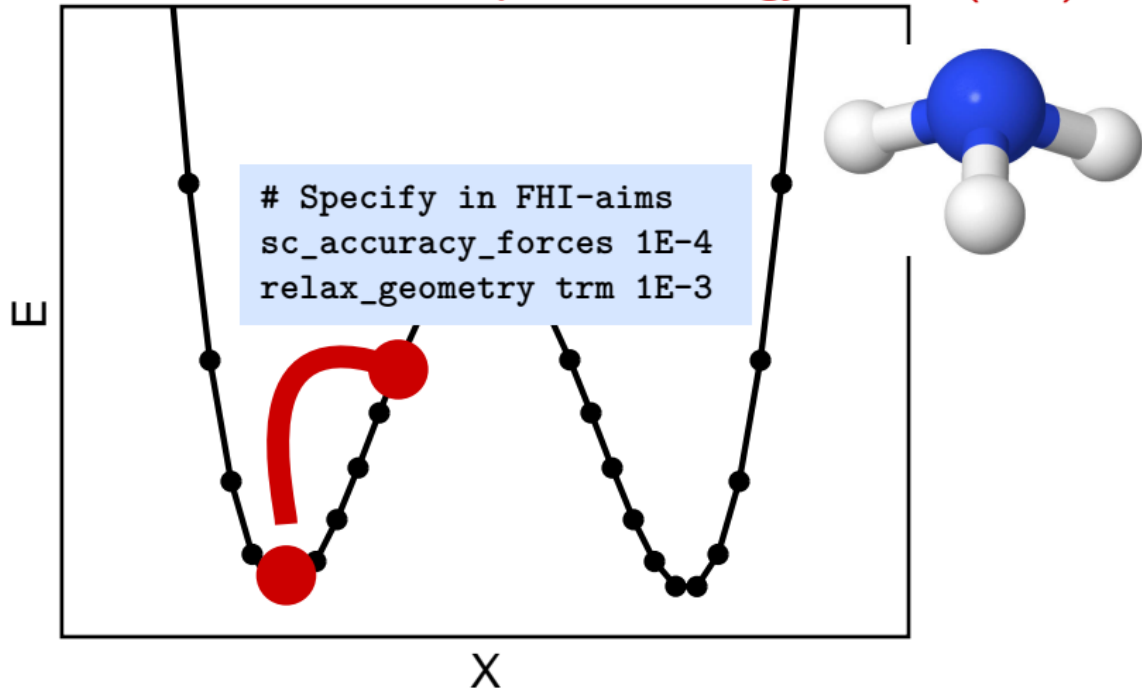
Industry standard: quasi Newton methods



## Local structure optimisation

## Forces

Structure optimization:  
Find local minimum on potential energy surface (PES)



Local structure optimisation

# Harmonic molecular motion

**How do atoms move in a potential  $V$ ?**

⇒ Solve equations of motion!

$$\frac{d}{dt} \frac{\partial T}{\partial \dot{R}_i} + \frac{\partial V}{\partial R_i} = 0$$

Kinetic energy:  $T$

Potential energy:  $V$

Local structure optimisation

# Harmonic molecular motion

**How do atoms move in a potential  $V$ ?**

⇒ Solve equations of motion!

$$\frac{d}{dt} \frac{\partial T}{\partial \dot{R}_i} + \frac{\partial V}{\partial R_i} = 0$$

Kinetic energy:  $T$

Potential energy:  $V$

⇒ Taylor expansion of  $V$

around equilibrium position  $R_0$  + harmonic approximation

$$V = V_0 - \underbrace{F(R_0)R}_{=0 \text{ equilibrium}} + \frac{1}{2} R^T H(R_0) R + \underbrace{\dots}_{=0 \text{ harmonic approximation}} \text{higher terms}$$

$F$ : Forces

$H$ : Hessian

Local structure optimisation

# Harmonic molecular motion

**How do atoms move in a potential  $V$ ?**

⇒ Solve equations of motion!

$$\frac{d}{dt} \frac{\partial T}{\partial \dot{R}_i} + \frac{\partial V}{\partial R_i} = 0$$

Kinetic energy:  $T$

Potential energy:  $V$

⇒ Solution

The Dynamic Matrix  $D_{ij}$ :  $D_{ij} = \frac{1}{\sqrt{M_i}\sqrt{M_j}} H_{ij}$

$$R \sim Q e^{i\omega t}, \quad \text{with} \quad DQ - \omega^2 Q = 0$$

Eigenmodes  $Q$

- If (1) Harmonic approximation is valid  
(2) Equilibrium geometry

Local structure optimisation

# Vibrations

**Solve**  $\mathbb{R}^{3N} \times \mathbb{R}^{3N}$  eigenvalue equation

$$\det(D - \omega^2 \mathbf{1}) = 0$$

**Hessian  $H$**       **Dynamic matrix  $D$**

$$H^{ij} := \frac{\partial^2 E}{\partial R_i \partial R_j} \quad D_{ij} = \frac{1}{\sqrt{M_i} \sqrt{M_j}} H_{ij}$$

**In practice:** finite central numerical differences (of forces)

**Wrapper**

```
> aims_vibrations.mpi.pl
```

**Manual, chap 4.6**

Local structure optimisation

# Vibrations

**Solve**  $\mathbb{R}^{3N} \times \mathbb{R}^{3N}$  eigenvalue equation

$$\det(D - \omega^2 \mathbf{1}) = 0$$

**Get**

- » Eigenmodes  $\{Q_i, i \in 1 \dots 3N\}$
- » Eigenfrequencies

Local structure optimisation

# Vibrations

**Solve**  $\mathbb{R}^{3N} \times \mathbb{R}^{3N}$  eigenvalue equation

$$\det(D - \omega^2 \mathbf{1}) = 0$$

## Get

- » **Eigenmodes**  $\{Q_i, i \in 1 \dots 3N\}$
- » **Eigenfrequencies**
- » **6 (almost) zero frequency modes** (if molecule non-linear)  
**translations + rotations**
- » **Imaginary frequency**  $\Rightarrow$  Saddle point

Local structure optimisation

# Vibrations

**Solve**  $\mathbb{R}^{3N} \times \mathbb{R}^{3N}$  eigenvalue equation

$$\det(D - \omega^2 \mathbf{1}) = 0$$

## Get

- » **Eigenmodes**  $\{Q_i, i \in 1 \dots 3N\}$
- » **Eigenfrequencies**
- » **6 (almost) zero frequency modes** (if molecule non-linear)  
**translations + rotations**
- » **Imaginary frequency**  $\Rightarrow$  Saddle point
- » **Infrared intensities** (derivative of dipole moment  $\mu$ )

$$I_i \sim \left| \frac{d\mu}{dQ_i} \right|^2$$

Local structure optimisation

# Vibrations

Based on harmonic approximation !  
Beyond: Tutorial 4 (MD) on Friday

## Get

- » **Eigenmodes**  $\{Q_i, i \in 1 \dots 3N\}$
- » **Eigenfrequencies**
- » **6 (almost) zero frequency modes** (if molecule non-linear)  
**translations + rotations**
- » **Imaginary frequency**  $\Rightarrow$  Saddle point
- » **Infrared intensities** (derivative of dipole moment  $\mu$ )

$$I_i \sim \left| \frac{d\mu}{dQ_i} \right|^2$$

## Visualising density differences

# Visualization

## Orbitals and densities

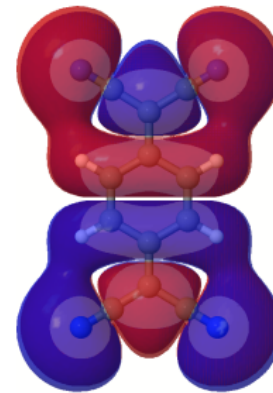
**Keyword in control.in**

```
output cube eigenstate homo
cube filename HOMO(cube)
output cube total_density
cube filename tot_dens_uc(cube)
```

**Get: \*.cube file** - values on a regular 3D grid.

**Software: molden** (jmol, gdis, xcrysden)

⇒ Appendix of handout



# Practical issues

- **Each calculation one directory**

```
> mkdir tutorial1  
> cd tutorial1  
> mkdir HF
```

- **2 input files**

```
geometry.in  
control.in
```

- **Launching FHI-aims calculation**

```
aims.x | tee aims.out
```

- ... scripting helps !

(Sample scripts in appendix of handout)

# Your Tutors for the afternoon



Oliver T. Hofmann



Lydia Nemec



Christian Carbogno



William P. Huhn



Volker Blum



Markus Schneider

