ASE: A programmable environment for calculations with many electronic structure codes

Ask Hjorth Larsen

Center for Atomic-scale Materials Design, Technical University of Denmark

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Atomic Simulation Environment

ASE is a set of tools for atomic-scale simulation tasks.

What can ASE do?

- Set up and view structures
- Run calculations such as structure optimizations and molecular dynamics
- Read and write structures in many file formats
- ASE is open source / free software (LGPLv2.1)
- Many contributors

What is so special about ASE?

- ASE unifies many electronic structure codes within a single environment
- Calculations are written as Python scripts



Motivation

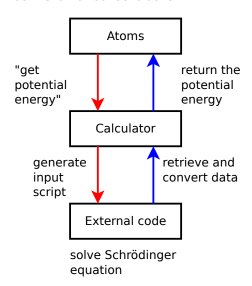
- Increase portability of calculations between codes
- Electronic structure codes usually take different input parameters and have incompatible file formats
- Much functionality is common to all codes: atomic geometries, structure optimization algorithms, . . .

- ASE identifies common functionality and separates it from the specific parameters of different codes
- ► Electronic structure codes are abstracted as calculators. Calculators implement a specific programming interface

A bit of history

- ► ASE was developed as an interface to the FORTRAN-based ultrasoft pseudopotential code Dacapo
- S. R. Bahn and K. W. Jacobsen, An object-oriented scripting interface to a legacy electronic structure code, Comput. Sci. Eng. 4, 56 (2002), ISSN 1521-9615.
- ASE is a part of the CAMPOS (CAMP Open Source) project https://wiki.fysik.dtu.dk/
- Other CAMPOS codes have been written with ASE interfaces: ASAP and GPAW
- ASE development is driven in part by our own requirements, so ASE/GPAW is a strongly correlated system

Atoms and calculators



- Atoms objects represent geometry: positions, atomic numbers, unit cell, momenta, ...
- Calculators can be based on anything that calculates energies, forces (DFT, classical potentials, ...)

Atoms and calculators

Example: The ASE/GPAW "hello world"

ASE calculators so far



Supported calculators

Grid-based real-space PAW code	DFT, HF
Highly efficient EMT code $(C++)$	EMT
planewave USPP code	DFT
Effective Medium Theory calculator	EMT
A planewave pseudopotential code	DFT
LCAO pseudopotential code	DFT
DftbPlus DFT based tight binding	DFT
Fast atomic orbital code	DFT, HF
Planewave pseodopotential code	DFT, HF
Planewave PAW code	DFT
NAO full-potential code	DFT, HF
Full Potential LAPW code	DFT, LAPW
Full Potential LAPW code	DFT, LAPW
Classical molecular dynamics code	
	Highly efficient EMT code (C++) planewave USPP code Effective Medium Theory calculator A planewave pseudopotential code LCAO pseudopotential code DftbPlus DFT based tight binding Fast atomic orbital code Planewave pseodopotential code Planewave PAW code NAO full-potential code Full Potential LAPW code Full Potential LAPW code

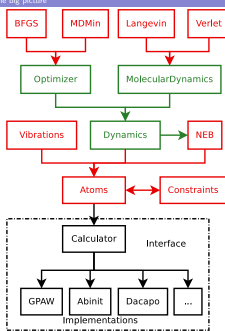
Scripting with Python in ASE



- Access to a complete language provides very high flexibility
- Python is interpreted, so no compilation required
- Well suited for scripting due to high-level features and simple syntax
- Cross-platform (although some supercomputers can be finicky)

Scientific programming in Python

- numpy: module for array calculations
- scipy: module for high-level mathematics with numpy (optimization, integration, root finding)
- matplotlib: module for plotting



Overview of ASE classes

- Only the most common objects shown
- Red: user interface Green: classes for internal use

The big picture

Demonstration: structure optimization

Structure generation

Molecules

▶ ASE includes G1, G2 and G3 datasets

```
from ase.data.molecules import molecule
system = molecule('CH3CH2OH')
```

Crystal structures

- Bulk
- Surfaces
- Clusters

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► ASE supports a couple dozen file formats

Demonstration: Gold 111 surface

Bulk rutile

Demonstration: Pt cluster with GUI

Future plans or work in progress

Electronic structure interface

- Get density of states
- Export wavefunctions, densities (basis set conversions)
- More uniform input parameters

Automation

- Bulk submit script generation?
- "Run this calculator on this set of structures"

And more

- Interface to deal with electronic excitations
- Debian packages for Debian Open Science
- "Scratch your own itch"



List of committers

Andrew Peterson David Landis Lars Grabow Jakob Blomquist Jens J Mortensen John Kitchin Jesper Kleis Michael Walter Christian Glinsvad Mikkel Strange Anthony Goodrow Troels K Jacobsen

Ask Hjorth Larsen Marcin Dulak Heine Anton Hansen Jon B. Maronsson Jesper Friis Jussi Enkovaara Karsten W Jacobsen Poul Georg Moses Jakob Schiotz Tao Jiang Carsten Rostgaard

Jonas Bjork George Tritsaris Felix Hanke Janne Blomqvist Jingzhe Chen Kristen Kaasbjerg Markus Kaukonen Marco Vanin Mattias Slabanja Thomas Olsen Janosch M Rauba

Conclusion

- Pluggable calculators
- Python scripting
- Object oriented interface
- Structure generation

Links and info

- CAMPOS: https://wiki.fysik.dtu.dk/
- ASE: https://wiki.fysik.dtu.dk/ase/
- Mailing list: ase-users@listserv.fysik.dtu.dk
- IRC: #gpaw on irc.freenode.net (some of the same people)
- Python: http://www.python.org/