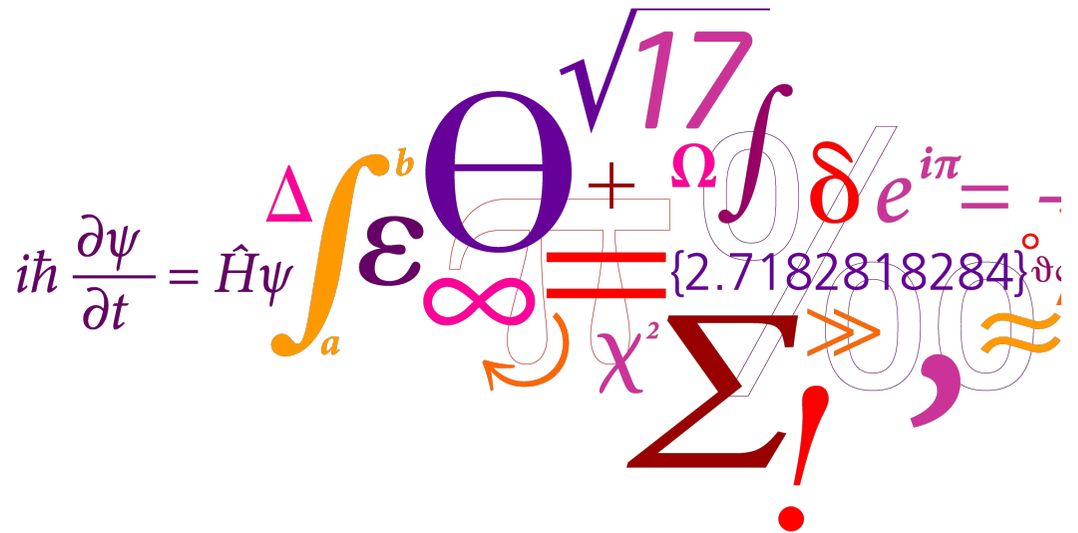


# Electronic structure tools: ASE and GPAW and introduction to the computer projects

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# Overview



- The Python language
- The Atomic Simulation Environment (ASE)
  - The anatomy of an atomic-scale simulation/calculation
  - The ASE
- Examples
  - Almost the simplest possible molecular dynamics simulation.
  - Almost the simplest possible GPAW calculation.
- Using the DTU “databar” (computer lab).
- Computational projects

Part I

# **BRIEF INTRODUCTION TO PYTHON**

# Why Python?



- For the programmer: Python is object-oriented
  - Object oriented and modular: Facilitates writing and maintaining complex problems.
  - Dynamically typed: Flexibility, facilitates code reuse.
  - Easy to write readable code: Code is maintainable.
  - Large libraries available (numerics, plotting, ...)
- For the user: Python is a scripting language.
  - Great for scripting a calculation
  - Great for small programs and prototypes.
  - Great for interactive experimenting.
  - Easy to learn.
  - Objects are powerful in scripts!
- Python can be extended in C/C++/Fortran
  - Solves performance problems of non-compiled languages.

# Learning Python



- Don't waste money on Python books!
  - It's not that hard, and online docs are good.
- Python tutorial: <http://docs.python.org/3/tutorial/>
  - More documentation at docs.python.org
- Learn Python + ASE + GPAW by example
  - Get a simple script, and modify it.
  - Simple scripts are almost like old-fashioned input files!
- Get Python from [www.python.org](http://www.python.org)
  - For Windows (only) get Anaconda.

# A few places where Python is different

- Blocks are marked by indentation, not by keywords or braces

```
n = 0
for i in range(10):
    n += i
    print(i, n)
print('After the loop')
```

- Loops are over an “iterable” (e.g. a list)

```
for i in ['apple', 'tomato', 'orange']:
    print(i)
```

# A few places where Python is different

- The = operator is naming objects, not assigning to variables!

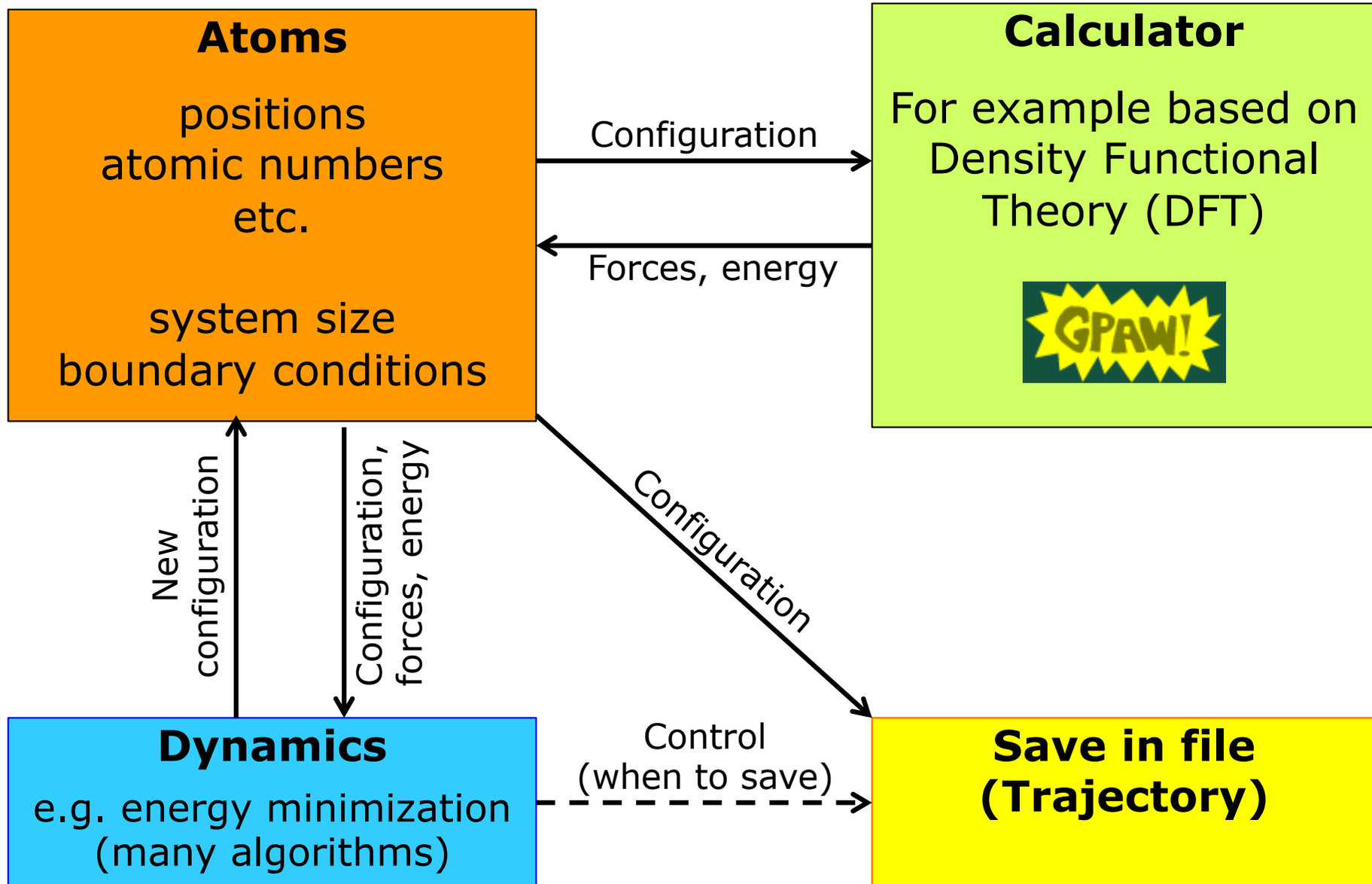
```
a = [1, 2, 3, 4, 5] # Create a list
b = a              # New name for list
a[2] = 42
print(b)          # [1, 2, 42, 4, 5]
```

```
c = 7
d = c
c += 42          # d is still 7, we just did
                 # c = c + 42
                 # creating a new object 49 and
                 # giving it the name c
```

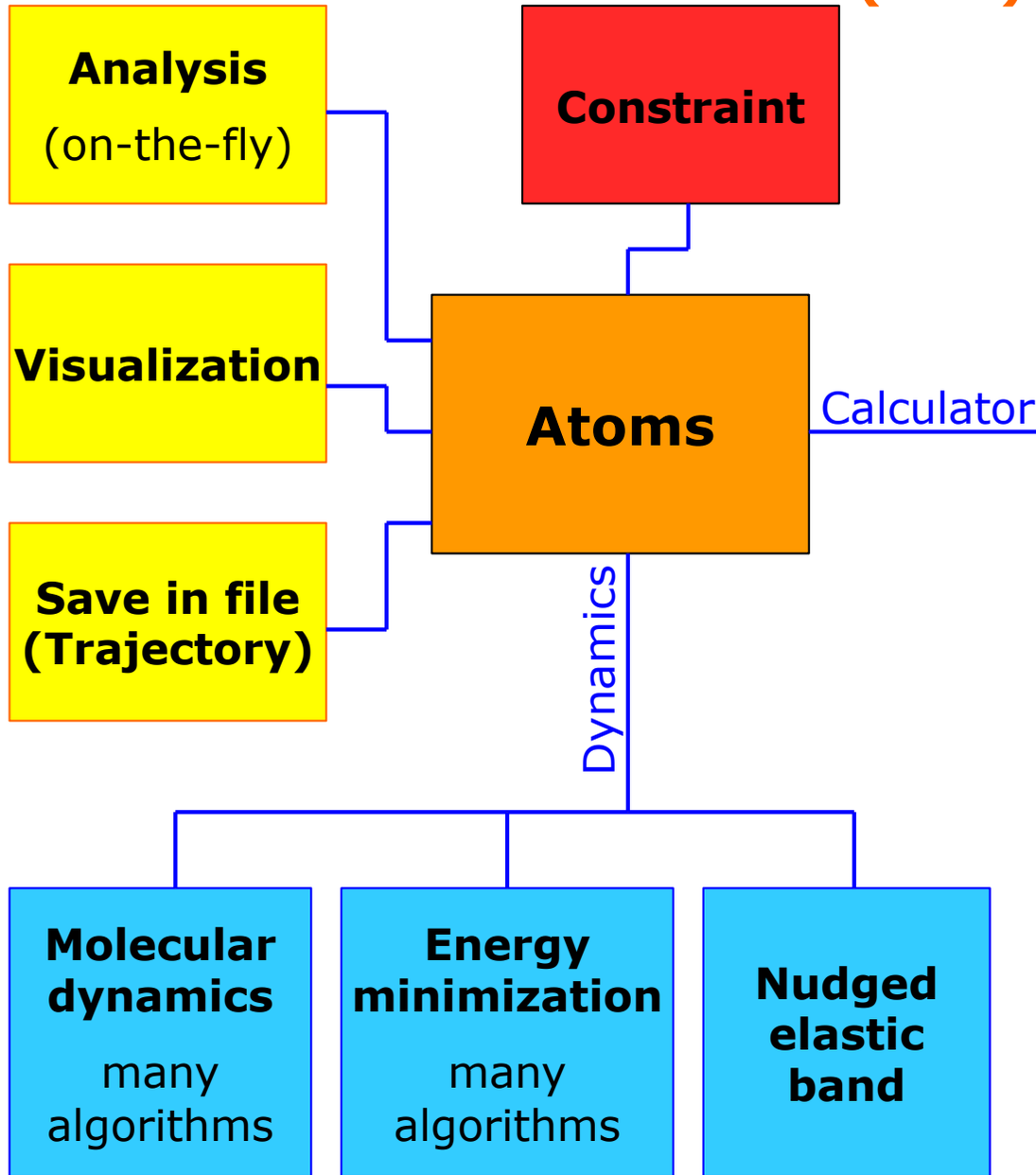
Part II

# **INTRODUCTION TO ASE AND GPAW**

# The "anatomy" of an atomic-scale computer simulation



# Atomic Simulation Environment (ASE)



**Density Functional Theory (DFT)**  


**DFT**  
  
**DFT**  


**DFT**  
  
**ELK**

And many other...

**Classical Potentials**  
**ASAP**

**Classical Potentials**  
**LAMMPS**

And some other...

# Example: Molecular dynamics



```
import numpy as np
from ase import Atoms, units
from ase.io.trajectory import Trajectory
from ase.build import bulk
from ase.md.verlet import VelocityVerlet
from asap3 import EMT

# Create the atoms
atoms = bulk("Cu", "fcc", cubic=1).repeat([3,3,3])
atoms.set_pbc(False)
atoms.center(vacuum=5.0)
# Give the first atom a non-zero momentum
atoms[0].momentum = np.array([0, -11.3, 0])

# Prepare to do molecular dynamics, forces described by EMT
atoms.calc = EMT()
dyn = VelocityVerlet(atoms, 5.0*units.fs)

# Make a trajectory writing output every fifth timestep.
trajectory = Trajectory("MD-output.traj", "w", atoms)
dyn.attach(trajectory, interval=5)

# Now do 1000 timesteps.
dyn.run(1000)
```

# GPAW Example: Atomization energy of Hydrogen (1/2)

```
from ase import Atoms, Atom
from gpaw import GPAW

a = 4. # Size of unit cell (Angstrom)
c = a / 2
# Hydrogen atom:
atom = Atoms('H',
             positions=[(c, c, c)],
             magmoms=[1],
             cell=(a, a, a))

# gpaw calculator:
calc = GPAW(h=0.18, nbands=1, xc='PBE', txt='H.out')
atom.calc = calc

e1 = atom.get_potential_energy()
calc.write('H.gpw')
```

Continued...

# GPAW Example: Atomization energy of Hydrogen (2/2)

Continued ...

```
# Hydrogen molecule:
d = 0.74 # Experimental bond length
molecule = Atoms('H2',
                  positions=([c - d / 2, c, c],
                             [c + d / 2, c, c]),
                  cell=(a, a, a))

calc.set(txt='H2.out')
molecule.calc = calc
e2 = molecule.get_potential_energy()
calc.write('H2.gpw')

print 'hydrogen atom energy:      %5.2f eV' % e1
print 'hydrogen molecule energy: %5.2f eV' % e2
print 'atomization energy:       %5.2f eV' % (2 * e1 - e2)
```

# The Atoms object



The `Atoms` object is the main simulation object. It contains:

- Per-atom data: positions, velocities, charges, magnetic moments, tags.
- Global data: Unit cell, boundary conditions.
- Refs to helper objects: Calculator, constraints, ...

Accessing the data:

`r = atoms.get_positions()` **or** `r = atoms.positions`

`atoms.set_positions(r)` **or** `atoms.positions = r`

`f = atoms.get_forces()` Will trigger a calculation, if needed.

# The Atoms object



## Manipulating the Atoms object:

`atoms.center(vacuum=5.0)` Centers in unit cell, possibly adjusting the amount of vacuum.

`atoms.rotate(45, 'z')` Rotate the atoms

`atoms.repeat([2,2,1])` Replicate atoms along x,y,z axes.

## The Atoms object is a Python sequence:

`atoms[i]` The  $i^{\text{th}}$  atom (starting at zero).

`atoms.extend(atoms2)` Add more atoms

`atoms1 + atoms2` Merge two atoms objects

`for a in atoms:`  
`print(a.position)` Loop over atoms

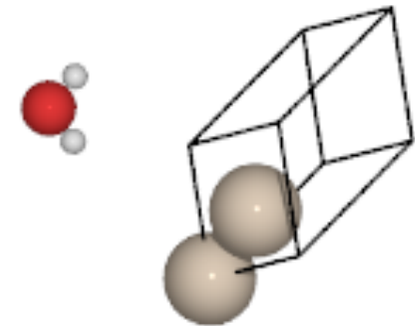
# Making systems

Manually specify the atoms:

```
from ase import Atoms
h2 = Atoms('H2', positions=[(0, 0, 0), (0, 0, 0.74)])
```

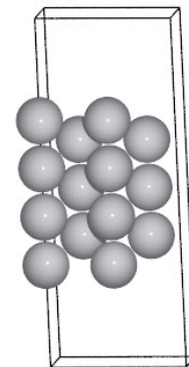
Known molecules and bulk structures:

```
from ase.build import molecule, bulk
water = molecule('H2O')
si2 = bulk('Si', 'diamond', a=5.4)
```



Simple surfaces:

```
from ase.build import fcc110
slab = fcc110('Pt', (2, 1, 7),
              a=4.0, vacuum=6.0)
```



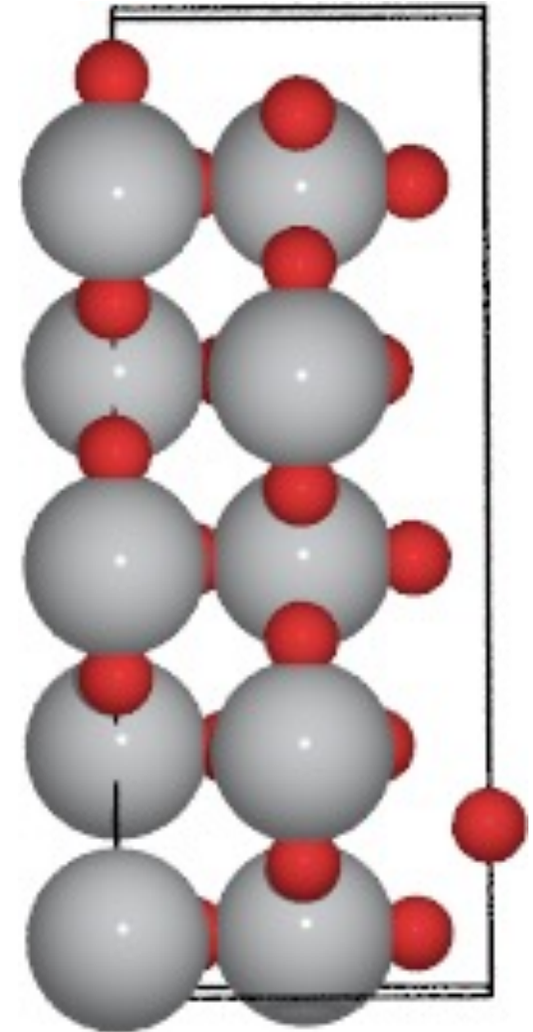
# Making systems (complicated example)

Make a Rutile crystal, then cut a surface from it:

```
from ase.spacegroup import crystal
from ase.build import surface

a, c, u = 4.584, 2.953, 0.3
rutile = crystal(['Ti', 'O'],
                 basis=[(0, 0, 0),
                        (u, u, 0)],
                 spacegroup=136,
                 cellpar=[a, a, c,
                          90, 90, 90])

nasty_cut = surface(rutile,
                    (1, 1, 0),
                    layers=5)
```



# GPAW essentials



- Our main tool for DFT calculations.
- Many advanced DFT features:  
*Spin-orbit coupling, DSCF, Bayesian Error Estimate, external potentials, TDDFT, ...*
- Beyond-DFT features:  
*GW, Random Phase Approximation, ...*

# GPAW essentials



- **Grid mode** – wavefunctions on a real-space grid

```
calc = GPAW(h=0.18, xc='PBE', ...)
```

Parallelizes very well for large systems. High accuracy

- **Plane wave mode** – wavefunctions in k-space

```
calc = GPAW(mode=PW(400), xc=..., ...)
```

Faster than grid mode for small/medium systems. High accuracy

- **LCAO mode** – wavefunctions in real space; with basis fct.

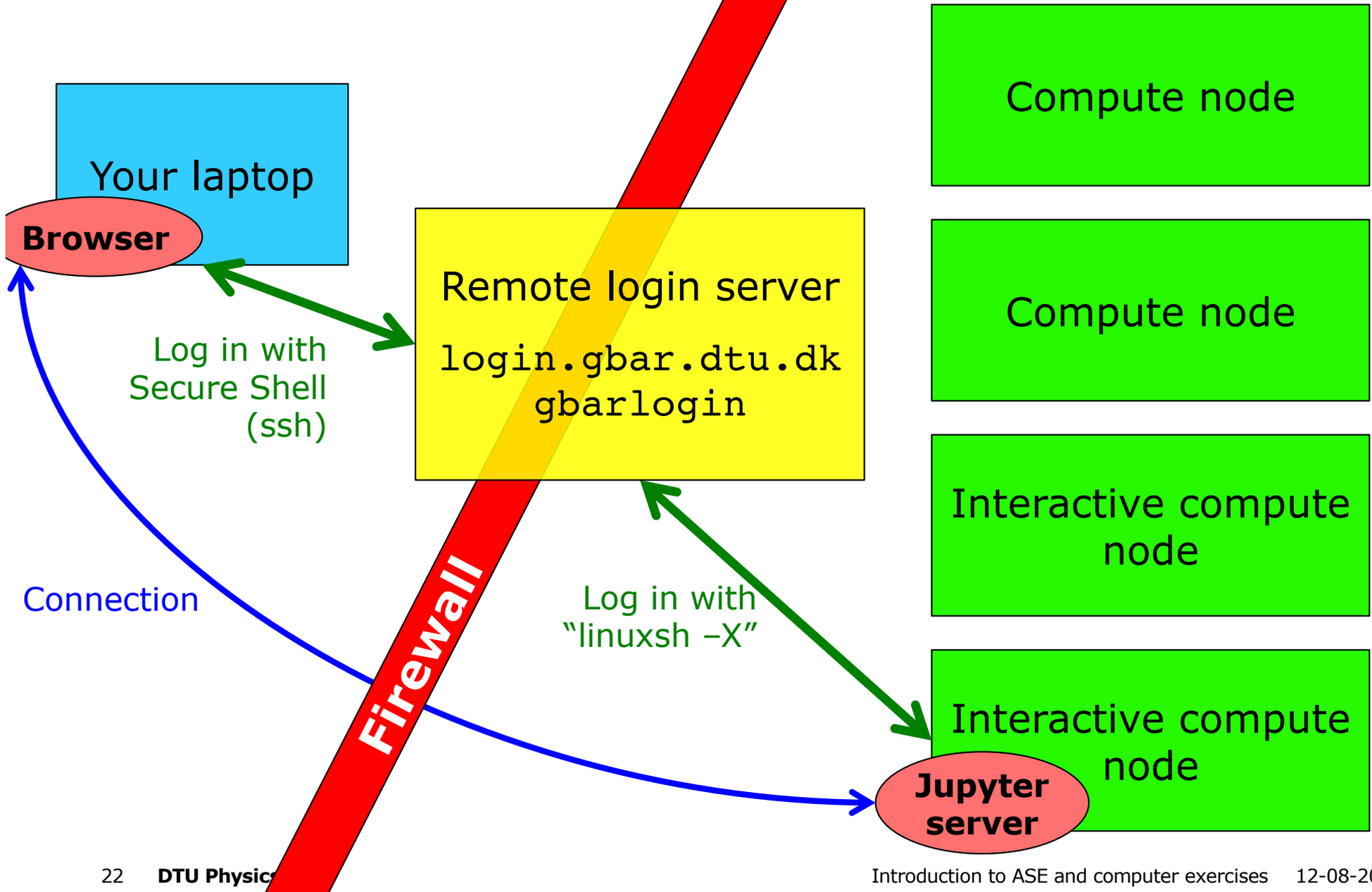
```
calc = GPAW(mode='lcao', basis='dzp', h=0.18, ...)
```

Fast but less accurate.

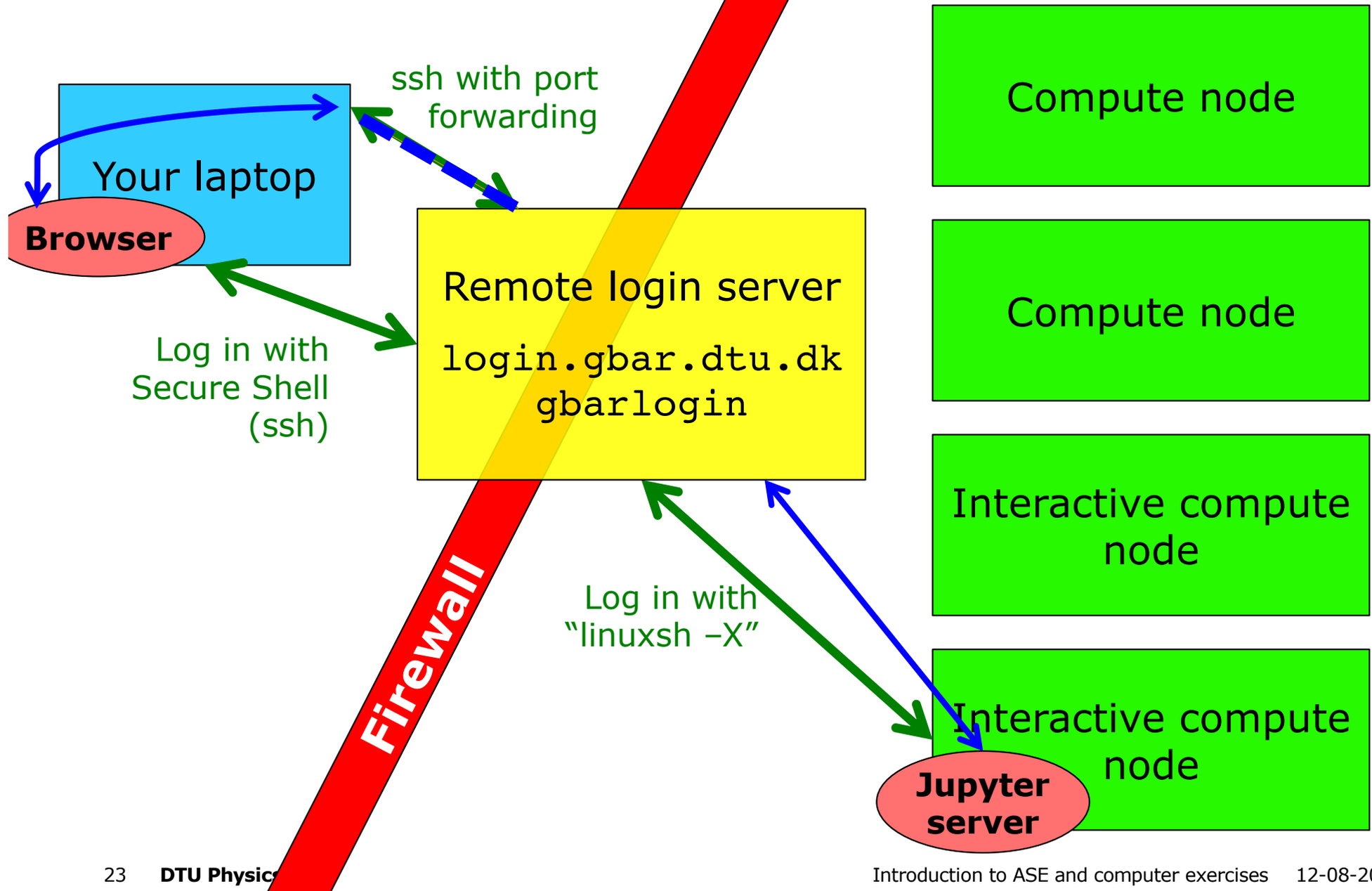
Part III

# COMPUTER EXERCISES AND THE DTU "DATABAR"

# The DTU "databar" (computer labs)



# The DTU "databar" (computer labs)



# The “databar” and the projects



- Information about the Summerschool projects:  
[wiki.fysik.dtu.dk/gpaw/](http://wiki.fysik.dtu.dk/gpaw/)

Includes detailed instructions on how to log in.

- Changing password in the HPC system (“databar”):
  - Change at [password.dtu.dk](http://password.dtu.dk) (takes up to 1 hour to sync)
- **SLIDES** available from the GPAW summerschool pages.

# Projects



- Photovoltaics
  - Mark Kamper Svendsen
- Catalysis
  - Cuauhtémoc Núñez Valencia
- Magnetism
  - Thorbjørn Skovhus
- Batteries
  - Steen Lysgaard
- Machine Learning
  - Benjamin Heckscher Sjølin
- Workflows
  - Ask Hjort Larsen

# Project: Photovoltaics



- Calculate electronic band structures and bandgaps
- Calculate optical absorption
  
- Special methods:
  - The Random Phase approximation
  - Optional: The Bethe-Salpeter Equation

# Project: Catalysis



- DFT calculations of adsorption geometries of molecules on surfaces.
- Calculations of the reaction path and the transition energy
- Special methods:
  - Nudged Elastic Band (for transition paths)
  - Optional: Vibrational analysis for adsorption entropy

# Project: Magnetism in 2D



- Calculations of the critical temperature of a  $\text{CrI}_3$  monolayer
- Calculations of the noncollinear ground state in  $\text{VI}_2$
- Autodiscovery of new magnetic monolayers in the Computational 2D Materials Database (C2DB)
- Special methods:
  - Energy mapping analysis
  - Noncollinear DFT calculations

# Project: Batteries



- Study the anode and cathode materials of a Li-ion battery with DFT
- Calculate the intercalation energy of Li in graphite, establish the equilibrium potential of a LiFePO<sub>4</sub>/C battery and determine important battery characteristics such as Li transport barriers and the voltage profile.
- Special methods:
  - Structure creation and modification with ASE
  - Unit cell relaxation
  - Bayesian error estimation
  - Nudged Elastic Band (NEB) calculations for estimating Li migration barriers

# Project: Machine Learning



- Machine Learning methods used on an example database
- Predicting band gaps and heat of formation using a structure “fingerprint”
- Predictions with several methods (e.g., ridge regression, decision tree, and gaussian process)
- Validation of predictions using DFT

# Project: Workflows with Atomic Simulation Recipes



- Learn to define tasks and workflows using Python
- Use command-line tools to create and manage tasks in a directory tree
- Run a materials workflow on multiple materials