3. Running & Performance

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National Supercomputer Centre (NSC), Linköping University NAISS-ENCCS training, online 4-5th Apr 2023



VASP - Best Practices Workshop







Introduction

- General considerations
- Focus on practical aspects of running VASP ...at specific supercomputer centres
- Influential parameters, NPAR/NCORE, ALGO, NSIM, KPAR, ...
- Memory usage
- Benchmarks, examples
- Common problems

... clickable links are underlined

Parallel calculations

Examples for different NAISS HPC clusters



Computation - considerations

Efficiency:

Running as many jobs as possible for a given allocation of computer time

Speed:

The amount of time (real, "human time") to run a specific simulation from when it starts

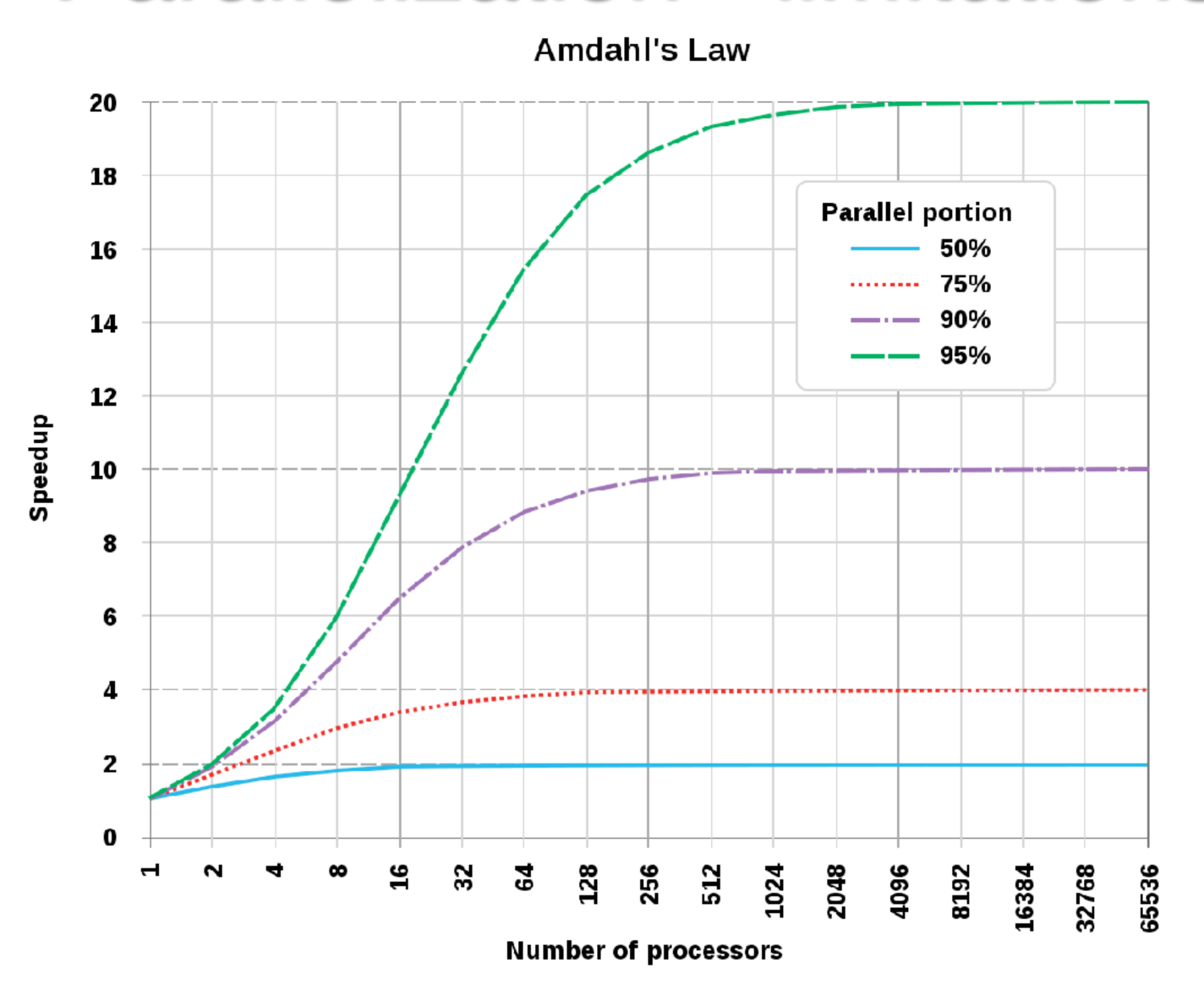
Time-to-solution:

Speed + the time waiting in queue

@Tetralith: wall-time limit 7 days

@Dardel: 24h (7 day option)

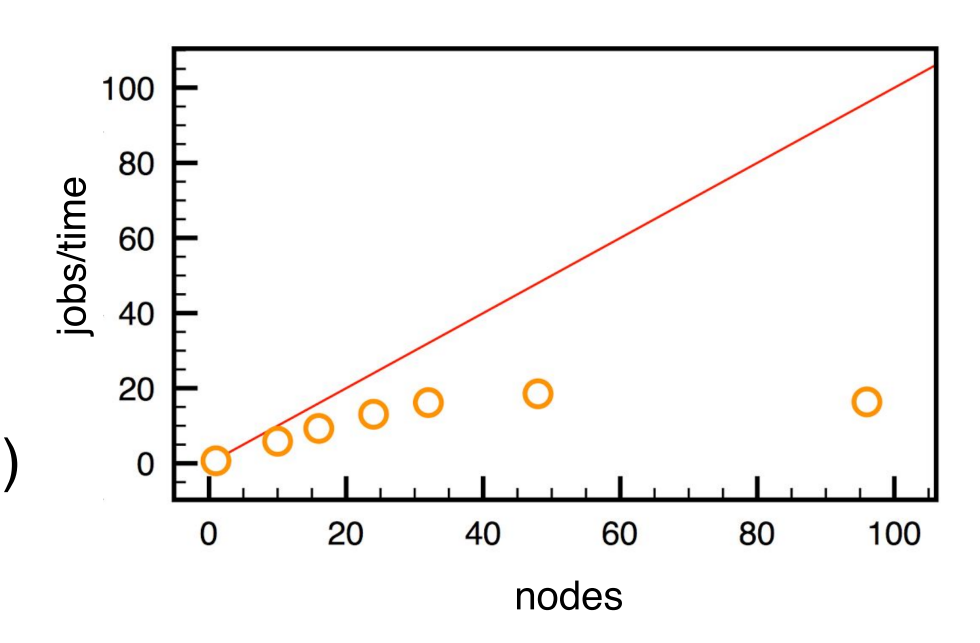
Parallelization - limitations

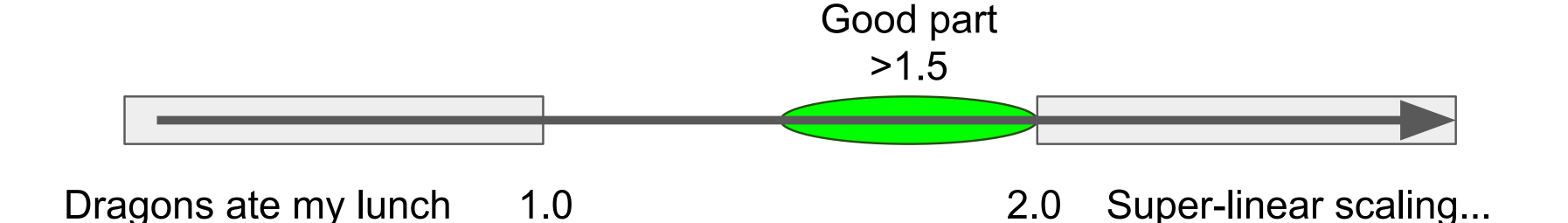


Simple scaling analysis

A minimal scaling analysis can save lots of allocated core hours...

- 1. Tool your runscript to time your simulation
- 2. Run an initial best guess number of cores (n)
- 3. Run the same test on half the number of cores (n/2)
- 4. Score = time(n/2) / time(n)





Hardware - affects best practices

• Tetralith (NSC), Intel Xeon Gold 6130 2.1GHz

I node = 32 cores (96GB RAM, fat node 384GB)

1832 x

60 x

• Dardel (PDC), HPE Cray EX, AMD EPYC 2.25 GHz

I node = 128 cores (256GB RAM, 512GB, 1024GB, 2048)

488 x 20 x 8 x 2 x

Hardware - affects best practices

Local systems, not supported by NAISS in 2023

Kebnekaise (HPC2N)

```
432 x | node (Intel Xeon E5-2690v4) = 28 cores (128GB RAM)

32 x + 2xGPU (NVidia K80) vasp-gpu
version!

52 x | node (Intel Xeon Gold 6132) = 28 cores (192GB RAM)

10 x + 2xGPU (Nvidia V100) vasp-gpu
version!

36 x | KNL (Intel Xeon Phi 7250) node = 68 cores (192GB RAM)
```

Abisko (HPC2N)

different

best

practices

```
318 x I node (AMD Opteron 6238) = 48 cores (128GB RAM)
```

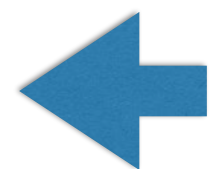
Running & performance

Important INCAR settings and benchmarks



Starting advice (reminder)

- Read the <u>documentation!</u>
- VASP default settings



good starting point

- Caution: "inherited" starting files
- Avoid overly complex INCAR
- Possible differences in centres installations

refer to respective webpages / documentation

Quick check your run

- How much/what resources to use?
 - Check NBANDS \$ grep NBANDS OUTCAR
 - Use ca. 8 bands/core
- How long will it take?
 \$ grep LOOP OUTCAR
 \$ grep LOOP+ OUTCAR
 - scales with k-points (IBZKPT) \$ grep k-points OUTCAR
- Does it converge? \$ cat OSZICAR
- Problems? \$ less slurm*.out

Quick check your run: tools

- sacct
- login to node & run top
- @Tetralith: jobload, jobstats & jobsh

```
$ man <command>
"-s r" for running job
```

```
$ sacct --user=<username> -X --format=Elapsed,State,AllocCPUS%9,CPUTimeRaw%20 --starttime=2019-10-01
$ sacct -e example: --format=JobID,Submit,Start,End,Elapsed,Nodelist,State,AllocCPUS%9,CPUTimeRaw%20
```

```
$ seff <jobid>
summary of run
```

```
$ squeue -u <username>
$ scancel <jobid>
```

```
@ Tetralith:
  $ jobload <jobid>
  $ jobsh <node>
```

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 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

PREC

- PREC = "precision", sets ENCUT and FFT grids
- PREC = Normal, default
- PREC = Accurate, highly accurate forces
- OBS: Recommended to set ENCUT by hand

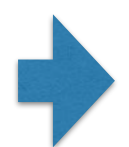
More on accuracy

- NGX, NGY, NGZ coarse plane wave FFT grid can edit directly (otherwise PREC)
- NGXF, NGYF, NGZF finer FFT grid
 - also see <u>ENAUG</u>
- LREAL=.FALSE.

default, might be needed for high accuracy

if proj. operators determined in real space, or not

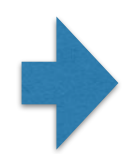
otherwise use faster: LREAL = Auto



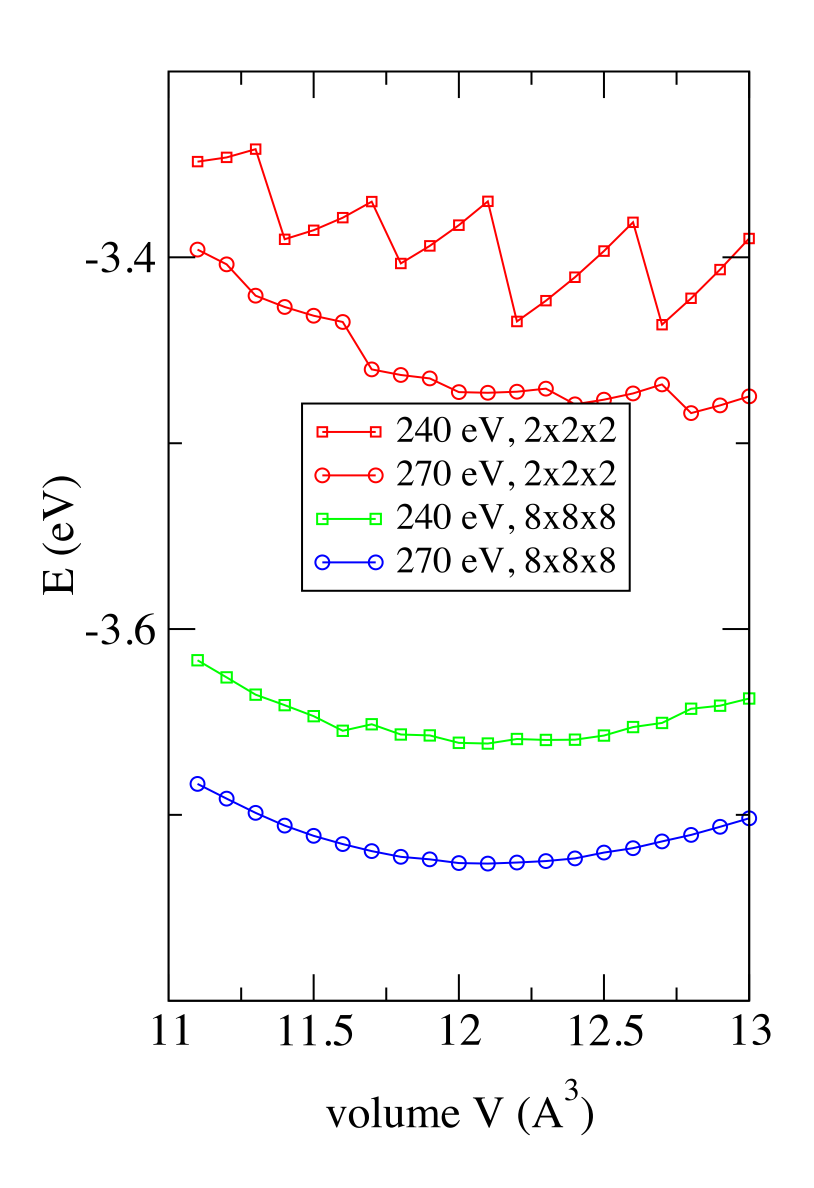
In some cases, need VASP installation with no special optimization flags

Convergence, ENCUT and k-mesh

• Cu example by G. Kresse



- Basis-set changes with volume
- Cell-shape relaxations, increase
 ENCUT = ENMAX x1.3
- Read section on <u>structure relaxation</u>



NBANDS

- NBANDS = NELECT/2 + NION/2 (ISPIN=1)
- May change due to parallelization!
- Easy to divide, 2ⁿ, 4, 8, 12, 16, ...
- select NBANDS = 511 or 512?
- Min limit, 1 band/core
- Affects Etot!

Run e.g. quick job to check NBANDS:

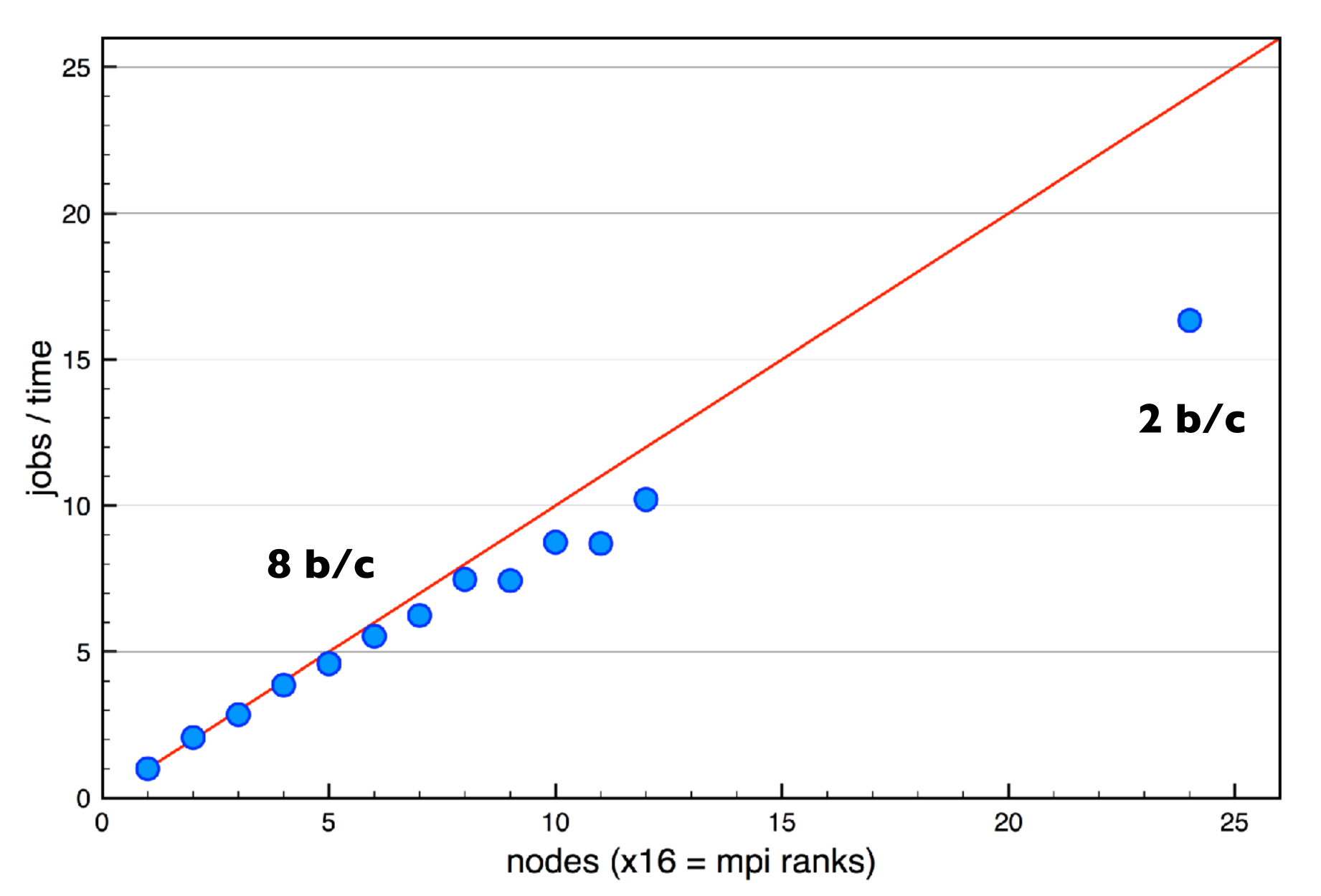
```
#SBATCH --reservation=devel @Tetralith
```

\$ grep NBANDS OUTCAR

How many cores - efficient and/or fast?

- Start from # of bands, NBANDS
- 1 band/core: typically inefficient
- 2 bands/core: ~50% parallel efficiency
- 8 bands/core: good starting point
 - try e.g. cores ≈ NBANDS / 8

Si-H/Ag(111) 129 atoms, VASP PBE @Triolith (old)

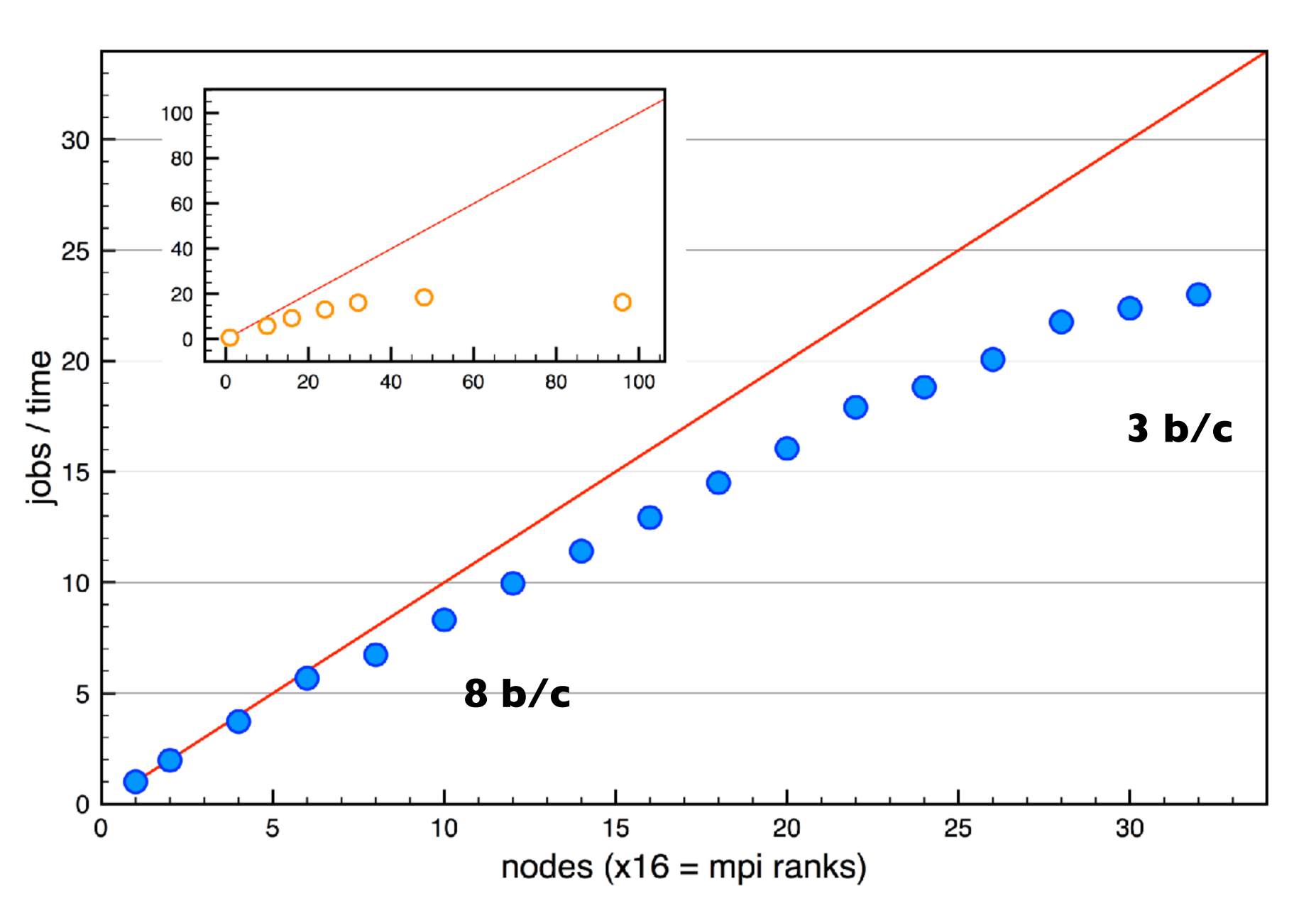


NBANDS=750 4 k-points

Triolith had 16 c/node Tetralith: 32 c/node

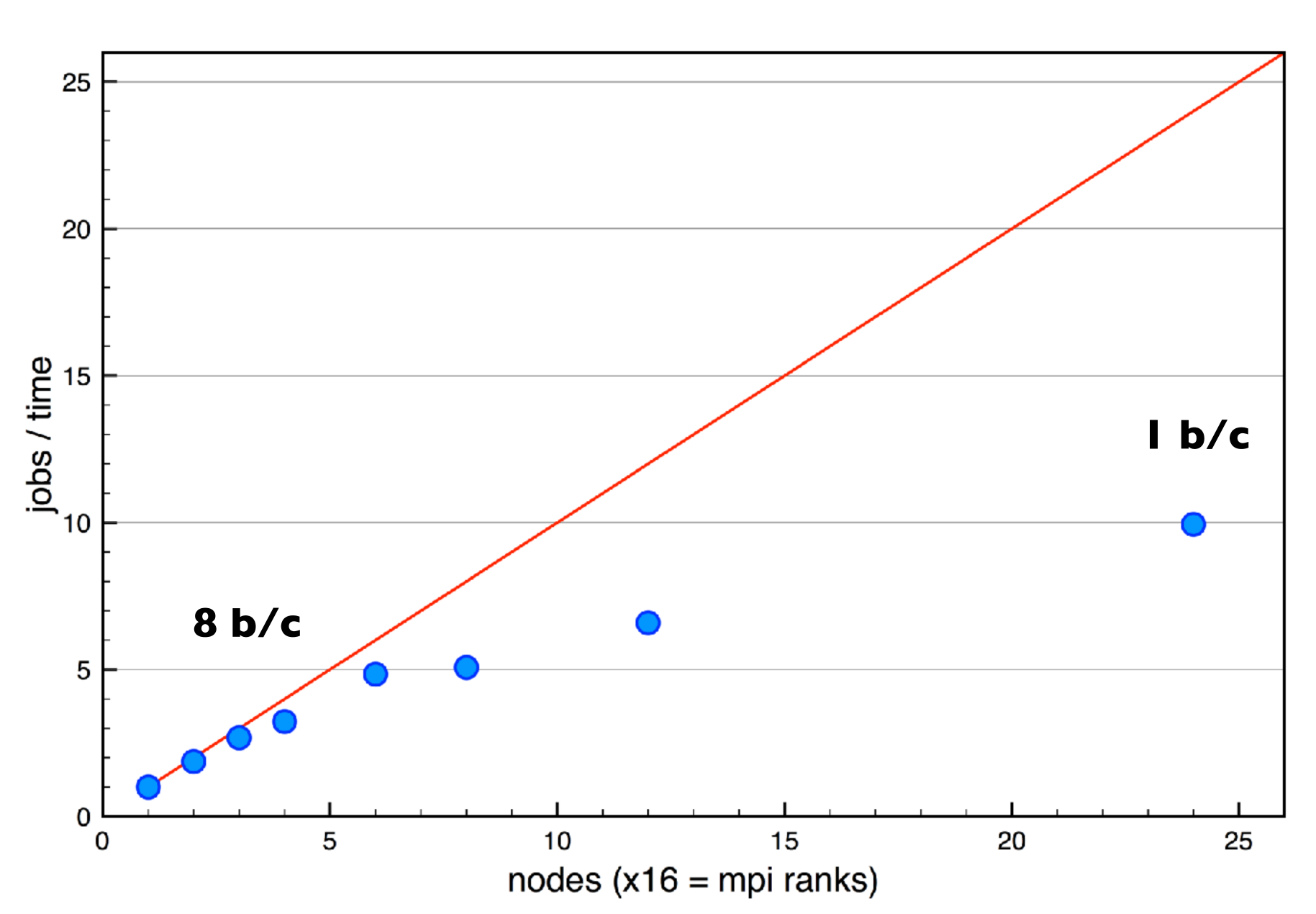
Kebnekaise: 28 c/node

GaAsBi 512 atoms, VASP PBE @Triolith (old)



NBANDS=1536 4 k-points

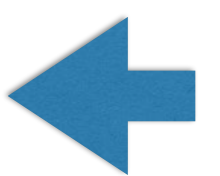
GaAsBi 128 atoms, VASP HSE06 @Triolith (old)



NBANDS=384 12 k-points

ALGO & NSIM

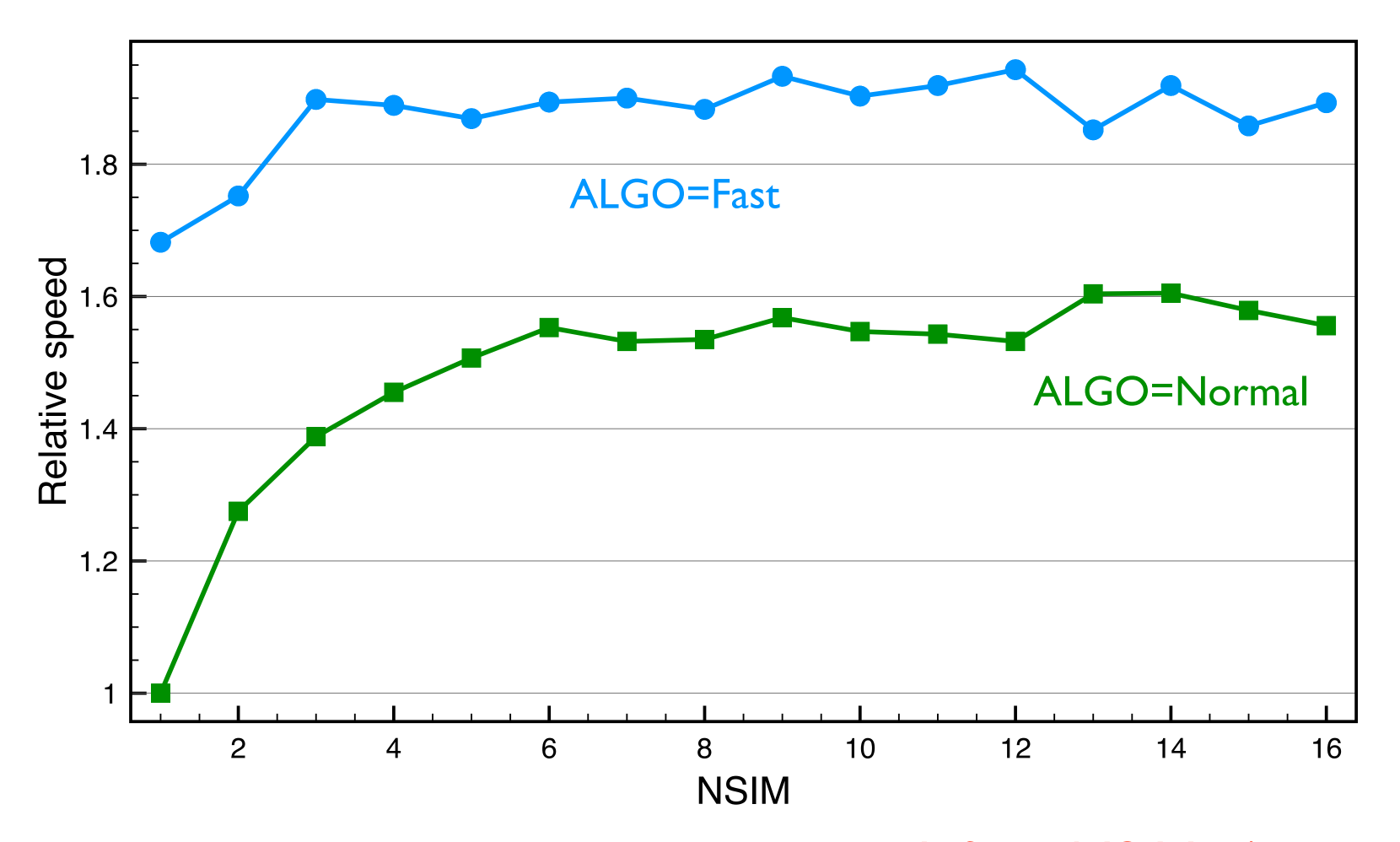
- Blocking mode for <u>RMM-DIIS</u> algorithm
- ALGO = Fast (Dav + R-D) / VeryFast (R-D)
- ALGO = Normal (<u>Davidson</u> algorithm), **default**
- not for hybrid-DFT, HSE06 (Damped, All, Normal)
- NSIM = 4, default



usually good (CPU)

- Kebnekaise/Tetralith: NSIM = 4 (or higher)
- Beskow: NSIM = 2

Si-H/Ag(111) 129 atoms, VASP PBE @Triolith (old)



default NSIM=4 seems OK here

NBANDS=750, 4 k-points

NCORE or NPAR (default)

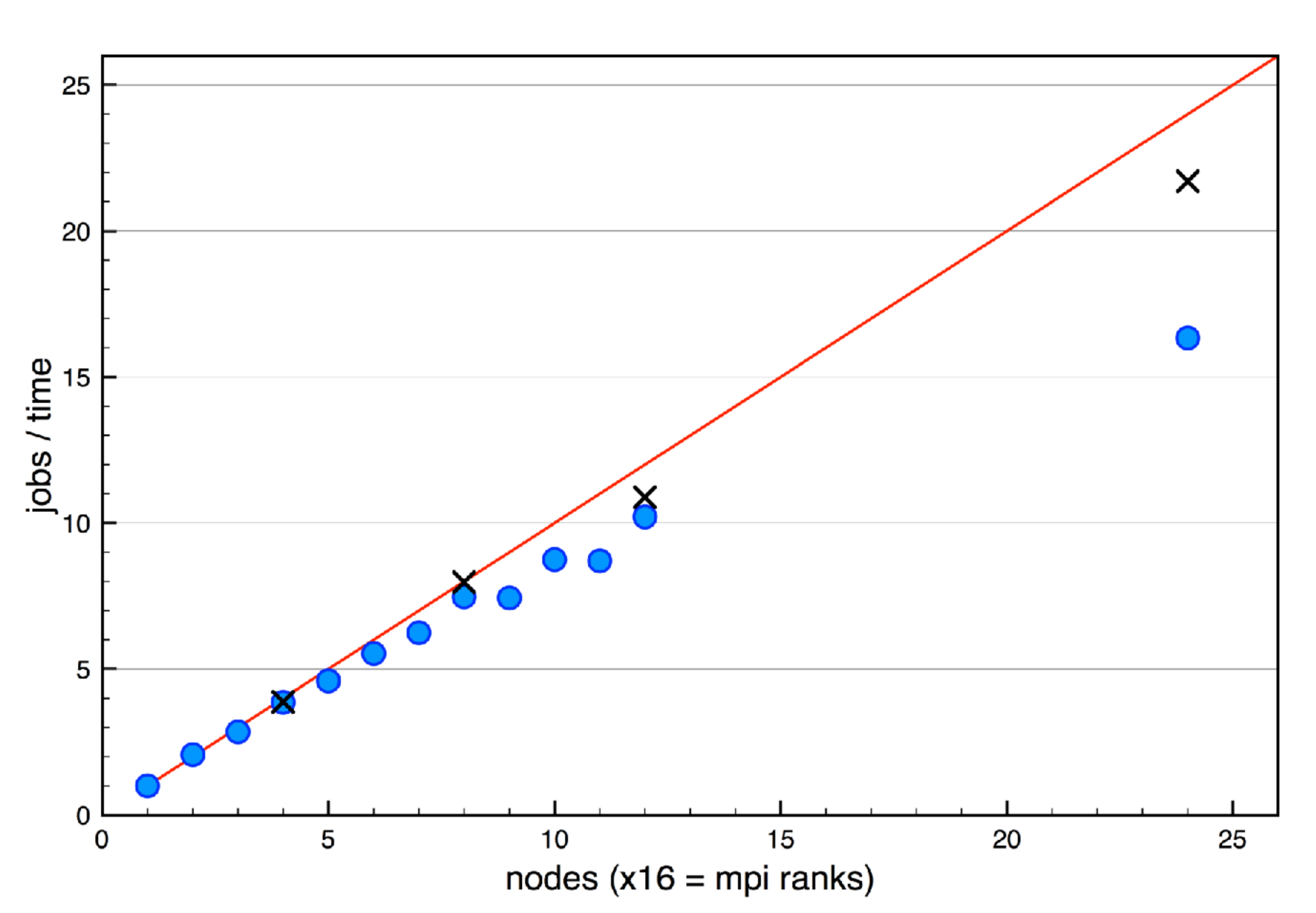
- cores per orbital / bands treated in parallel
- Davidson & RMM-DIIS algorithm
- ALGO = Normal & Fast, VeryFast
- NPAR = 1, saves memory
- NPAR = number of compute nodes
- NCORE = cores per compute node (or socket)

I find it easier to use NCORE, e.g. on Tetralith (if full node): NCORE=32 or 16

KPAR

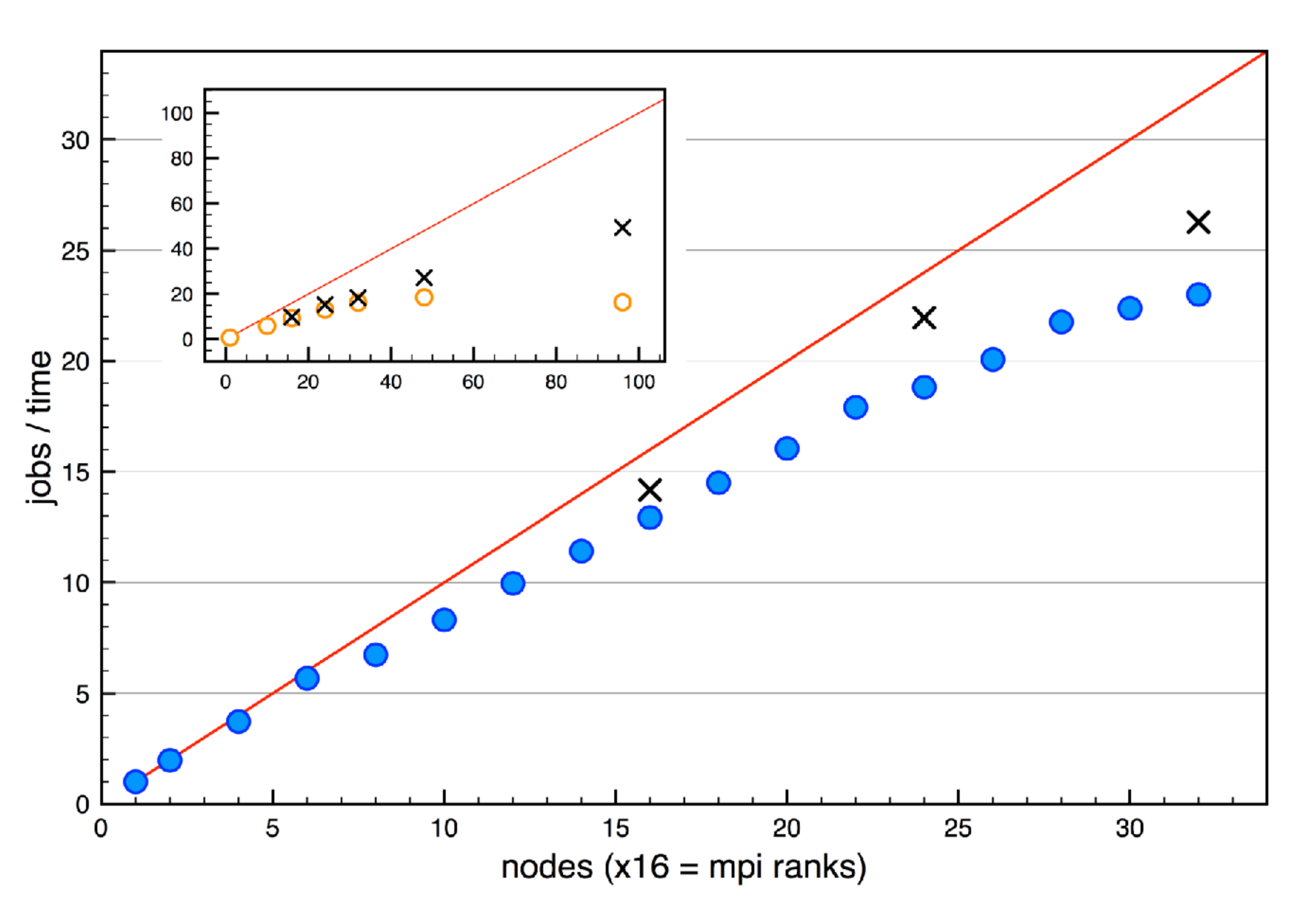
- KPAR = number of k-points treated in parallel
- in particular, good for hybrid-DFT jobs
- increase cores at least 2x
- try KPAR = min (nodes, k-points)

Si-H/Ag(111) 129 atoms, VASP PBE @Triolith (old)



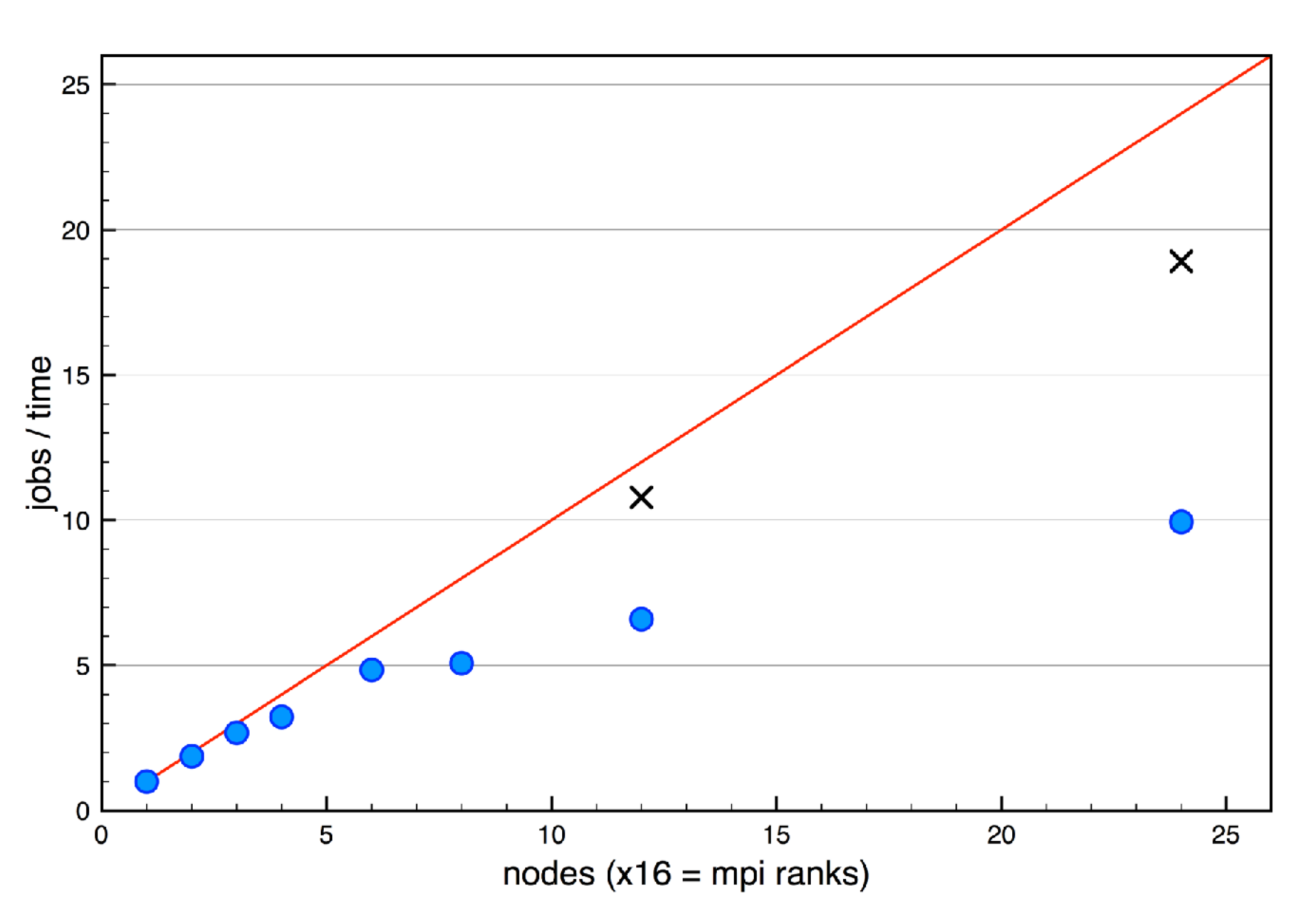
NBANDS=750 4 k-points

GaAsBi 512 atoms, VASP PBE @Triolith (old)



NBANDS=1536 4 k-points

GaAsBi 128 atoms, VASP HSE06 @Triolith (old)



NBANDS=384 12 k-points

Quick comparison

GaAsBi 512 atoms, VASP PBE, NBANDS = 1536, 4 k-points

@Tetralith, 6 nodes, NCORE=32, NSIM=30: 576s

4: 625s

© Kebnekaise, 7 nodes, NCORE=28, NSIM=30: 707s

4: 768s

retired @Beskow, 6 nodes, NCORE=32, NSIM=4: 1074s

24 2: 1593s

GaAsBi 128 atoms, VASP HSE06, NBANDS = 384, 3 k-points

@Kebnekaise, 6 nodes on 24c, NSIM=1: 1598s

retired @Beskow, 6 nodes on 24c, NSIM=1: 2146s

on 32c : 2044s

Peter Larsson's Tetralith scaling test

10

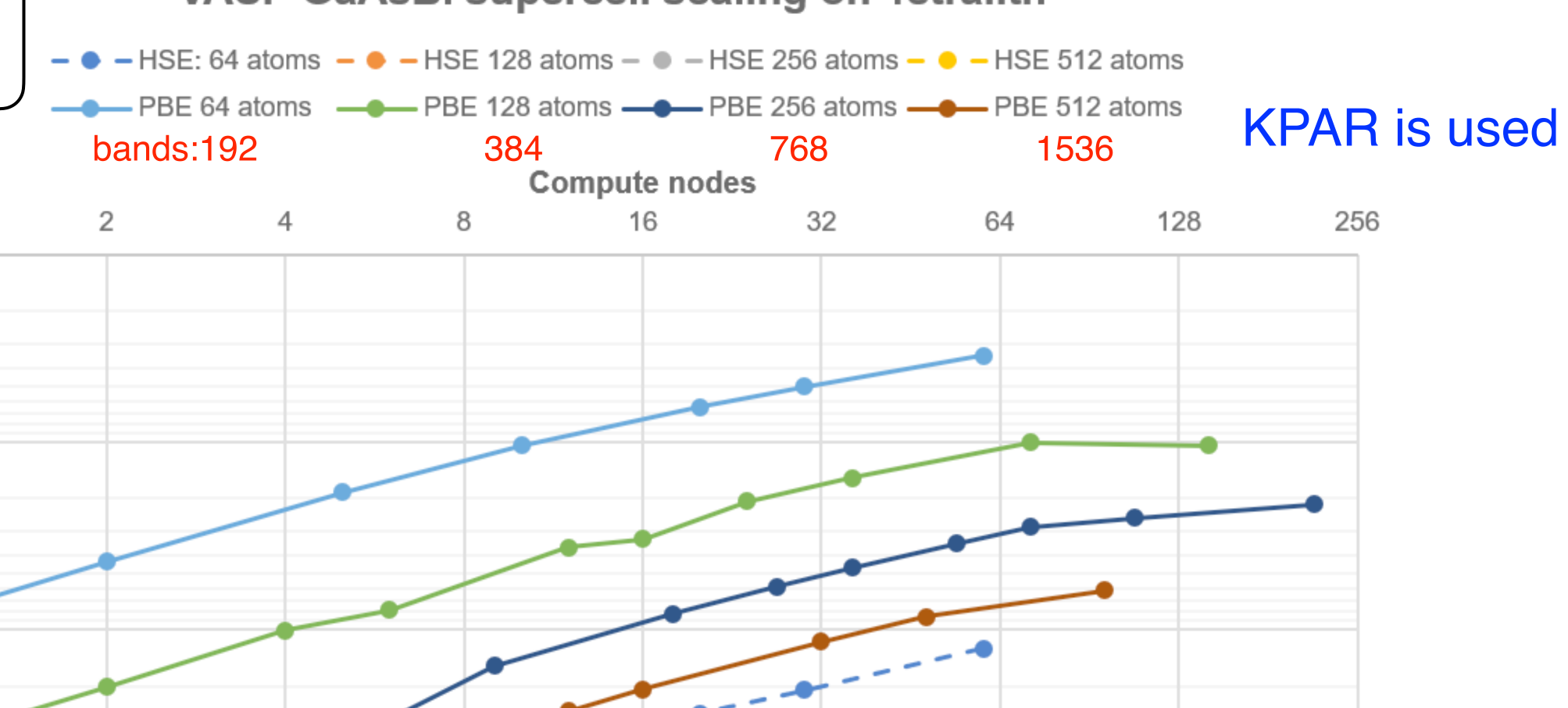
100

1000

10000

Time (s)

VASP GaAsBi supercell scaling on Tetralith



As a practical example, let us calculate how many core hours that would be required to run 10,000 full SCF cycles (say 100 geometry optimizations, or a few molecular dynamics simulations). The number of nodes has been chosen so that the parallel efficiency is > 90%:

Atoms	Bands	Nodes	Core hours
64	192	5	8,000
128	384	12	39,000
256	768	18	130,000
512	1536	16	300,000

The same table for 10,000 SCF cycles of HSE06 calculations looks like:

Atoms	Bands	Nodes	Core hours
64	192	10	400,000
128	384	36	2,000,000
256	768	36	6,900,000
512	1536	24	13,000,000

For comparison, a typical large SNAC project might have an allocation of 100,000-1,000,000 core hours per month with several project members, while a smaller personal allocation might be 5,000-10,000 core hours/month. Thus, while it is technically possible to run very large VASP calculations quickly on Tetralith, careful planning of core hour usage is necessary, or you will exhaust your project allocation.

Testing & Benchmarks, CPU & GPU

Example for HPC clusters

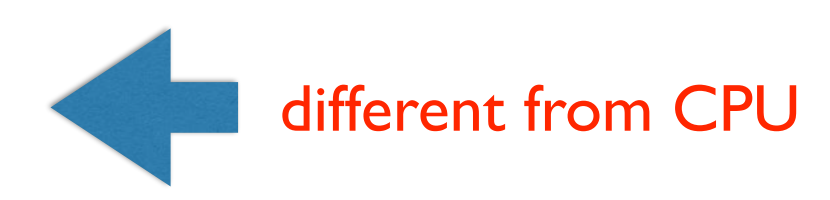


VASP testing @NSC

- From VASP6, test-suite included
- Peter Larsson's test-suite (old)
- Memory bandwidth important for VASP
- 10-100 runs for same job -> statistics
- Different builds (flags) for comparison
- Different (new) compiler suites
- Use collection of old problematic jobs... (BRMIX)

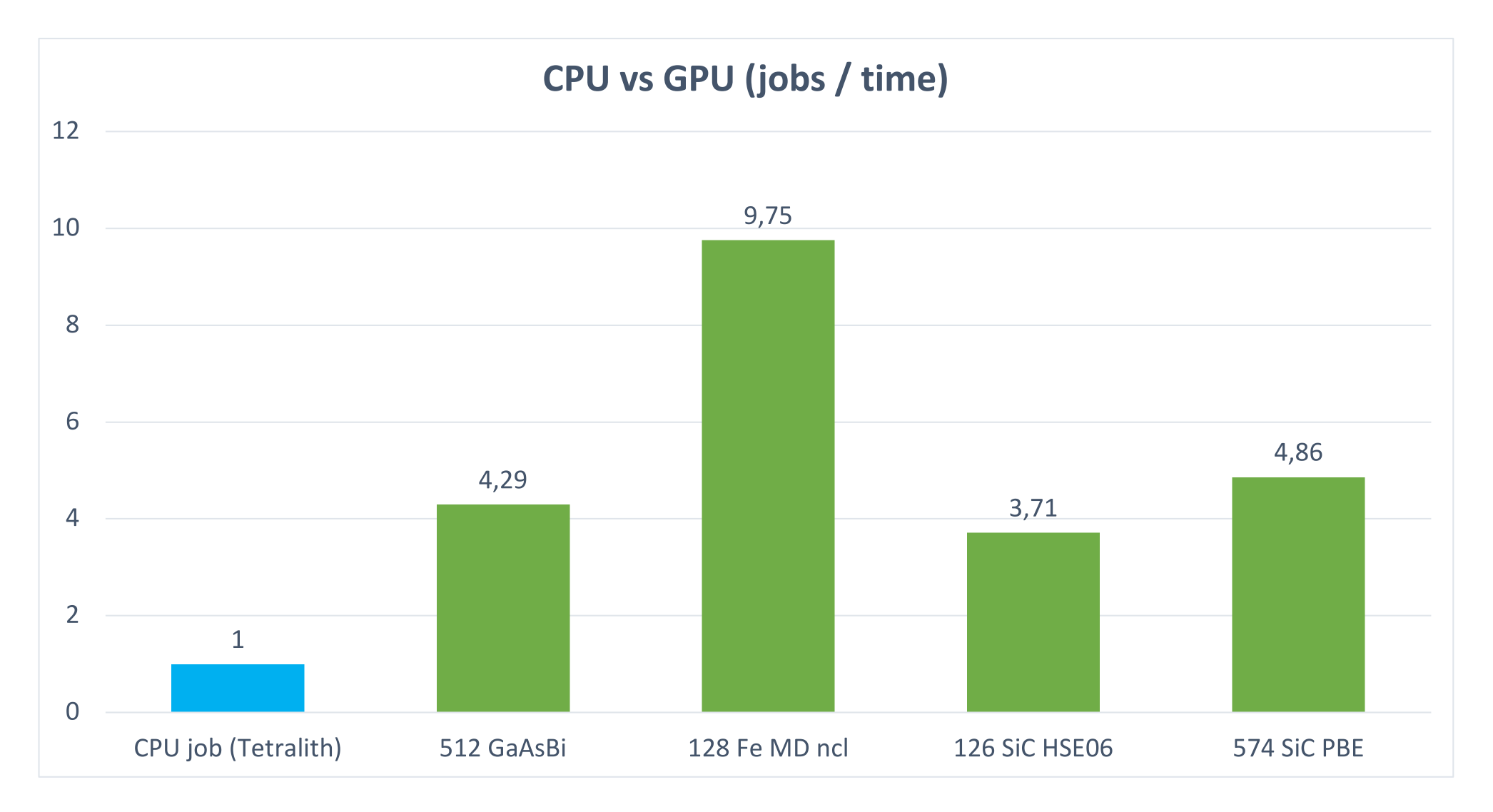
GPU

- VASP6 GPU OpenACC (old: CUDA)
- Different optimization than CPU
- !KPAR k-point parallelization ok
- NSIM very important!
 - e.g. test with NSIM = 16



- NCORE not supported
- GPU RAM possible bottleneck
- VASP6 faster with OpenACC, see link

Max throughput: number of jobs per time, 1(2) Tetralith nodes vs 1(2) A100 GPUs



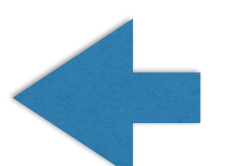
Problems & Summary

Different problems and their (possible) solution "Rules of thumb" summary



Possible problems

• Input related many kinds



- Memory (too little)
- Job size vs. allocation (mismatch)
 - Inefficient use (wasting core-hours)
- "Difficult" calculations (too costly, not possible)
- Bugs (compilers, old versions, ...)
 - sometimes from choice of compiler flags which in theory ought to be OK...

Memory (RAM) issues

starting with regular PBE - running HSE06, GW

- changing type of calcs.



2x2x2 k-mesh \rightarrow 4x4x4 k-mesh

x 8 k-points

- increasing the k-mesh

$$ENCUT = 400 eV \rightarrow ENCUT = 600 eV$$

 $\times 1.8$

- increasing energy cutoff

$$n_{pw} = \propto \text{ENCUT}^{3/2}$$

Memory (RAM) issues ...solutions

- Reduce cores/node, e.g. 24c/node, 16c/node
- More nodes (and reduce cores)
 #SBATCH --ntasks-per-node=16
 INCAR: NCORE=16
- Tetralith: use "fat" memory nodes
 #SBATCH -C fat
- Reduce k-mesh, ENCUT?
- Simplify system?
- Don't use —mem flag

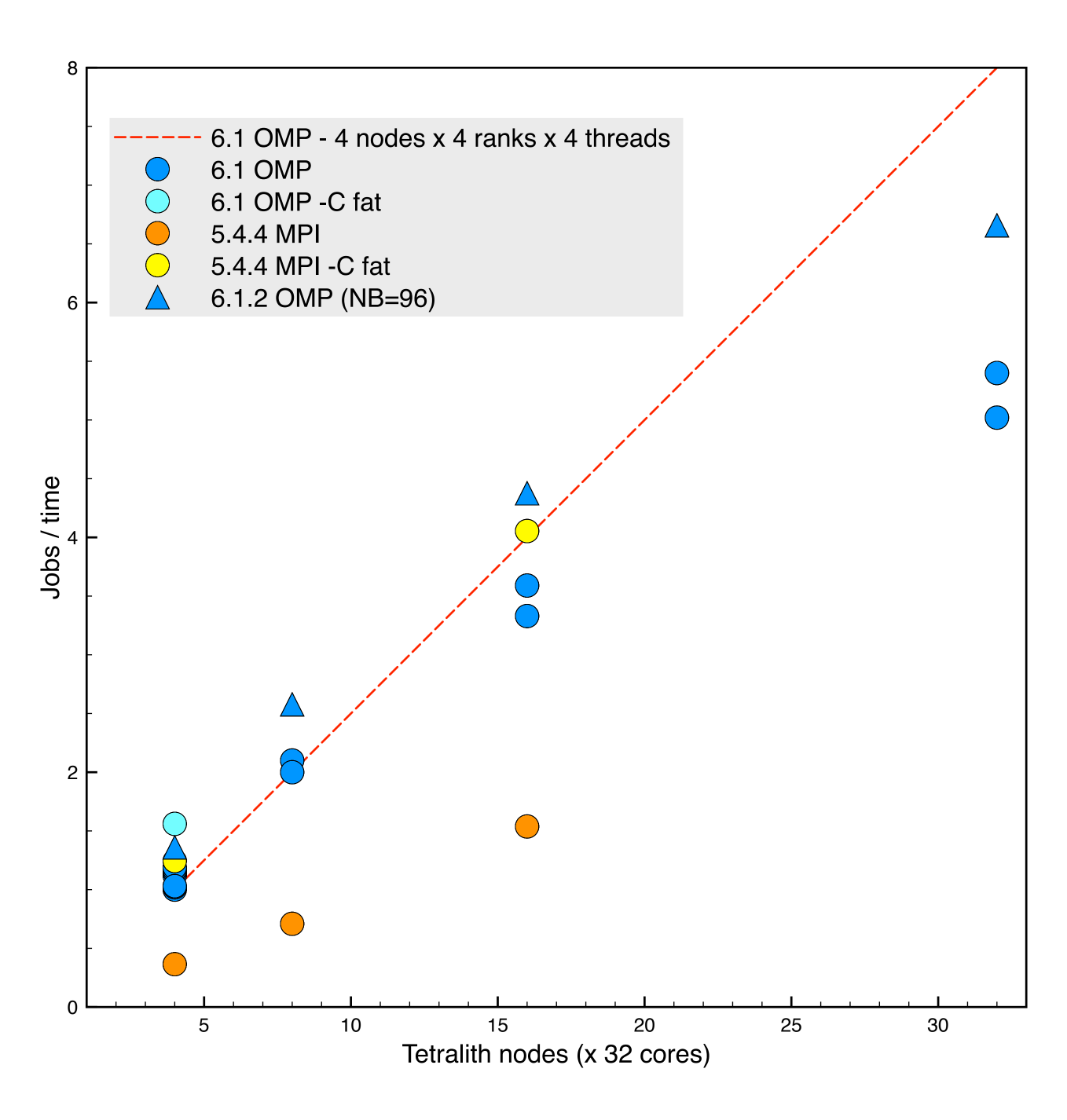
Memory (RAM) issues ...OpenMP?

- OpenMP availble from VASP6
- May combine with "fat" nodes, reducing cores, ...
- Test no. of threads

```
#!/bin/bash
#SBATCH -J vasp
#SBATCH -N 4
#SBATCH --ntasks-per-node=8
#SBATCH -t 4:00:00
#SBATCH -A naiss-xxx-yyy

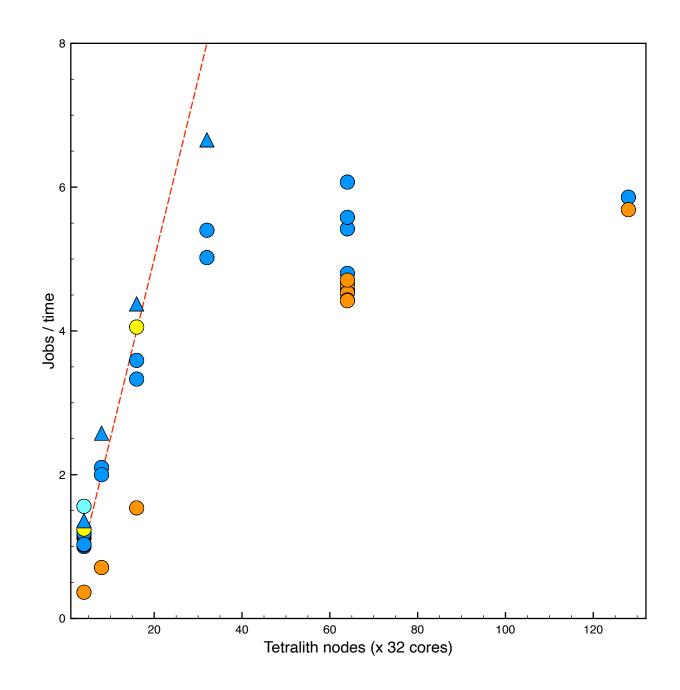
export OMP_NUM_THREADS=4
export OMP_STACKSIZE=512m

module add VASP/6.4.0.14022023-omp-nsc1-intel-2018a-eb
mpprun vasp_std
```



Notes:

Fe 2000 atoms PBE MD
Memory heavy job
VASP6 vs VASP5
OpenMP more efficient
"Fat" node -> 4x 96 GiB RAM
testing with NB=96 in src/scala.F



Warning/advice output

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

```
GGGG
        W A A R R NN N II NN
                                                   !!!
                       R N N N II N N N G
                                                   !!!
    W WW W AAAAAA RRRRR
                A R R N NN II N
                A R R N
                              N II N
                                             GGGG
                                                   !!!
ALGO=A and IALGO=5X tend to fail with the tetrahedron method
(e.g. Bloechls method ISMEAR=-5 is not variational)
please switch to IMSEAR=0-n, except for DOS calculations
For DOS calculations use IALGO=53 after preconverging with ISMEAR>=0
   I HOPE YOU KNOW, WHAT YOU ARE DOING
```

Common support cases

- complicated INCAR...
- structure (POSCAR)
- k-mesh (KPOINTS)
- NCORE/NPAR, KPAR
- VASP version
- cores
- memory

Common support cases

ALGO=N

- complicated INCAR... simplify & try again!
- structure (POSCAR) reasonable/correct?
- k-mesh (KPOINTS) \ \Gamma\text{-centered?}
- NCORE/NPAR, KPAR simplify (possibly remove)!
- VASP version try latest (possibly "vanilla" version)!

\$ module add VASP/5.4.4.16052018-nsc1-intel-2018a-eb

• cores too few/many?



memory
• ENCUT

k-mesh

larger memory nodes:

#SBATCH -C fat

reduce cores/node:

#SBATCH --ntasks-per-node=16

INCAR: NCORE=16

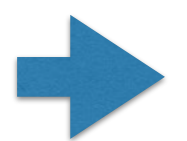
Notes / Reminders

- Same NBANDS when comparing Etot?
- Large enough NBANDS? e.g. increase for higher temp.
- Sufficient accuracy for your case?
- Use vasp_gam for 1 k-point calcs.
- LWAVE=.FALSE. WAVECAR might grow very large, don't output if not needed

Summary "rules of thumb"

- job size (max): total cores ≈ NBANDS / 8
- NSIM = 4 (default), or more
- NCORE = cores/node
- PREC = Accurate if forces important
- ENCUT = ENMAX x1.5 "max setting"
- KPAR = min (nodes, k-points) HSE06, especially useful
- In general, INCAR default settings OK
- GPU: important to increase NSIM

Resources



Wiki and Manual

Check in detail!

• Examples, tutorials

Presentations

Forum



Find all the links: https://vasp.at/

- Also other resources, materials and tools for VASP (see presentation 4.)
- Peter Larsson's old blog at NSC: https://www.nsc.liu.se/~pla/
- NSC VASP installations: https://www.nsc.liu.se/software/installed/tetralith/vasp/

Questions / trouble @NSC clusters? support@nsc.liu.se

VASP Refs.

- Good presentations by <u>Marsman</u> and <u>Blöchl</u>
- Blöchl PRB 50, 17953 (1994)
- Blöchl et al. https://arxiv.org/abs/cond-mat/0201015v2
- Kresse & Joubert PRB 59, 1758 (1999)
- Holzwarth et al. PRB 55, 2005 (1997)
- Martin, Electronic Structure, Chapter 11.1, 13.2
- https://vasp.at