

JUWELS & JURECA Tuning for the platform

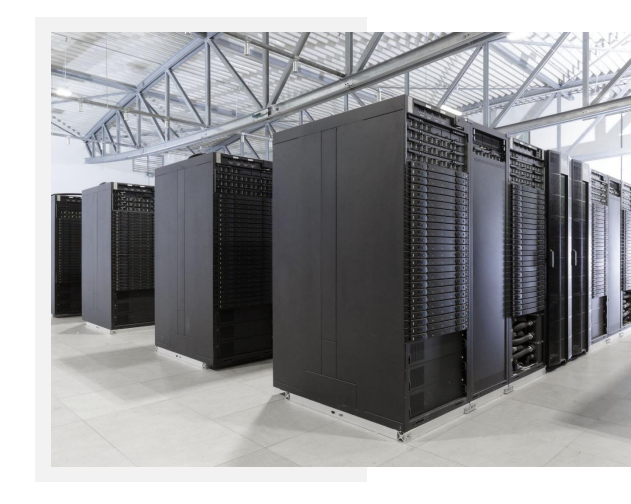


Usage of ParaStation MPI

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JUWELS & JURECA

Tuning for the platform

- 1. ParaStation MPI
- 2. Compiling your program
- 3. Running your program
- 4. Tuning parameters
- 5. Resources





History of ParaStation

- 1995: University project (→ University of Karlsruhe)
- 2005: Open source (→ ParaStation Consortium)
- Since 2004: Cooperation with JSC
 - various precursor clusters
 - DEEP-System (MSA prototype)
 - JuRoPA3 (J3)
 - JUAMS
 - JURECA (Cluster/Booster)
 - JUWELS (Cluster/Booster)
 - JURECA DC







ParaStation MPI



- based on MPICH (4.1.1)
 - supports all MPICH tools (tracing, debugging, ...)
- proven to scale up to 3,300 nodes and 136.800 procs per job running ParaStation MPI
 - JUWELS: No. 127 (Top500 Nov 2023)
 - JURECA DC: No. 82 (Top500 Nov 2023)
 - JUWELS Booster: No. 18 (Top500 Nov 2023)
- supports a wide range of interconnects, even in parallel
 - InfiniBand on JURECA DC and JUWELS
 - Omni-Path on JURECA Booster (deprecated)
 - Extoll on DEEP projects research systems (deprecated)
- tight integration with Cluster Management (e.g. healthcheck)
- MPI libraries for several compilers
 - especially for GCC and Intel

ParaStation MPI: Modularity

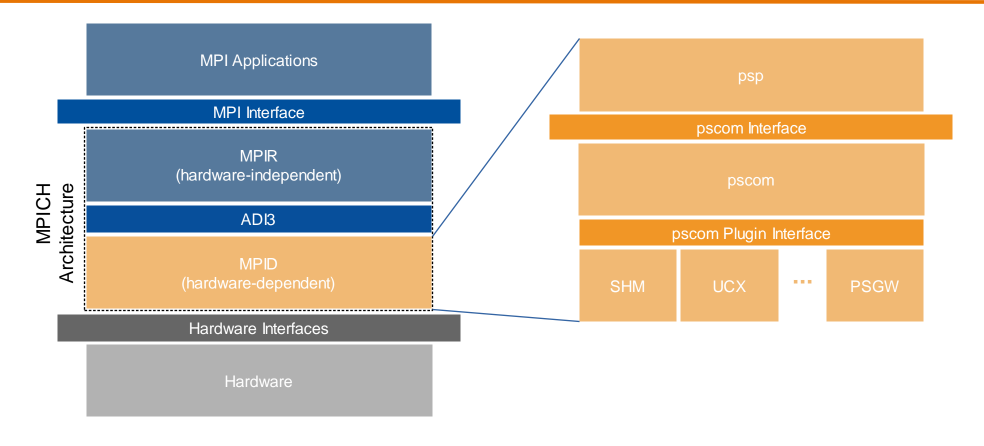


- 2 or more different modules with different hardware
- a job can execute dynamically on all modules
- you can pick the best out of all the worlds in a single job

- e.g. JURECA:
 - DC: AMD EPYC + Nvidia A100 + Infiniband
 - Booster: Intel KNL + Omni-Path
- how do these modules communicate with each other?

ParaStation MPI: pscom

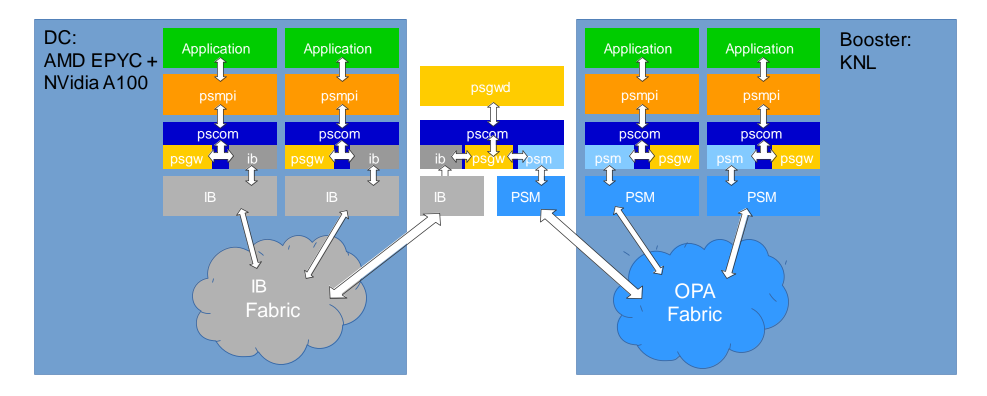




- low-level communication layer supporting various transports and protocols
- applications may use multiple transports at the same time

ParaStation MPI: pscom





- for the JURECA DC-Booster System, the ParaStation MPI Gateway Protocol bridges between Mellanox IB and Intel Omni-Path
- in general, the ParaStation MPI Gateway Protocol can connect any two low-level networks supported by pscom
- implemented using the psgw plugin to pscom, working together with instances of the psgwd

ParaStation MPI: Modular Jobs



- two processes communicate through a gateway, if they are not directly connected by a high-speed network (e.g. IB or OPA)
- static routing to choose a common gateway
- high-speed connections between processes and gateway daemons
- virtual connection between both processes through the gateway, transparent for application
- virtual connections are multiplexed through gateway connections
- further information: apps.fz-juelich.de/jsc/hps/jureca/modular-jobs.html

ParaStation MPI: CUDA awareness



- CUDA awareness supported by the following MPI APIs
 - Point-to-point (e.g. MPI SEND, MPI RECV, ...)
 - Collectives (e.g. MPI_Allgather, MPI_Reduce, ...)
 - One-sided (e.g. MPI Put, MPI Get, ...)
 - Atomics (e.g. MPI_Fetch_and_op, MPI_Accumulate, ...)
- CUDA awareness for all transports via staging
- CUDA optimization: UCX
- ability to query CUDA awareness at compile- and runtime

ParaStation MPI: CUDA awareness



- activate CUDA awareness by meta modules
 - default configurations
- query CUDA awareness:

Compiling on JUWELS



- currently MPI-4 version (5.9.2-1) available
- single thread tasks
 - module load Intel ParaStationMPI
 - module load GCC ParaStationMPI
- multi-thread tasks (mt)
 - module load Intel ParaStationMPI/5.9.2-1-mt
 - no multi-thread GCC version available
- ChangeLog available with
 - less \$(dirname \$(which mpicc))/../ChangeLog
- Gnu and Intel compilers available
- module spider for getting current versions
- see also the previous talk JUWELS Introduction

Wrapper



Wrappers

- mpicc (C)
- mpicxx (C++)
- mpif90 (Fortran 90)
- mpif77 (Fortran 77)
- when using OpenMP and the need to use the "mt" version, add
 - -fopenmp (GNU)
 - -qopenmp (Intel)

Did I use the wrapper correctly?



- libaries are linked at runtime according to LD_LIBRARY_PATH
- 1dd shows the libraries attached to your binary
- look for ParaStation libraries

```
ldd hello mpi:
libmpi.so.12 => /p/software/juwels/stages/2020/
software/psmpi/5.5.0-1-iccifort-2020.2.254-GCC-9.3.0/
lib/libmpi.so.12 (0x000015471ea43000)
VS.
libmpi.so.12 => /p/software/juwels/stages/2020/
software/psmpi/
5.5.0-1-iccifort-2020.2.254-GCC-9.3.0-mt/lib/
libmpi.so.12 (0x000014f110e58000)
```

JUWELS: start via srun



- use srun to start MPI processes
- srun -N <nodes> -n <tasks> spawns task
 - directly (-A <account>)
 - via salloc
 - from batch script via sbatch
- exports full environment
- stop interactive run with (consecutive) ^c
 - passed to all tasks
- no manual clean-up needed
- you can log into nodes which have an allocation/running job step
 - squeue -u <user>
 - sgoto <jobid> <nodenumber>
 - e.g. sgoto 2691804 0



```
/* C Example */
#include <stdio.h>
#include <mpi.h>
int main (int argc, char **argv) {
  int numprocs, rank, namelen;
  char processor name[MPI MAX PROCESSOR NAME];
 MPI Init (&argc, &argv);
 MPI Comm rank (MPI COMM WORLD, &rank);
 MPI Comm size (MPI COMM WORLD, &numprocs);
 MPI Get processor name (processor name, &namelen);
 printf ("Hello world from process %d of %d on %s\n",
         rank, numprocs, processor name);
 MPI Finalize ();
  return 0;
```

Running on JUWELS (Intel chain)



- module load Intel
- module load ParaStationMPI
- mpicc -03 -o hello_mpi hello_mpi.c
- Interactive:
- salloc -N 2 -A partec # get an allocation
- srun -n 2 ./hello_mpi

Hello world from process 0 of 2 on jwc08n188.juwels Hello world from process 1 of 2 on jwc08n194.juwels

- Batch:
- sbatch ./hello mpi.sh
- Increase verbosity:
 - PSP_DEBUG=[1,2,3,...] srun -n 2 ./hello_mpi

Process Placement



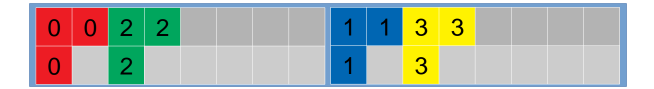
- ParaStation process pinning:
 - avoid task switching
 - make better use of CPU cache and memory bandwidth
- JUWELS is pinning by default:
 - so --cpu-bind=threads may be omitted
- manipulate pinning:
 - e.g. for "large memory / few task" applications
- manipulate via
 - --cpu-bind=threads|sockets|cores|mask_cpu:<mask1>,<mask2>,...
 - CPU masks are always interpreted as hexadecimal values
 - --distribution=*|block|cyclic|arbitrary|plane=<options>
 [:*|block|cyclic|fcyclic[:*|block|cyclic|fcyclic]][,Pack|NoPack]
- further information: https://apps.fz-juelich.de/jsc/hps/juwels/affinity.html

Process Placement



• Example:

- --ntasks-per-node=4
- --cpus-per-task=3
- --cpu-bind=threads



● --cpu-bind=mask_cpu:0x7,0x700,0xE0,0xE000



Process Placement



- best practice depends not only on topology, but also on characteristics of application:
- putting threads far apart is
 - improving the aggregated memory bandwidth available to your application
 - improving the combined cache size available to your application
 - decreasing the performance of synchronization constructs
- putting threads close together is
 - improving the performance of synchronization constructs
 - decreasing the available memory bandwidth and cache size

Hybrid MPI/OpenMP



```
#include <stdio.h>
#include <mpi.h>
                                                              Example:
#include <omp.h>
                                                              2 Nodes, 2x2 Procs,
int main(int argc, char *argv[]) {
                                                              2x2x24 Threads
 int numprocs, rank, namelen;
 char processor name[MPI MAX PROCESSOR NAME];
 int iam = 0, np = 1;
                                                                Node x
                                                                              Node y
 MPI Init(&argc, &argv);
 MPI Comm size (MPI COMM WORLD, &numprocs);
 MPI Comm rank (MPI COMM WORLD, &rank);
 MPI Get processor name (processor name, &namelen);
#pragma omp parallel default(shared) private(iam, np)
   np = omp get num threads();
   iam = omp get thread num();
   printf("Hello from thread %02d out of %d from process %d out of %d on %s\n",
          iam, np, rank, numprocs, processor name);
 MPI Finalize();
```

On JUWELS



- module load Intel ParaStationMPI/5.9.2-1-mt
- salloc -N 2 -A partec -cpus-per-task=24
- export OMP_NUM_THREADS=\${SLURM_CPUS_PER_TASK}
- srun -n 4 ./hello hybrid | sort

```
Hello from thread 00 out of 24 from process 0 out of 4 on jwc01n238.juwels
Hello from thread 00 out of 24 from process 1 out of 4 on jwc01n238.juwels
Hello from thread 00 out of 24 from process 2 out of 4 on jwc01n247.juwels
Hello from thread 00 out of 24 from process 3 out of 4 on jwc01n247.juwels
Hello from thread 01 out of 24 from process 0 out of 4 on jwc01n238.juwels
Hello from thread 01 out of 24 from process 1 out of 4 on jwc01n238.juwels
Hello from thread 01 out of 24 from process 2 out of 4 on jwc01n247.juwels
Hello from thread 01 out of 24 from process 3 out of 4 on jwc01n247.juwels
Hello from thread 23 out of 24 from process 0 out of 4 on jwc01n238.juwels
Hello from thread 23 out of 24 from process 1 out of 4 on jwc01n238.juwels
Hello from thread 23 out of 24 from process 2 out of 4 on jwc01n247.juwels
Hello from thread 23 out of 24 from process 3 out of 4 on jwc01n247.juwels
```

Pinning



• JUWELS:

- 2 Sockets, 24 Cores per Socket
- 2 HW-Threads per Core
- − → 96 HW-Threads possible
- normally (SMT):
 - HW-Threads 0-23, 48-71 → CPU0
 - HW-Threads 24-47, 72-95 → CPU1

"Package"

Node										
	Socket 0					Socket 1				
Core 0	Core 1		Core 22	Core 23	Core 24	Core 25		Core 46	Core 47	
HWT 0	HWT 1		HWT 22	HWT 23	HWT 24	HWT 25		HWT 46	HWT 47	
HWT 48	HWT 49		HWT 70	HWT 71	HWT 72	HWT 73		HWT 94	HWT 95	

Pinning



• JURECA DC:

- 2 Sockets, 64 Cores per Socket
- 2 HW-Threads per Core
- → 256 HW-Threads possible
- normally (SMT):
 - HW-Threads 0-63, 128-191 → CPU0
 - HW-Threads 64-127, 192-255 → CPU1

"Package"

Node										
	Socket 0					Socket 1				
Core 0	Core 1		Core 62	Core 63	Core 64	Core 65		Core 126	Core 127	
HWT 0	HWT 1	÷	HWT 62	HWT 63	HWT 64	HWT 65		HWT 126	HWT 127	
HWT 128	HWT 129		HWT 190	HWT 191	HWT 192	HWT 193		HWT 254	HWT 255	

Pinning



- no thread pinning by default on JURECA and JUWELS
- allow the Intel OpenMP library thread placing
 - export KMP_AFFINITY=[verbose, modifier,...]
 - compact: place threads as close as possible
 - scatter: as evenly as possible
- full environment is exported via srun on JURECA and JUWELS
- for GCC: set GOMP CPU AFFINITY (see manual)

Large Job Considerations



- every MPI process talks to all others:
 - (N-1) x 0.55 MB communication buffer space per process!
- example 1 on JUWELS:
 - job size 256 x 96 = 24,576 processes
 - 24,575 x 0.55 MB \rightarrow ~ 13,516 MB / process
 - x 96 processes / node \rightarrow ~ 1,267 GB communication buffer space
 - but there is only 96 GB of main memory per node
- example 2 on JURECA DC:
 - job size 256 x 256 = 65,536 processes
 - 65,535 x 0,55 MB \rightarrow ~ 36,044 MB / process
 - x 256 processes / node \rightarrow ~ 9,011 GB communication buffer space
 - but there is only 512 GB of main memory per node

On Demand / Buffer Size



- Three possible solutions:
- 1. Try using alternative meta modules
- 2. Create buffers on demand only:
 - export PSP_ONDEMAND=1
 - activated by default!
- 3. Reduce the buffer queue length:
 - (default queue length is 16)
 - export PSP_OPENIB_SENDQ_SIZE=3
 - export PSP OPENIB RECVQ SIZE=3
 - do not go below 3, deadlocks might occur!
 - trade-off: performance penalty
 - (sending many small messages)

16k

16k

16k

16k

queue length

On Demand / Queue Size Guidelines



- On-Demand works best with nearest neighbor communications
 - (Halo) Exchange
 - Scatter/Gather
 - All-reduce
 - ...
- but for All-to-All communication:
 - queue size modification only viable option...
- example

```
rank 0: for (;;) MPI_Send ()
rank 1: for (;;) MPI_Recv ()
• PSP_OPENIB_SENDQ/RECVQ_SIZE=4: 1.8 seconds
• PSP_OPENIB_SENDQ/RECVQ_SIZE=16: 0.6 seconds
• PSP_OPENIB_SENDQ/RECVQ_SIZE=64: 0.5 seconds
```

Resources



- www.par-tec.com
- www.fz-juelich.de/en/ias/jsc/systems/supercomputers
- opt/parastation/doc/pdf
- by mail: sc@fz-juelich.de
- by mail: support@par-tec.com
- download ParaStation MPI at github:
 - https://github.com/ParaStation/psmgmt
 - https://github.com/ParaStation/pscom
 - https://github.com/ParaStation/psmpi

Summary



- you now should be able to
 - compile
 - run your application
 - tune some runtime parameters
 - diagnose and fix specific errors
 - know where to turn to in case of problems

Thank you for your attention!



Questions?