

2. VASP - Basics

Weine Olovsson

National Supercomputer Centre (NSC), Linköping University

SNIC-PRACE training, online @NSC 19-20th Oct 2020

VASP - Best Practices Workshop



<https://www.nsc.liu.se/>

<https://www.snic.se/>

<https://training.prace-ri.eu/>



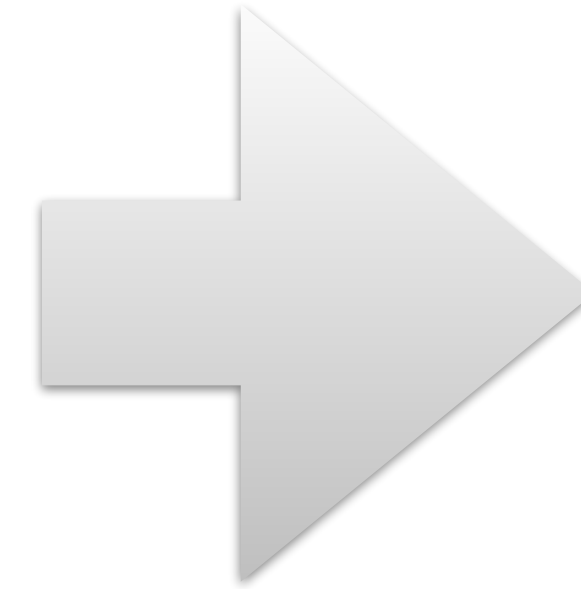
Introduction

- Where to find information
 - VASP at different SNIC HPC centers (examples)
- Starting files
- Important parameters
- Input/output
- Examples

... clickable links are [underlined](#)

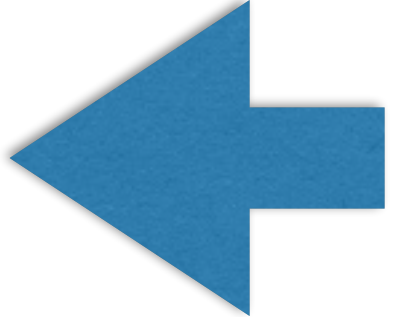
Short background

- Software license
- PAW-method
- DFT, **post-DFT** (HSE06, GW, ...)
- Born-Oppenheimer Molecular Dynamics
- **widely used** in Academia/Industry
 - Efforts from **Intel** & **Nvidia** for optimization
- 20-25% of Tetralith usage
- **VASP6** is available since 2020



<http://vasp.at/>

Starting advice

- Read the [documentation!](#)
- VASP default settings  good starting point
- Caution: “inherited” input files
- Avoid overly messy INCAR...
- Possible differences in installations & versions
refer to respective center webpages / documentation

Resources

- Wiki / Manual

Check in detail!

- Wiki
examples,
presentations
- Forum

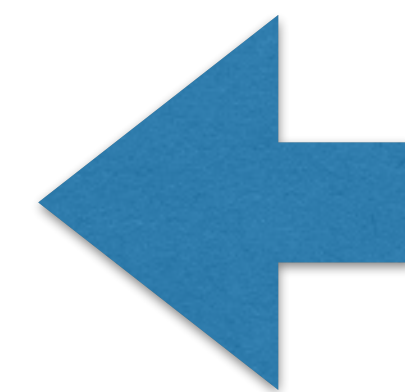


Find the links:

<http://vasp.at/>

- Peter Larsson's (old) blog at NSC:

<https://www.nsc.liu.se/~pla/>

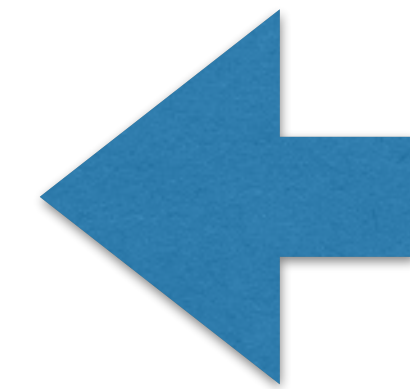


info &
discussion

Questions / trouble? support@nsc.liu.se, ...

VASP at SNIC HPC centers

- **Tetralith** / Sigma, NSC, LiU



this course

- Kebnekaise, HPC2N, UmU

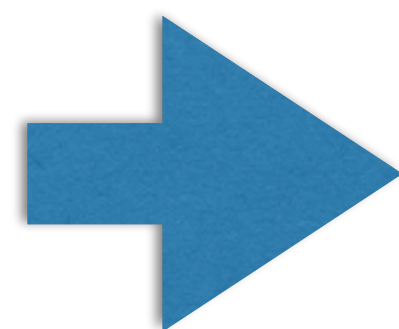
- Beskow, PDC, KTH

- Also available at other systems/centers

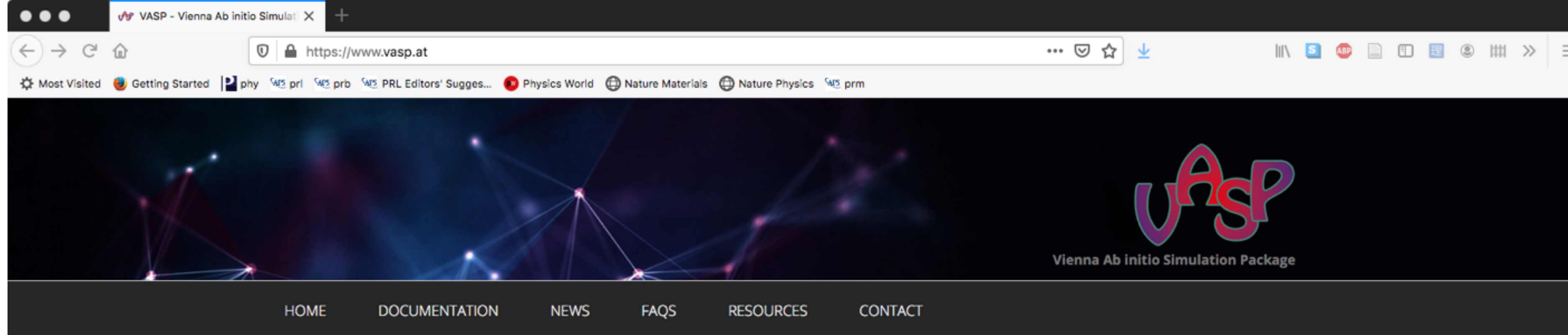
- **On many systems:** \$ module avail vasp
\$ module spider vasp

VASP versions & utilities

- **Latest:** 6.1.0 (6.1.1 to be installed)
- Check centre [webpages](#) for details!
- [wannier90](#): maximally localized wannier functions
- [VTST](#): transition state tools for VASP
- [VASPsol](#): solvation model for VASP
- [Beef](#): Bayesian error estimation functionals
- constrained relaxation



...more about utilities tomorrow



The Vienna Ab initio Simulation Package: atomic scale materials modelling from first principles.

[READ MORE](#)

****Important bug in the testsuite of VASP6 affecting BSD and MacOS: please read this item!****

GET A LICENSE

Not a license holder yet? [Apply for a license here!](#)
Unsure how to proceed? Have a look at the [FAQs](#).

THE VASP WIKI

All documentation: the user manual, tutorials, examples, and lecture notes you will find on our wiki.

VASP SUPPORT FORUM

Our main support channel is a forum. Everybody is welcome to have a look, but posting questions and answers is reserved for licensed user only.

VASP NEWS

New features and releases, bugfixes, highlighted papers, and more.

THE VASP TEAM

Who we are and how we may be reached.

THE VASP PORTAL

Licensees please login here:

Username
Password

[LOGIN](#)

[Forgot my password/username](#)

Latest News

- [Important Notice CoViD-19](#)

Latest Bugfixes

- [Bugfix: testsuite in VASP.6.1.0 to 6.1.2 on](#)

Latest Highlights

- [NEW RELEASE: PAW datasets v.54!](#)

<https://www.vasp.at/>

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Next: [Introduction](#) [Contents](#) [Index](#)

N.B. This document is no longer maintained, please visit our [wiki](#).



The VASP Manual - Vaspwiki

https://cms.mpi.univie.ac.at/wiki/index.php/The_VASP_Manual

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page discussion view source history

All requests for technical support from the VASP group must be addressed to: vasp.materialphysik@univie.ac.at

The VASP Manual

Contents [hide]

- 1 Getting started
- 2 Input and Output
- 3 Featured topics
- 4 Support

Getting started

How to Install VASP	First install VASP.
Workshops	The collection of workshops is a good place for the introduction to the basics of VASP.
Lectures	The collection of lectures is a good place to start as well.
Tutorials and Examples	The collection of tutorials and examples is a good place to learn the usage of VASP.

Input and Output

INCAR tags	All INCAR tags at a glance.
Input Files	
Output Files	

navigation

- [Main page](#)
- [Recent changes](#)
- [Random page](#)
- [Help](#)


search

Search

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tools

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


https://cms.mpi.univie.ac.at/wiki/index.php/The_VASP_Manual

Featured topics

Category	<i>subtopics (amongst others)</i>
Electronic Minimization	Davidson, RMM-DIIS, Conjugate-gradient, preconditioning, mixing.
XC Functionals	LDA, GGA, metaGGA, HF/DFT hybrid functionals, vdW functionals.
Structural Optimization	Ionic minimization methods.
Molecular Dynamics	Barostats, thermostats, ensembles, different MD methods.
Lattice Vibrations	Phonons, electron-phonon interactions.
Dielectric Properties	Static and frequency dependent dielectric properties, Berry phases, spectroscopy (UV, VIS, X-ray).
Linear Response	Static dielectric properties, phonons form linear response.
NMR	Chemical shifts, electric field gradients.
Many-Body Perturbation Theory	ACFDT, BSE, GW, MP2, CRPA.
Magnetism	Spin-orbit coupling, non-collinear magnetism.
Defects	Dipole corrections for defects in solids.
Atoms and Molecules	Monopole, dipole, and quadrupole corrections.
Transition States	Elastic band method, improved dimer method.
Wannier Functions	Interfacing to Wannier90.
Calculational issues	Most important parameters, typical sources of errors.
Theoretical Background	DFT, PAW.
VASP.6 / VASP6	Features that will only be available in VASP.6.X.

Support

If you have questions or run into trouble, please have a look and/or post a question at the [VASP forum](#) 

Please note: we offer support on a courtesy base only, not as a contractual service.

[Back to the top](#)

https://cms.mpi.univie.ac.at/wiki/index.php/The_VASP_Manual

Category:Examples - Vaspwiki

https://cms.mpi.univie.ac.at/wiki/index.php/Category:Examples

Most Visited Getting Started phy prl prb PRL Editors' Sugges... Physics World Nature Materials Nature Physics prm

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category discussion view source history

All requests for technical support from the VASP group must be addressed to: vasp.materialphysik@univie.ac.at

Category:Examples

All articles related to VASP example calculations

Contents

Pages in category "Examples"

The following 81 pages are in this category, out of 81 total.

A

- Adsorption of H2O on TiO2
- Alpha-AIF3
- Alpha-SiO2
- At and mol further

B

- Bandgap of Si in GW
- Bandgap of Si using different DFT+HF methods
- Bandstructure and CRPA of SrVO3
- Bandstructure of Si in GW (VASP2WANNIER90)
- Bandstructure of SrVO3 in GW
- Beta-tin Si

C

- Calculate U for LSDA+U
- Cd Si
- Cd Si relaxation
- Cd Si volume relaxation
- CO

E cont.

- Estimation of J magnetic coupling

F

- Fcc Ni
- Fcc Ni (revisited)
- Fcc Ni DOS
- Fcc Ni DOS with hybrid functional
- Fcc Si
- Fcc Si bandstructure
- Fcc Si DOS

G

- Graphite interlayer distance
- Graphite MBD binding energy
- Graphite TS binding energy

H

- H2O
- H2O molecular dynamics

N cont.

- NiO GGA
- NiO GGA+U
- NiO HSE06
- NiO L(S)DA+U
- NiO LSDA
- NiO LSDA+U
- Nucleophile Substitution CH3Cl - Standard MD
- Nucleophile Substitution CH3Cl - BM
- Nucleophile Substitution CH3Cl - mMD1
- Nucleophile Substitution CH3Cl - mMD2
- Nucleophile Substitution CH3Cl - mMD3
- Nucleophile Substitution CH3Cl - SG

O

- O atom
- O atom spinpolarized
- O atom spinpolarized low symmetry
- O dimer

navigation

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search

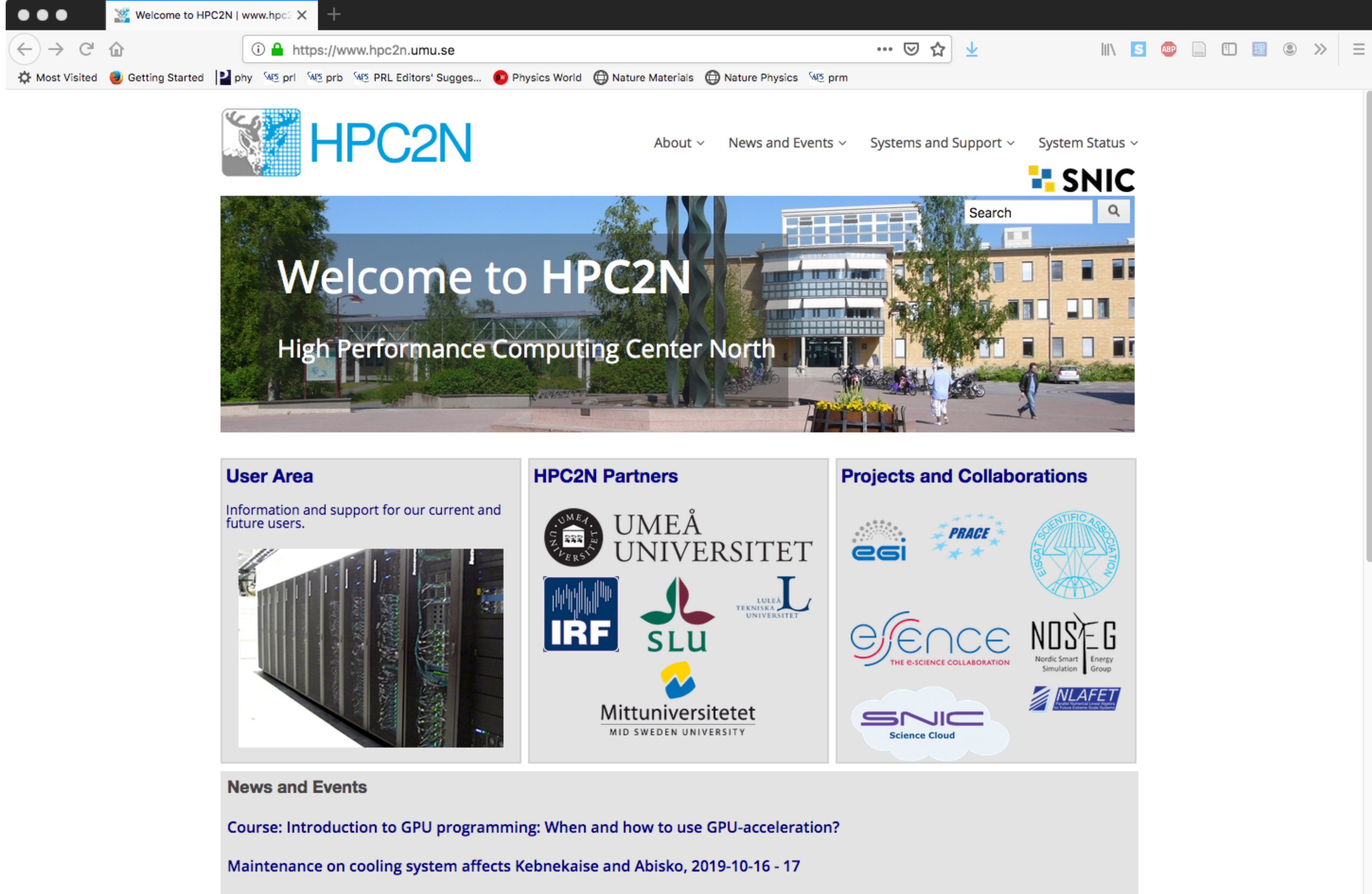
Search

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tools

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<https://www.vasp.at/wiki/index.php/Category:Examples>



<https://www.hpc2n.umu.se/> [Systems and Support > Software > VASP](#)

<https://www.hpc2n.umu.se/resources/software/vasp>



User Area

Information and support for our current and future users.



HPC2N Partners



Projects and Collaborations



News and Events

[Course: Introduction to GPU programming: When and how to use GPU-acceleration?](#)

[Maintenance on cooling system affects Kebnekaise and Abisko, 2019-10-16 - 17](#)

Software www.hpc2n.umu.se		
https://www.hpc2n.umu.se/resources/software		
Most Visited Getting Started phy S prl prb PRL Editors' Sugges... Physics World Nature Materials Nature Physics prm		
	models of macromolecular complexes using Electron Microscopy. External info: ↗	
Siesta	Siesta performs electronic structure calculations and ab initio molecular dynamics simulations of molecules and solids. External info: ↗	Abisko, Kebnekaise
Turbomole	TURBOMOLE is a quantum chemical program package Extern info: ↗	Abisko, Kebnekaise
VASP	Performs ab-initio quantum-mechanical molecular dynamics (MD) using pseudopotentials and a plane wave basis set. Extern info: ↗	Abisko, Kebnekaise
VMD	VMD is a molecular visualization program for displaying, animating, and analyzing large biomolecular systems using 3-D graphics and built-in scripting. Extern info: ↗	Abisko, Kebnekaise
WRF	Advanced Research WRF (ARW) Modeling System is a flexible, state-of-the-art atmospheric simulation system. Extern info: ↗	Abisko, Kebnekaise
Compilers		
GCC	GNU Compiler Collection. Has C, C++, Fortran 77, Fortran 90, and Fortran 95. Available for serial, OpenMP, and MPI code for all languages, with the exception of Fortran 95 MPI programs. Part of several compiler toolchains . External info: ↗	Abisko, Kebnekaise
Intel	Intel Compilers. Has C, C++, Fortran 77, Fortran 90, and Fortran 95. Available for serial, OpenMP, and MPI code for all	Abisko, Kebnekaise



```
freesurfer/5.3.0      mpfr/gcc/3.1.3      turbomole/6.4.0
fsl/5.0.5            mumps/gcc/4.10.0   valgrind/3.12.0
ga/intel-mpi-mkl-i8/5.3  mumps/pgi/4.10.0  vasp/5.3.5
gamess-us/20130501-r1  mumps/psc/4.10.0  vasp-data/5.4
gate/5.0.0_p01       namd/2.9           vasp-tools/0.2
gate/7.1.0           netcdf/4.1.3      verify-module
gaussian/09.0.01     nwchem/6.5        vesta/3.3.2
geant4/10.01.00      octave/3.6.4      visit/2.10.0
geant4-data/10.01    octopus/5.0.0     vmd/1.9.2
gmp/gcc/6.1.0        openblas/0.2.13   voro/gcc/0.4.6
gpaw/0.11.0.13004    openbugs/3.2.3    vtune/2016.2.0
gromacs/5.1.1        openfoam/3.0.1    wrf/3.7.1
gurobi/6.5           openmpi/1.10.1
t-mn01 [~]$
```

[Home](#) » [VASP](#)

VASP

Policy

The Vasp program is not distributed via site licences. However, HPC2N have access to the VASP code to be able to support any research groups that have a valid VASP license.

See the VASP license for information regarding terms for published work.

When you have gotten access to a license, the license holder should either add the license info into SUPR (or contact support@hpc2n.umu.se with the following information: license number and list of users who should have access). You will then be given access to using VASP.

Note: only the owner of the license can add/delete users to/from the access list.

General

VASP is a package for performing ab-initio quantum-mechanical molecular dynamics (MD) using pseudopotentials and a plane wave basis set.

Description

VASP is a complex package for performing ab-initio quantum-mechanical molecular dynamics (MD) simulations using pseudopotentials or the

NSC

START SYSTEMS STORAGE SOFTWARE ABOUT USER AREA

Welcome to National Supercomputer Centre at Linköping University

NSC is a provider of leading edge national supercomputing resources. We provide a wide range of high performance computing and data services to members of academic institutions throughout Sweden and to our partners SMHI, MET Norway, and Saab.

OUR PARTNERS

SMHI
The Swedish Meteorological and Hydrological Institute (SMHI) is an expert agency under the

MET Norway
MET Norway is the meteorological service in Norway, serving the public, the civil services

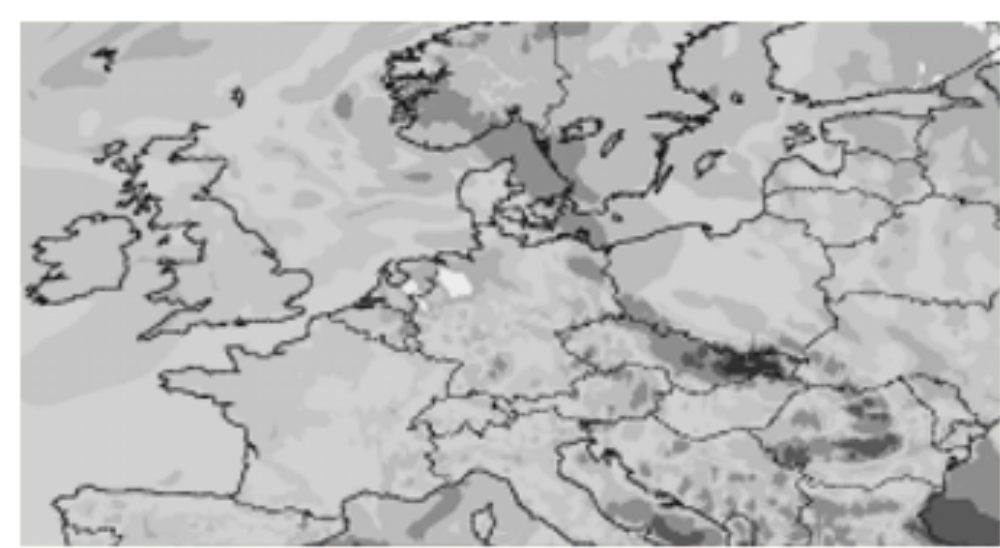
<https://www.nsc.liu.se/> Software > Installed software > Tetralith & Sigma software list > VASP
<https://www.nsc.liu.se/software/installed/tetralith/vasp/>



Welcome to National Supercomputer Centre at Linköping University

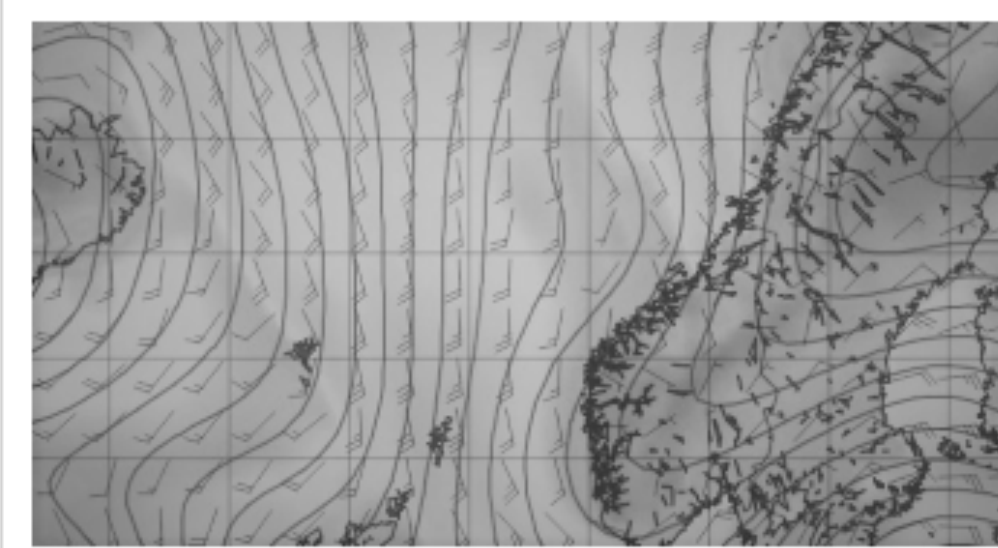
NSC is a provider of leading edge national supercomputing resources. We provide a wide range of high performance computing and data services to members of academic institutions throughout Sweden and to our partners SMHI, MET Norway, and Saab.

OUR PARTNERS



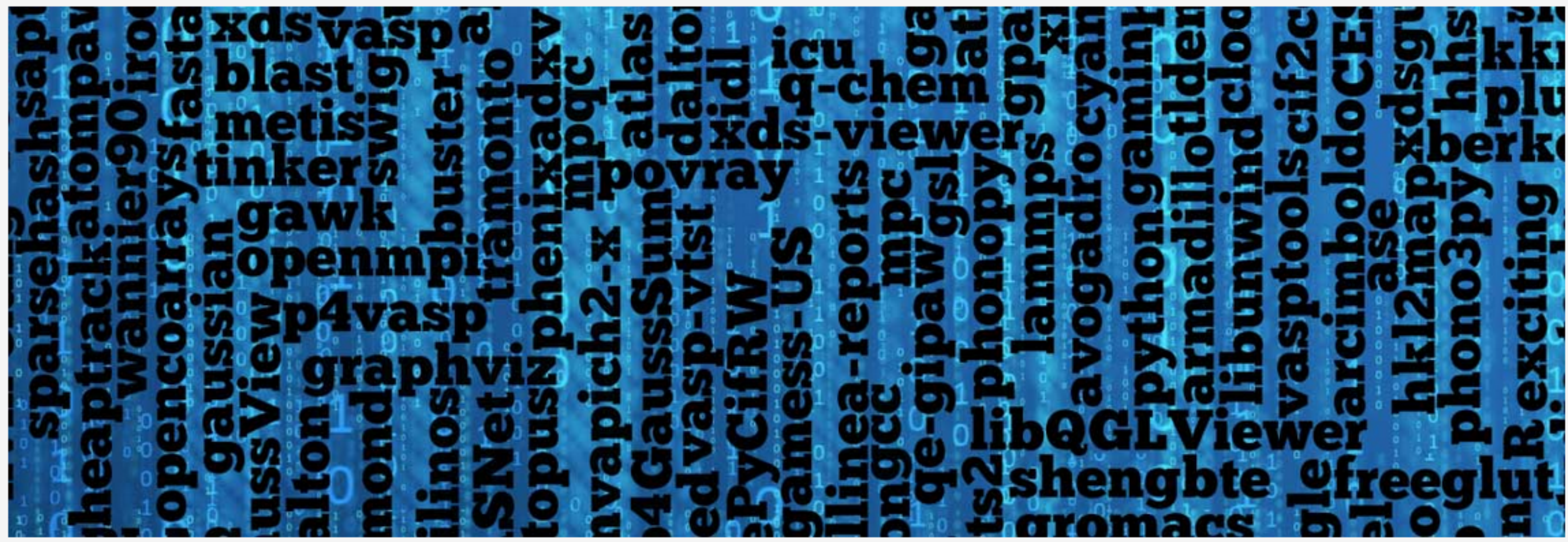
SMHI

The Swedish Meteorological and Hydrological Institute (SMHI) is an expert agency under the



MET Norway

MET Norway is the meteorological service in Norway, serving the public, the civil services



Software

Explore this part to find out about all software environment related matters. Please proceed to the [installed software](#) page to find out which scientific software we have on our clusters. If you are going to compile software from source code, we suggest that you read the [NSC build environment](#) introduction and the [compilers](#) section.

Installed software

Which software is available on what systems and how to run it

Software installation policy

What to do if software you need isn't installed

Software licensing

How we handle software licensing of commercial software.

Compilers

NSC build environment

Modules



Installed software

NSC has a large number of software installations available, often in multiple versions to suit the needs of various user communities. For a list of installed software, please see the corresponding resource page below. If you need software that is presently not installed, please see our [software installation policy](#).

Software portfolios by cluster

- [Tetralith & Sigma Software List](#).
- [Bi Software List](#). This is the main resource for finding out what software is available on Bi.

Module system

You can also query the [module system](#) for available software and recommendations on what versions to use, e.g:

```
module avail
module add vasp/recommendation
```

SNIC knowledge base

Information on software and availability for all of SNIC is also available in the [SNIC knowledge base software section](#). There is specific information for these NSC resources:

Tetralith & Sigma Software
 A list of software installed on Tetralith and Sigma and links to further information

Tetralith & Sigma Software

NSC / Software / Installed software / Tetralith & Sigma Software

Tetralith & Sigma Software List

DISCLAIMER: Please note that the software catalogue is a work in progress! If your application is missing, please request it by sending e-mail to [NSC Support](#)

The following scientific applications have been installed centrally under `/software/sse/`. This list may not always be 100% up to date. The most reliable source is running the command `module avail` while logged into Tetralith or Sigma, possibly augmented by `ls /software/sse/manual/` to show additional manually performed installations without modules. Please note that some of this **software is licensed**, and may not be available for everyone. You need ask NSC for access, which is typically granted upon some proof of having a license.

The list was last updated: 2019-02-21

Electronic structure

- [Abinit](#)
- [ASE](#)
- [Elk](#)
- [EPW](#)
- [exciting](#)
- [GPAW](#)
- [phonopy](#)
- [phono3py](#)
- [p4vasp](#)
- [Quantum Espresso](#)
- [vasptools](#)
- [VASP \(licensed\)](#)
- [WIEN2k \(licensed\)](#)

- ABINIT AMBER ANSYS ASE ATAT Alinea Performance Reports Alinea-DDT Alinea-MAP CDO COMSOL
- CP2K CPMD DL_POLY Dalton/LSDalton EC-Earth EPW Eik FERRET GPAW Grace Gurobi Optimizer
- HDF5 Julia LAMMPS MATLAB MOLDEN Mathematica NAMD NCO NCVIEW Open Babel OpenFOAM
- ParaView Quantum ESPRESSO STAR-CCM+ Siesta VMD WEST WIEN2K Yambo ecCodes exciting
- grib_api netCDF p4vasp parallel phono3py phonopy vasptools **VASP** Clang Gaussian and GaussView

NSC / Software / Installed software / Tetralith & Sigma Software / VASP

VASP Installations on Tetralith & Sigma

First of all, VASP is licensed software, your name needs to be included on a VASP license in order to use NSC's centrally installed VASP binaries. [Read more about how we handle licensing of VASP at NSC.](#)

Some problems which can be encountered running VASP are described at the end of this page.

How to run: quick start

A minimum batch script for running VASP looks like this:

```
#!/bin/bash
#SBATCH -J jobname
#SBATCH -N 4
#SBATCH --ntasks-per-node=32
#SBATCH -t 4:00:00
#SBATCH -A SNIC-xxx-yyy

module add VASP/5.4.4.16052018-nsc1-intel-2018b-eb
mpprun vasp_[std/gam/ncl]
```

This script allocates 4 compute nodes with 32 cores each, for a total of 128 cores (or MPI ranks) and runs VASP in parallel using MPI. Note that you should edit the jobname and the account number before submitting.



KTH / PDC

Software

General information about VASP

Licenses

Available Software - General information about VASP

General information about VASP

System	Available versions
Beskow	5.4.4-wannier90 , 5.4.4 , 5.4.1.patched , 5.3.5-vtst3.1 , 5.3.5-31Mar14
Tegner	5.4.1 , 5.3.5

The Vienna Ab initio Simulation Package (VASP) is a computer program for atomic scale materials modeling, e.g. electronic structure calculations and quantum-mechanical molecular dynamics, from first principles.

For more information see: <http://vasp.at>

Licenses

VASP is not free software and requires a software license. VASP licenses are managed in [SUPR](#). All people who want to use VASP should have SUPR accounts and be a member of a VASP group in SUPR. VASP groups have owners, typically a principal investigator of a project, and that owner can add and remove people using the SUPR interface. If you are Ph.D student, we suggest that you check with your supervisor.



KTH / PDC

Software

Using VASP 5.4.4 on Beskow

General observations

NPAR, NCORE and NSIM

How to choose the number of cores

Vasp Filenames

BEEF functionals

VASP TST Tools

VTST Scripts

VASPsol

Using vdW functionals

Running Vasp

Available Software - General information about VASP - Using VASP 5.4.4 on Beskow

Using VASP 5.4.4 on Beskow

General observations

- VASP is not helped by hyper-threading (64 virtual cores per compute node).
- No GPU/OpenMP-support.
- Running on fewer than 32 cores per node allocates more memory to each MPI task. This can in some cases improve performance and is necessary if your job crashes with an OOM error. See the example submit script below on how to do this correctly.

NPAR, NCORE and NSIM

From initial testing, we recommend:

- NPAR = number of compute nodes
- NCORE = cores / node, typically 16,24 or 32.
- NSIM = 2
- KPAR = number of compute nodes (if applicable)

How to choose the number of cores

Rule of thumb:

- 1 atom per core = Good
- 0.5 atom per core = Could work (but had efficiency and time wasted)

Example of day-to-day tools

- less / gedit / vi / nano reading/editing files
- grace / gnuplot plotting tools
- Bash simple scripts
- cif2cell convert from .cif, create structures
- p4vasp analysis of VASP output
- xcrysden / vesta view structure
- ASE different tools (Python), create structures

Also of interest:

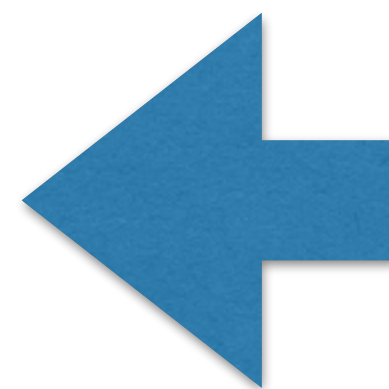
- Python / R analysis etc. (check out e.g. using jupyter)
- Matlab / Octave analysis etc.
- Schrödinger create/view structure @Kebnekaise, Tetralith

[Quick guide to Tetralith for the workshop](#)
(Presentations from old Tetralith training)



Different types of calculations

- Structural relaxation (different ways)
- Regular E_{tot} scf run using PBE, HSE06, GW, ...
- Density of states, bandstructure, charge density, ...
- Born-Oppenheimer MD
- Used within a special framework (VTST, ...)
- See [VASP wiki examples](#)



Input files

- [INCAR](#) - input parameters
- [POSCAR](#) - structure (generate using *e.g.* cif2cell)
- [POTCAR](#) - PAW potentials (how to select?)
- [KPOINTS](#) - k-mesh (or list)
- + job script

SLURM batch queue system &
settings used by SNIC centers

INCAR parameters

- [PREC](#) - “precision”, ENCUT and FFT grids
- [ENCUT](#) - plane wave energy cutoff
- [ALGO](#) - wf optimisation
- [NBANDS](#) - if not set, auto-determined
- [NSIM](#) - for RMM-DIIS algorithm (ALGO)
- [NCORE](#) or [NPAR](#) - bands treated in parallel
- [KPAR](#) - k-point parallel

INCAR parameters

accuracy /
method

- PREC - “precision”, ENCUT and FFT grids
- ENCUT - plane wave energy cutoff **Completeness of basis-set**
Recommended to set!
- ALGO - wf optimisation
- NBANDS - if not set, auto-determined
Must be the same for Etot comparison!

parallel
calcs.

- NSIM - for RMM-DIIS algorithm (ALGO)
- NCORE or NPAR - bands treated in parallel
- KPAR - k-point parallel

- We will get back to the settings in part 3!

INCAR defaults

- PREC = Normal Might want “Accurate”
- ENCUT = ? **Always set!** ENMAX x1.0 - x1.5
- ALGO = Normal ^{good tradeoff} Can use “Fast” and “VeryFast”
- NBANDS = ? can be **overridden** by VASP
- sometimes extra empty states needed
- NSIM = 4 Typically OK
- NCORE = 1 **Adjust** (if not hybrid-functional, HSE06, etc.)
- KPAR = 1 for k-point parallel calcs.

Will discuss in more detail later on...

INCAR defaults

in very brief,
refer to
VASP wiki
for details

- NSW = 0 max ionic steps, also MD steps
- NELM = 60 max electronic selfconsistency steps
- NELMIN = 2 min steps. For relaxation/MD set 4-8
- EDIFF = 1E-4 converge to 4 last digits, sometimes higher accuracy is needed
- EDIFFG = EDIFF x10 ionic relaxation break condition,
if negative value, break if forces < |EDIFFG|
- ISMEAR = 1 how to treat partial electron occupancy:
1 = metals, 0 = bandgap, -5 = for accurate E_{tot}
- ISPIN = 1 2 = spin-polarized calc.
- IBRION = -1 (NSW=-1,0) or 0 how ions are updated & moved
no update MD =2 ionic relaxation

POSCAR

A simple case of fcc Ni, refer to the [VASP wiki example](#)

(hopefully) useful → comment →

lattice vectors →

number of atoms
per type →

position for first atom →

```
Ni fcc
3.53
0.5 0.5 0.0
0.0 0.5 0.5
0.5 0.0 0.5
Ni
1
Cartesian
0 0 0
```

← lattice constant (Å)

← element symbols

← Cartesian or Direct coordinates

negative value:
cell volume

optional, useful for
clarity & plotting

First letter is sufficient, i.e.
“C” for “Cartesian”

Direct coordinates: expressed in terms of the lattice vectors (no lattice constant, scaling)

Cartesian coordinate: expressed as (x,y,z) with the scaling factor included

POSCAR

From the course examples, H and Si on Ag(111) surface:

100% H on Si on Ag(111)

10.007900

1.0000000000000000 0.0000000000000000 0.0000000000000000

0.5000000000000000 0.866025403784439 0.0000000000000000

0.0000000000000000 0.0000000000000000 4.352531500114909

H Si Ag

14 14 108

← Note order of atoms

Selective dynamics

← Relax for different directions

Direct

first H atom →

0.7583380000000000 0.0528816000000000 0.6059450000000000

0.5052440000000000 0.1182310000000000 0.6059450000000000

0.0507845000000000 0.1960170000000000 0.6059450000000000

0.8003590000000000 0.2519030000000000 0.6059450000000000

0.3333333333333333 0.3389020000000000 0.6059450000000000

0.1141910000000000 0.3862440000000000 0.6059450000000000

0.6184740000000000 0.4980580000000000 0.6059450000000000

0.3812620000000000 0.5113130000000000 0.6059450000000000

0.8830710000000000 0.6246840000000000 0.6059450000000000

Selective dynamics
always for Direct coord.
T = relax
F = fixed

Rest of H,
Si & Ag atoms
following

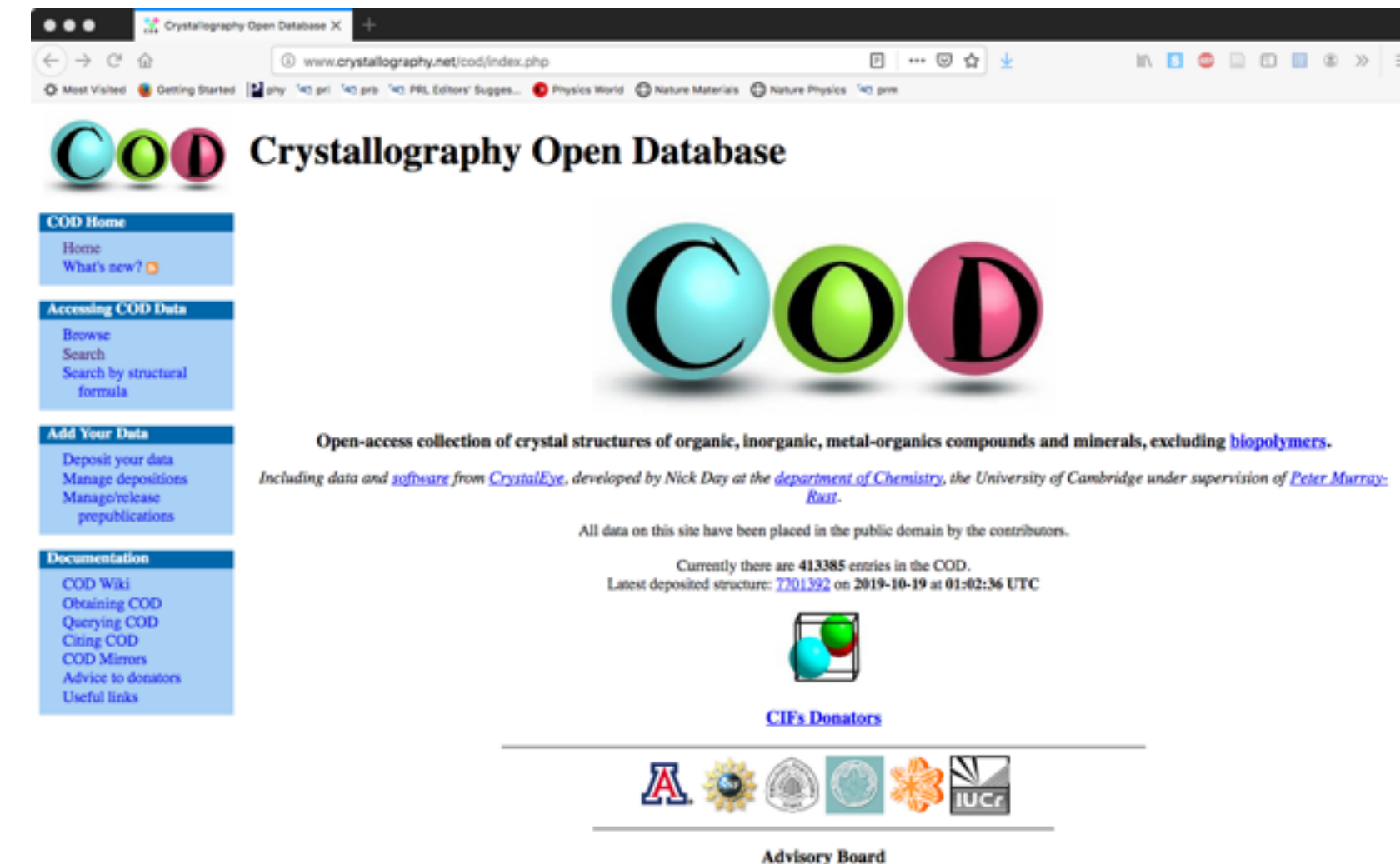


POSCAR

Some useful resources:

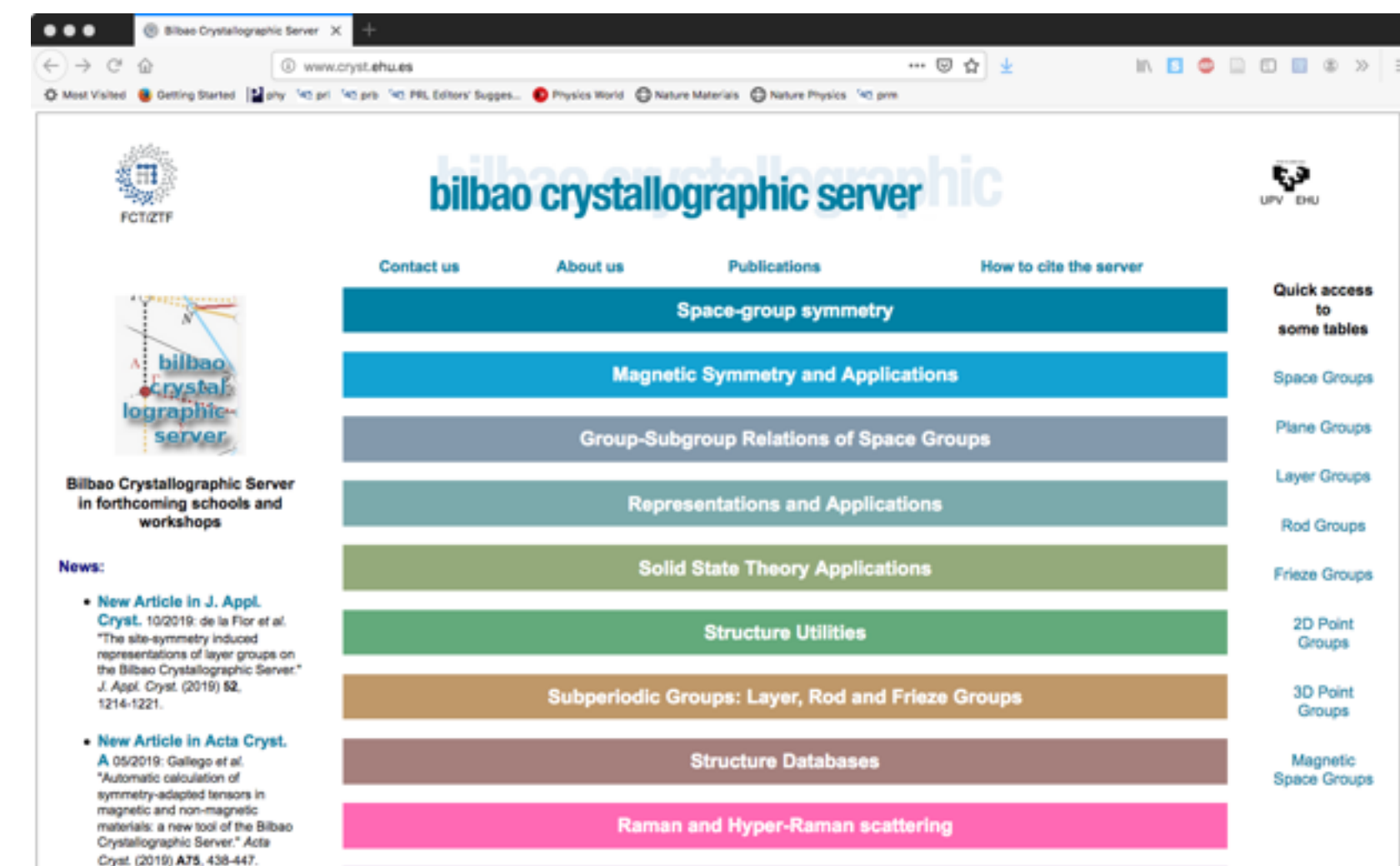
[Crystallography Open Database](http://www.crystallography.net/cod/index.php)

Database with published structures from experiment .cif



[Bilbao Crystallographic Server](http://www.crysl.ehu.es)

Many Crystallographic tools, e.g. check BZ of fcc cell



POSCAR

A few examples on how to visualize and/or edit POSCAR:

[Atomic Simulation Environment \(ASE\)](#)

Handle structures (and much more) using python scripts, also GUI

[cif2cell](#)

Versatile script, reads .cif
saves to many formats including
POSCAR - also build supercells

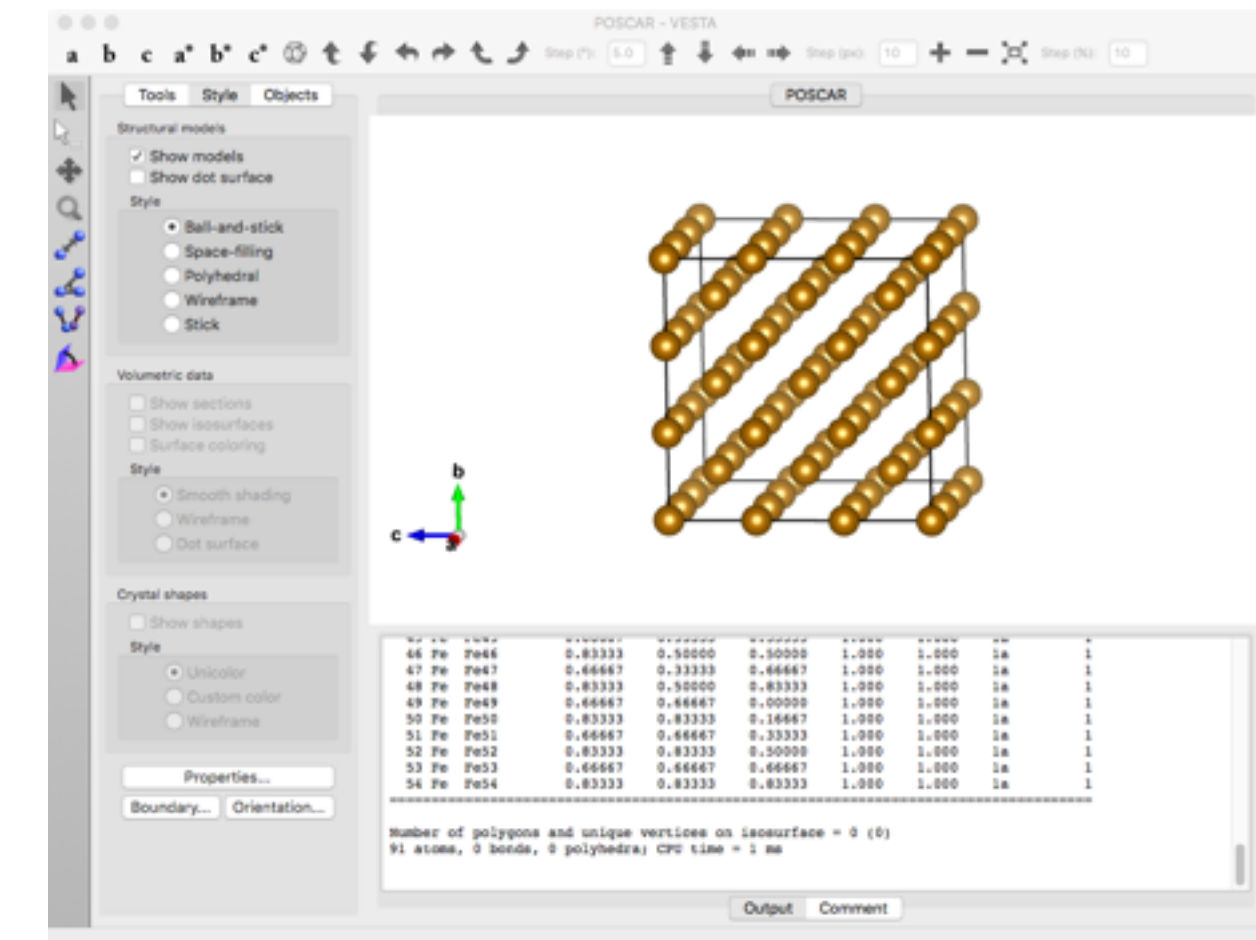
Commercial software:

[NanoLab](#)

[MedeA](#)

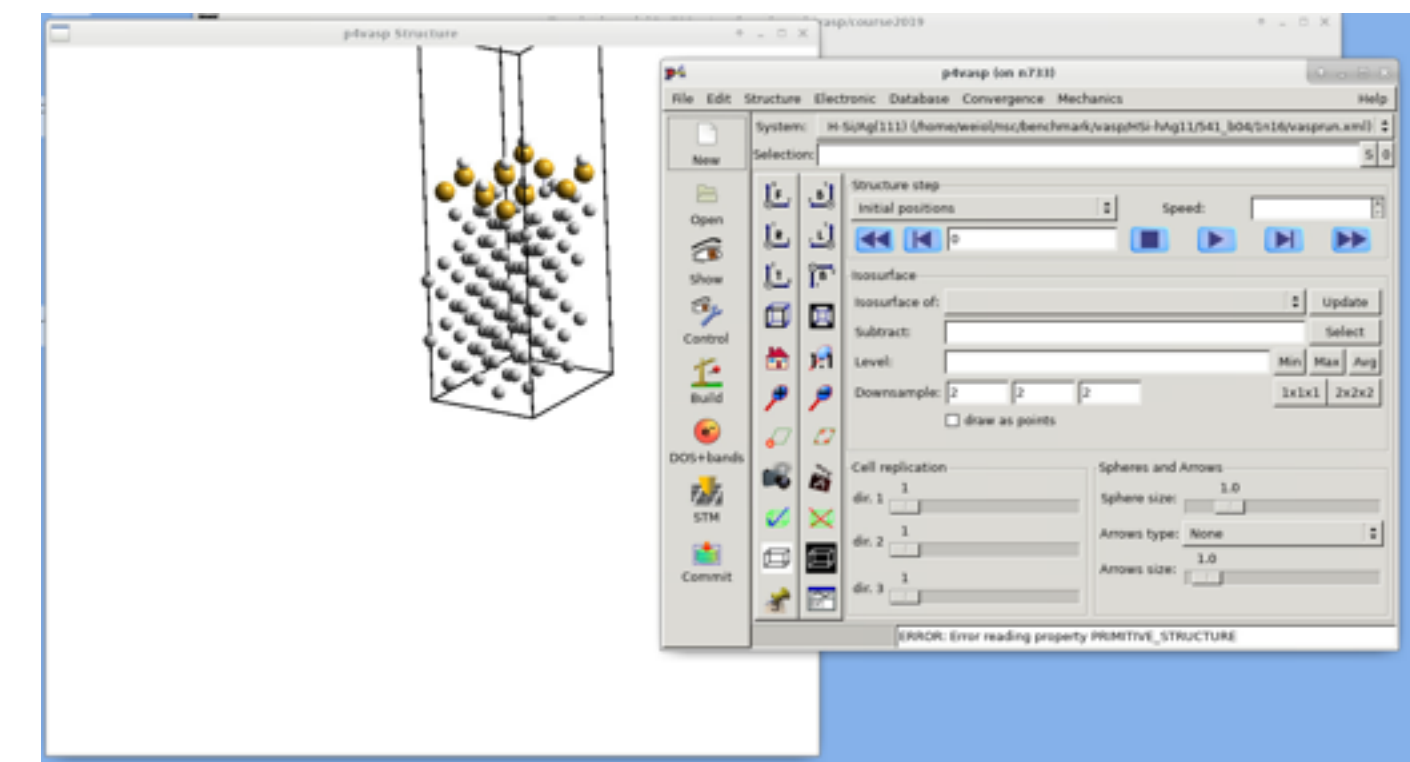
[MaterialsStudio](#)

[VESTA](#)



Opens .cif displays structure, save as POSCAR

[p4vasp](#)



Apart from analysis and visualization, also edit POSCAR

POTCAR

PAW potentials - non-trivial to tailor, select with care

```
PAW_PBE Cu 22Jun2005 ← type, element, date
valence → 11.000000000000000
parameters from PSCTR are:
XR-type → VRHFIN =Cu: d10 p1 ← atomic configuration
LEXCH = PE
EATOM = 1390.9808 eV, 102.2342 Ry

TITEL = PAW_PBE Cu 22Jun2005
LULTRA = F use ultrasoft PP ?
IUNSCR = 1 unscreen: 0-lin 1-nonlin 2-no
RPACOR = 2.000 partial core radius
POMASS = 63.546; ZVAL = 11.000 mass and valenz
RCORE = 2.300 outmost cutoff radius
RWIGS = 2.200; RWIGS = 1.164 wigner-seitz radius (au A)
energy cutoff → ENMAX = 295.446; ENMIN = 221.585 eV ← smallest energy cutoff
ICORE = 3 local potential
LCOR = T correct aug charges
LPAW = T paw PP
EAUG = 586.980
DEXC = 0.000
RMAX = 2.344 core radius for proj-oper
RAUG = 1.300 factor for augmentation sphere
RDEP = 2.302 radius for radial grids
RDEPT = 1.771 core radius for aug-charge
```

POTCAR

- Check [recommendations](#), LDA, PBE

- for short bonds: `_h`
strong pressure

- for GW: `_GW`

- States in valence: `_sv`, `_pv`, `_d`

- “soft” (no short bonds): `_s`

- Where?

@Kebnekaise: `$ echo $VASP_PP_PATH`
`/hpc2n/eb/software/Core/VASP-data/5.4/potPP`

@Tetralith: `/software/sse/manual/vasp/POTCARs`

Note several choices, e.g.:

Ga, `Ga_d`, `Ga_d_GW`,
`Ga_GW`, `Ga_h`, `Ga_sv_GW`

Useful commands:

```
$ grep PAW POTCAR
```

```
$ grep ENMAX POTCAR
```

KPOINTS

A simple case of fcc Ni, 1 atom

0 = automatic generation of mesh →

```
k-points ← comment
0
Monkhorst Pack ← Monkhorst-Pack method (M)
11 11 11 ← odd kmesh - includes  $\Gamma$ -point
0 0 0 ← optional shift of k-mesh
```

My example, H and Si on Ag(111) surface, 136 atoms

First letter is sufficient, i.e.
“G” for “Gamma”

```
Automatic mesh
0
Gamma ← Gamma method (G)
2 2 1
0. 0. 0.
```

- Γ -point included by default
- hexagonal structures only use this!

KPOINTS

For **bandstructure** calculations, provide a list of k-points, [see example](#)

k-points for bandstructure L-G-X-U K-G ← comment

k-points per line-segment → 10

line ← k-points per line-segment

Reciprocal / Cartesian → Reciprocal

0.50000	0.50000	0.50000	1	← symmetry point + weight
0.00000	0.00000	0.00000	1	
0.00000	0.00000	0.00000	1	
0.00000	0.50000	0.50000	1	
0.00000	0.50000	0.50000	1	
0.25000	0.62500	0.62500	1	
0.37500	0.7500	0.37500	1	
0.00000	0.00000	0.00000	1	

KPOINTS

- Metal - “many” k-pts
 - Band gap materials - “few” k-pts
 - Unit cell (few atoms) - more k-pts
 - Supercell (100s atoms) - few/one, k-pt
 - No guarantee for convergence...
 - MP method popular, G “safest” to apply
 - 1x1x3 cell geometry → 3x3x1 k-mesh
- real vs. reciprocal space
- real vs. reciprocal space

VASP binaries

- `vasp_std` - regular version
- `vasp_gam` - one k-point (Gamma), **faster**
- `vasp_ncl` - noncollinear magnetism
- `vasp_gpu`, `vasp_gpu_ncl` - for GPU
- + modifications

e.g. constrained relaxation

Job script - Kebnekaise (HPC2N)

```
#!/bin/bash
#SBATCH -A snic2019-3-203
#SBATCH -J test
#SBATCH -t 3:59:00
#SBATCH -n 56

ml icc/2017.4.196-GCC-6.4.0-2.28
ml ifort/2017.4.196-GCC-6.4.0-2.28
ml impi/2017.3.196
ml VASP/5.4.4-18Apr17-hpc2n

mpirun vasp_std
```

#SBATCH -N 2

Example: running on 2 nodes (28x2 cores) @Kebnekaise

<https://www.hpc2n.umu.se/resources/software/vasp> running @Abisko: note differences

Job script - Kebnekaise (HPC2N)

```
#!/bin/bash
#SBATCH -A snic2019-3-203
#SBATCH -J test
#SBATCH -t 3:59:00
#SBATCH -N 2
#SBATCH --ntasks-per-node=14

ml icc/2017.4.196-GCC-6.4.0-2.28
ml ifort/2017.4.196-GCC-6.4.0-2.28
ml impi/2017.3.196
ml VASP/5.4.4-18Apr17-hpc2n

mpirun vasp_std
```

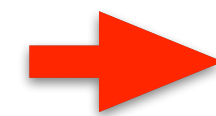
#SBATCH -n 56

Example: running on 2 nodes (28x2 cores), using half the cores
for more memory @Kebnekaise

<https://www.hpc2n.umu.se/resources/software/vasp> running @Abisko: note differences

Job script - Kebnekaise (HPC2N)

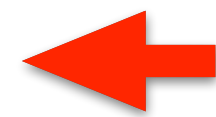
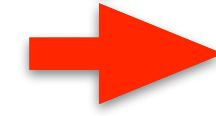
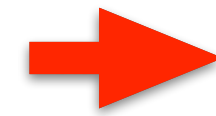
GPU calc.



```
#!/bin/bash
#SBATCH -A snic2019-3-203
#SBATCH -J test
#SBATCH -t 3:59:00
#SBATCH -n 28
#SBATCH --gres=gpu:v100:2,gpuexcl

ml icc/2017.4.196-GCC-6.4.0-2.28
ml ifort/2017.4.196-GCC-6.4.0-2.28
ml impi/2017.3.196
ml CUDA/9.1.85
ml VASP/5.4.4-18Apr17-p01-hpc2n

mpirun vasp_gpu
```



Example: running on 1 node (28 cores) with 2xV100 GPUs @Kebnekaise

<https://www.hpc2n.umu.se/resources/software/vasp>

Job script - Tetralith (NSC)

#SBATCH -n 64

note "mpprun"

```
#!/bin/bash
#SBATCH -A snic2019-3-203
#SBATCH -J test
#SBATCH -t 3:59:00
#SBATCH -N 2

module load VASP/5.4.4.16052018-nsc1-intel-2018a-eb
mpprun vasp
```

Example: running on 2 nodes (32x2 cores) @Tetralith

To increase available memory, reduce cores/node, e.g:
#SBATCH --ntasks-per-node=16

Alternatively, use "fat" memory nodes:
#SBATCH -C fat

<https://www.nsc.liu.se/software/installed/tetralith/vasp/>

Job script - Beskow (PDC)

```
#!/bin/bash -l
#SBATCH -A 2019-3-203
#SBATCH -J test
#SBATCH -t 3:59:00
#SBATCH --nodes=2

module unload cray-mpich/7.0.4
module load vasp/5.4.4

aprun -n 48 -N 24 vasp
#aprun -n 64 -N 32 vasp
```

Example: running on 2 nodes (24x2 cores) @Beskow
alternatively on 2 nodes (32x2 cores)

<https://www.pdc.kth.se/software/software/VASP/beskow/5.4.4/index.html#running-vasp>

Output files

- OUTCAR - main, detailed output
- OSZICAR - iteration summary
- **slurm-***.out** - **stdout**, iteration summary, **warnings**
- CONTCAR - updated structural data (at finish)
structural relaxation / MD
- XDATCAR - positions at each ionic step
- ...

Output files

- [DOSCAR](#) - total, partial density of states (DOS)
- [CHGCAR](#) - charge density
output can also be switched off
- [WAVECAR](#) - plane wave coefficients (for restart)
- ...

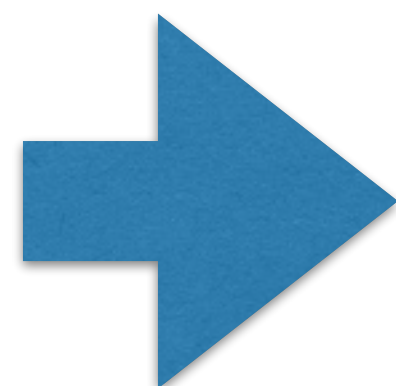
OSZICAR

Min. algo	Step	Total free Energy	Energy diff.	Eigenvalue diff.	Charge density residual vector		
	N	E	dE	d eps	ncg	rms	rms (c)
DAV:	1	-0.189343666468E+01	-0.18934E+01	-0.20040E+03	904	0.422E+02	
DAV:	2	-0.108926039335E+02	-0.89992E+01	-0.87586E+01	1440	0.554E+01	
DAV:	3	-0.109805531666E+02	-0.87949E-01	-0.87949E-01	1208	0.675E+00	
DAV:	4	-0.109807517982E+02	-0.19863E-03	-0.19863E-03	1368	0.313E-01	
DAV:	5	-0.109807519113E+02	-0.11307E-06	-0.11310E-06	1256	0.684E-03	0.519E+00
DAV:	6	-0.108723496529E+02	0.10840E+00	-0.69164E-02	1064	0.137E+00	0.317E+00
DAV:	7	-0.108218097854E+02	0.50540E-01	-0.13575E-01	1120	0.205E+00	0.163E-01
DAV:	8	-0.108228444695E+02	-0.10347E-02	-0.32972E-03	944	0.419E-01	0.706E-02
DAV:	9	-0.108230614389E+02	-0.21697E-03	-0.22028E-04	1312	0.111E-01	0.557E-02
DAV:	10	-0.108230846187E+02	-0.23180E-04	-0.25743E-05	560	0.381E-02	

1 F= -.10823085E+02 E0= -.10823085E+02 d E =-.431458E-08

Final total free energy

Total steps: NELMIN to NELM



Need to check if convergence is reached!

In particular if NELM was reached (default = 60 steps)

Stdout (slurm-***.out)

running on 2 total cores
distrk: each k-point on 2 cores, 1 groups
distr: one band on 1 cores, 2 groups
using from now: INCAR
vasp.5.4.4.18Apr17-6-g9f103f2a35 (build Sep 13 2019 06:30:52) complex

POSCAR found type information on POSCAR Si
POSCAR found : 1 types and 2 ions
scaLAPACK will be used
LDA part: xc-table for Pade appr. of Perdew
POSCAR, INCAR and KPOINTS ok, starting setup
FFT: planning ...
WAVECAR not read
entering main loop

- Check for warnings!

	N	E	dE	d eps	ncg	rms	rms (c)
DAV:	1	-0.189343666468E+01	-0.18934E+01	-0.20040E+03	904	0.422E+02	
DAV:	2	-0.108926039335E+02	-0.89992E+01	-0.87586E+01	1440	0.554E+01	
DAV:	3	-0.109805531666E+02	-0.87949E-01	-0.87949E-01	1208	0.675E+00	
DAV:	4	-0.109807517982E+02	-0.19863E-03	-0.19863E-03	1368	0.313E-01	
DAV:	5	-0.109807519113E+02	-0.11307E-06	-0.11310E-06	1256	0.684E-03	0.519E+00
DAV:	6	-0.108723496529E+02	0.10840E+00	-0.69164E-02	1064	0.137E+00	0.317E+00
DAV:	7	-0.108218097854E+02	0.50540E-01	-0.13575E-01	1120	0.205E+00	0.163E-01
DAV:	8	-0.108228444695E+02	-0.10347E-02	-0.32972E-03	944	0.419E-01	0.706E-02
DAV:	9	-0.108230614389E+02	-0.21697E-03	-0.22028E-04	1312	0.111E-01	0.557E-02
DAV:	10	-0.108230846187E+02	-0.23180E-04	-0.25743E-05	560	0.381E-02	

1 F= -.10823085E+02 E0= -.10823085E+02 d E =-.431458E-08

writing wavefunctions

Warning/advice output

Check stdout (slurm-***.out)
for warnings!

Typical warnings:

Reminder to set (if applicable):
[NCORE](#)
typically = used cores/nodes

For high accuracy (default) keep:
[LREAL=.FALSE.](#)

```
W   W   AA   RRRRR   N   N   II   N   N   GGGG   !!!  
W   W   A   A   R   R   NN   N   II   NN   N   G   G   !!!  
W   W   A   A   R   R   N   N   N   II   N   N   N   G   !!!  
W   WW  W   AAAAAA  RRRRR   N   N   N   II   N   N   N   G   GGG   !  
WW  WW  A   A   R   R   N   NN   II   N   NN   G   G  
W   W   A   A   R   R   N   N   II   N   N   GGGG   !!!
```

For optimal performance we recommend to set
NCORE= 4 - approx SQRT(number of cores)
NCORE specifies how many cores store one orbital (NPAR=cpu/NCORE).
This setting can greatly improve the performance of VASP for DFT.
The default, NCORE=1 might be grossly inefficient
on modern multi-core architectures or massively parallel machines.
Do your own testing !!!!
Unfortunately you need to use the default for GW and RPA calculations.
(for HF NCORE is supported but not extensively tested yet)

ADVICE TO THIS USER RUNNING 'VASP/VAMP' (HEAR YOUR MASTER'S VOICE ...):

You have a (more or less) 'large supercell' and for larger cells
it might be more efficient to use real space projection operators
So try LREAL= Auto in the INCAR file.
Mind: If you want to do a very accurate calculations keep the
reciprocal projection scheme (i.e. LREAL=.FALSE.)

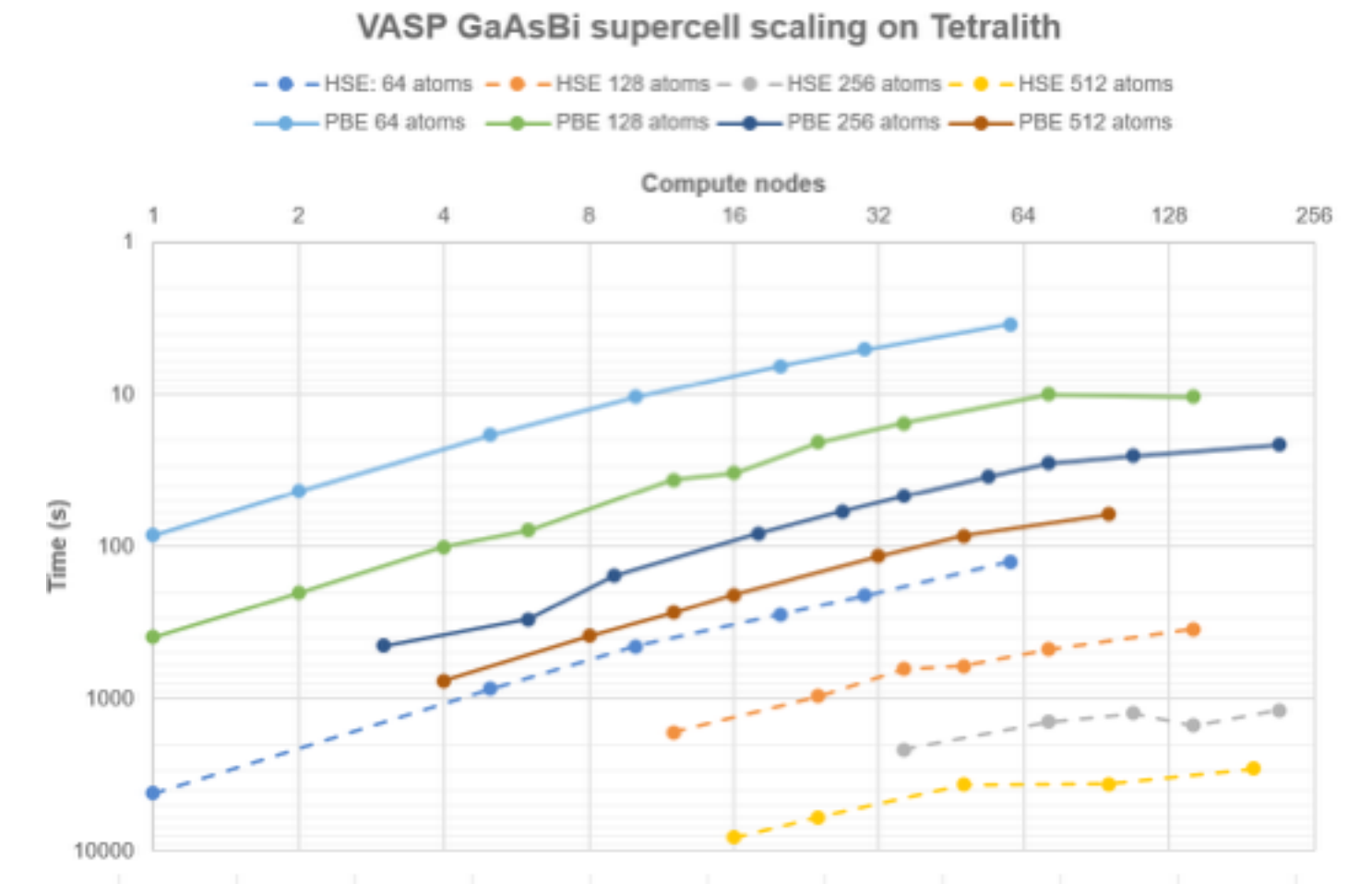
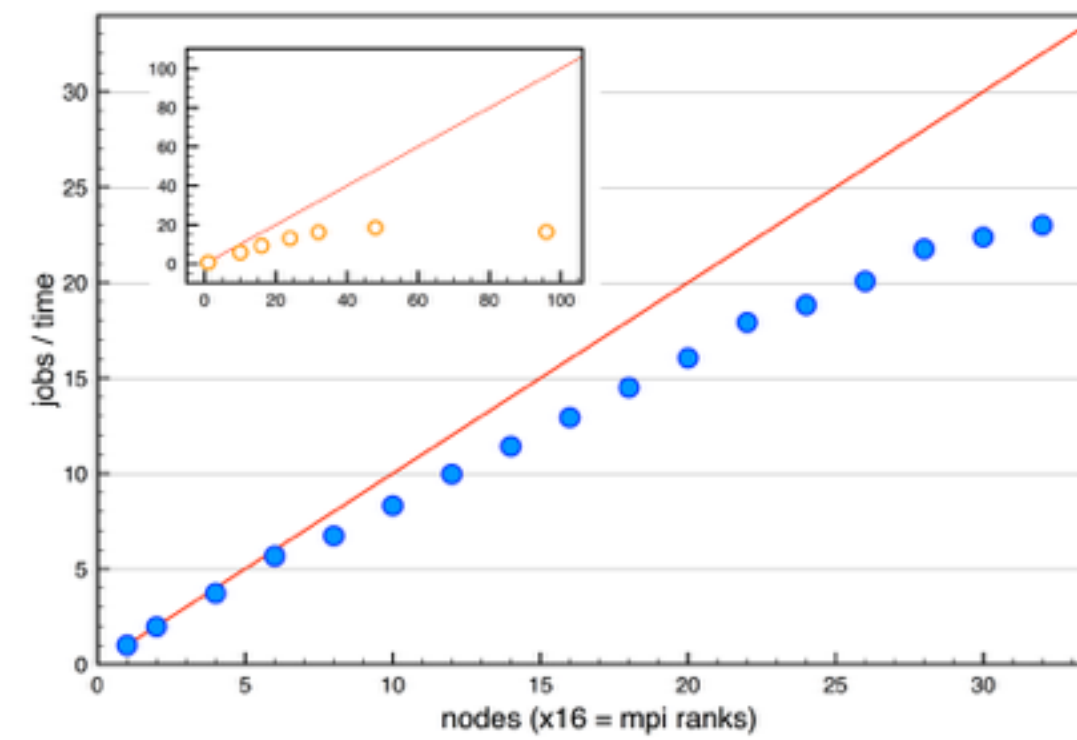
VASP6

- Link to features in VASP6.X
- Hybrid **OpenMP** & MPI parallelization
- **OpenACC** for GPU
- Cubic scaling RPA and GW
- Electron-phonon coupling using stochastic displacements of atoms
- Test suite - Note! potential issue!

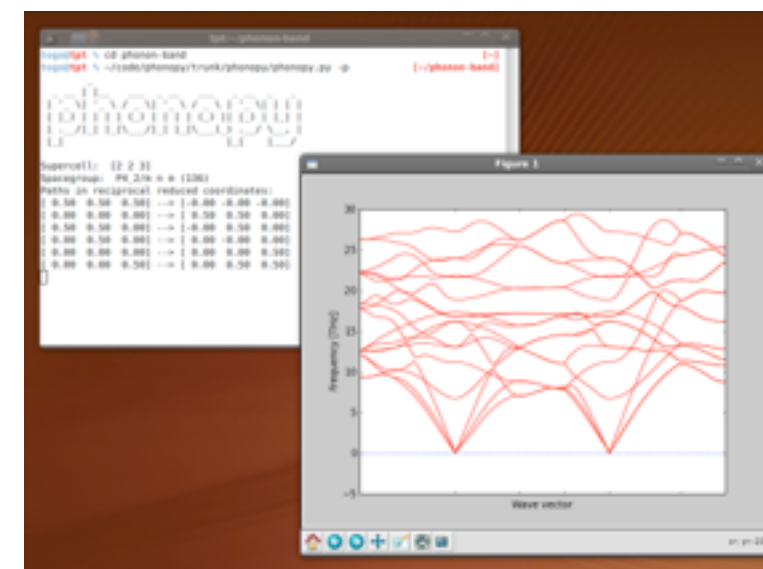
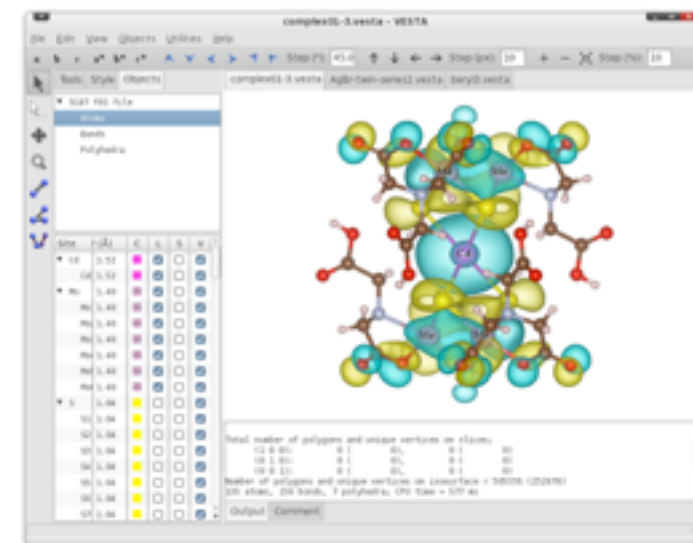
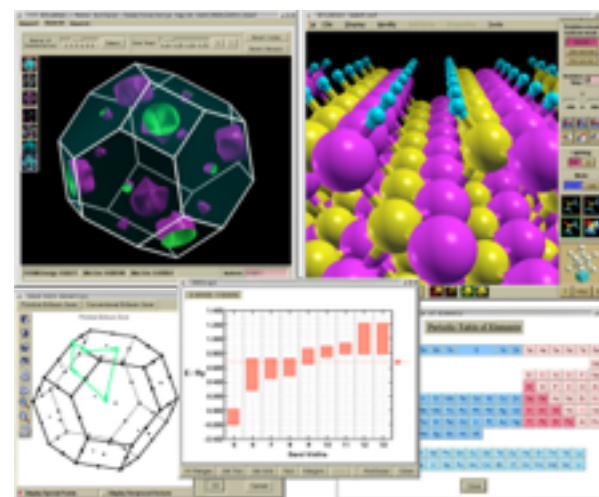
Tomorrow:

- Running & Performance

- Parallelization
- Efficient settings
- **Problems**



- Utilities & Summary



VTST•Tools

USPEX Computational Materials Discovery