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Application Performance Evaluation



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Executive Summary: As documented in the original proposal, the Energy oriented Centre of Excellence (EoCoE) is a user driven consortium dedicated to tackling modelling challenges in the field of renewable energy. Consequently, the implementation and organization of the project places High Performance Computing (HPC) applications of the four chosen user communities at the heart of the project.

In the EoCoE context, which requires close collaboration between domain scientists and HPC experts, application performance evaluation is a key instrument: it permits the initial status of an application code to be objectively determined before more detailed examination or modification; enables monitoring of the impact of each code modification during the optimization process; and quantitative assessment of the impact of such support activity. This deliverable report describes the performance metrics definition, the first four-day workshop event bringing together HPC experts and application scientists, and a progress summary of the first seven codes evaluated.



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1. Motivation

Within in its transversal basis (WP1), the EoCoE project has gathered a comprehensive range of HPC expertise that aims to enhance the performance of applications from the four domain pillars, thereby enabling them to effectively exploit the existing European computing infrastructure. Close interaction between WP1 and the application domains WP2-WP5 is a key feature of EoCoE, with the ultimate goal of expediting advances in simulations of low-carbon energy systems and technology.

In this context, application performance evaluation is an instrument of key importance, since it permits us to:

- 1. define the status of an application code at the moment when EoCoE HPC experts start to examine it,
- 2. monitor the impact of each code modification during the optimization process,
- 3. quantitatively assess the impact of such support activity when it comes to an end.

This deliverable report describes the status of performance evaluation activity over the first 6 months of the project, beginning with a dedicated workshop for this purpose, and various follow-up actions such as Section 3, which presents the definition of the EoCoE performance evaluation report and the performance metrics it uses; Subsection 3.4, on the establishment of an automated and reproduceable process that delivers all the required metrics; Section 4, which describes the system for monitoring progress in application optimisation.



2. First joint EoCoE-PoP benchmarking workshop

The first EoCoE-POP workshop on benchmarking and performance analysis brought together code developers of community codes associated with WP 2-5 with HPC experts associated with WP 1 and HPC experts from the CoE "POP". The goal of this 4-day event held at Jülich Supercomputing Centre from 8th-11th December, 2015 was to familiarise the developers from WP2-5 with state-of-the-art HPC performance analysis tools, enabling the teams to make a preliminary identification of bottlenecks, and to initiate the standardisation of benchmark procedures for these codes within the EoCoE project. The workshop comprised 4.5 hours of presentations on the benchmarking and performance tools followed by 12 hours of hands-on work supervised by the WP1 and PoP HPC experts.



Figure 1: Workshop participants and support activity during the first benchmarking workshop

As an initial step, all code developers were instructed on how to perform benchmarking within the JUBE¹ workflow environment, which will permit measurements to be documented, shared and rigorously reproduced over the project lifetime and beyond. Developers were then able to begin analysing their applications using specific HPC tools under the guidance of HPC experts (Score-P, Scalasca, Vampir, Paraver, Extrae, Darshan, VTune and others). Based on this face-to-face collaboration and common training, small teams of code developers and HPC experts from WP 1 were established, who have begun to follow up on the promising initial work to provide comprehensive benchmarks and performance data by the time the next workshop is held in June.

Each of the participating developer teams was allocated a WP1 mentor, tasked with assisting any follow-up benchmarking and tuning work, and acting as an initial contact point for enquiries going beyond the initial assessment (I/O issues, data management, visualisation etc). A summary of the participating codes is given in table 1. Four of these (ALYA, Metallwalls, PARFLOW and Gysela) belong to the set of codes already prioritised (triggered) for WP1 optimisation activity.

A further valuable outcome was the exchange of respective ideas and needs between code developers and HPC experts, as this helped clarifying the issues from either perspective and enabled both sides to interact more smoothly with a well defined focus on the next actions to be taken. For example, the requirements for a full code 'audit' from the EoCoE and POP perspectives were clarified: here it was decided that the initial benchmarking would take place within and immediately after the workshop by EoCoE WP1 members, whereas more in-depth follow-up analyses could be channelled via a formal request to POP

¹www.fz-juelich.de/jsc/jube

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WP	Context	Code	Developer	WP1 contact
2	Wind farms	ALYA	Houzeaux (BSC)	Ould-Rouis (MdlS)
2	Ensemble forecasting	ESIAS	Bernd (FZJ)	Lührs (JSC)
3	Photovoltaics	PVnegf	Aeberhard (FZJ)	Napoli (JSC)
3	Materials	Metallwalls	Salanne	Haefele (MdlS)
4	Hydrology	PARFLOW	Kollet (FZJ)	Sharples
4	Geothermics	SHEMAT	Qu (RWTH)	Sharples
5	Plasma transport	Gysela	Latu (MdlS)	Guillame Latu

Table 1: Codes participating in first EoCoE benchmarking workshop

at a later stage.



3. EoCoE performance evaluation report and metrics definition

Performance evaluation has the obvious purpose to uncover bottlenecks and possibly other technical areas of improvement for the codes under consideration. In order to verify the impact and success of code changes it is mandatory to apply it *iteratively and continuously* in a regular manner. In particular, it is *not* sufficient to analyse a code once and from the results create an optimised version of a code in a single step.

3.1 Organizational structure and reporting

The EoCoE management has carefully engineered a lean yet efficient organisational structure which ensures that such an ongoing and continuous process involving code developers and HPC-experts can be achieved and monitored, with a minimum of bureaucratic overhead. The elements and ingredients for this collaborative micro-community are

- 1. Permanent code teams, consisting of at least one developer and one HPC-experts, to corroborate the collaboration between in a sustainable manner.
- 2. Code identity card filled by the application developer to initiate the analysis.
- 3. A well-defined set of global performance metrics to have a common perspective on progress and development. Ideally, most of the initial measures are obtained during an EoCoE performance workshop.
- 4. The possibility to add further application-specific performance metrics if necessary.
- 5. A Code Diary for each code that allows to track and document the evolution.
- 6. A technical infrastructure based on Git which allows all code teams to share their reports and to provide a basis from which best practice methods can be deduced.

Appendix A shows the full information for the two most advanced codes in this performance evaluation process: Metalwalls and ESIAS.

3.2 Performance tools

Several tools are used to extract the metrics described in section 3.3. The following list provides the ones that are currently used in the EoCoE performance evaluation process.

- Alinea performance report² provides a nice performance overview of a code with information gathered from a single run
- The UNIX *time* command: wall time measurement of the duration of an application
- Darshan³ provides IO measurements
- Score-P/Scalasca⁴ provides application profiling and trace based on source code instrumentation methods as well as trace analysis tools
- Vampir trace analysis⁵ provides trace visualization tools

²http://www.allinea.com/products/allinea-performance-reports ³http://www.mcs.anl.gov/research/projects/darshan/

⁴http://www.scalasca.org/

⁵https://www.vampir.eu/



- Extrae / Paraver⁶ provides application profiling and trace based on runtime hardware counter sampling methods as well as trace visualization tools
- PAPI⁷, used through Scalasca, provides hardware counters measurements
- SLURM⁸ scheduling system is able to provide the memory footprint of the first MPI rank of the application
- IdrMem⁹ library is used to retrieve the memory footprint on systems where Slurm is not available.
- Intel VTune¹⁰ provides application profiling and trace based on runtime hardware counter sampling methods as well as trace visualization tools. It is also the only tool able to extract the measure of the memory throughput on recent Intel CPUs
- Intel Vectorization Advisor¹¹ is used to evaluate the quality of the code vectorization

3.3 Metrics definition

The definition of all global performance metrics is given in the table 2.

Metrics Global.1, Global.2 and Global.3 might exhibit some inconsistencies as these three measures are extracted from three different runs performed with different binaries. This should not change the global picture as long as similar run times are observed for these three runs. These inconsistencies might also impact metric Node.2 as computations involve these three measures.

<u>Memory vs Compute Bound metric (Global.4)</u> is computed with the runtime coming out of two dedicated runs. The two runs uses the same amount of MPI ranks and threads but on twice the number of nodes. This leads to depleted resources, and, by using specific deployments, one has the chance to observe memory bandwidth effects. Typically on current dual socket systems, a compact and a scatter run are performed. The compact run packs all the MPI processes and threads on a single socket, whereas the scatter run distributes them evenly on the two sockets. Going from the compact run to the scatter one, the available computing power is kept constant while doubling the available memory bandwidth. As a consequence, if both runs exhibit the same wall time, this means that the memory bandwidth available has no impact on the application. So the code is strongly compute bound and the ratio run time compact / run time scatter is 1.0. On the other hand, if the scatter run is twice as fast, the ratio is than 2.0 and this means that the code is strongly memory bound.

<u>The load imbalance metric</u> gives the potential for code improvement if the load imbalance would be perfectly fixed. Thanks to the trace analysis, Scalasca is able to compute the critical path of the application and the overhead due to load imbalances between ranks/threads. The metric used here is simply the ratio overhead / critical path. For instance, if a 20% load imbalance is measured, fixing perfectly this load imbalance would

⁶http://www.bsc.es/computer-sciences/performance-tools/paraver

⁷http://icl.cs.utk.edu/papi/

⁸http://slurm.schedmd.com/

 $^{^9 \}tt https://gitlab.maisondelasimulation.fr/dlecas/IdrMem$

¹⁰https://software.intel.com/en-us/intel-vtune-amplifier-xe

¹¹https://software.intel.com/en-us/intel-advisor-xe



		Metric name	Definition	Measure
	1	Total Time (s)	Total application wall time	time
ba	2	Time IO (s)	Average time spent in doing IO for each	Darshan
Ē			process	
	3	Time MPI (s)	Average time spent in MPI for each process	Scalasca
	4	Memory vs Compute Bound	1.0 means strongly compute bound, 2.0	cf text
			means strongly memory bound	
	1	IO Volume (MB)	Total amount of data read and written	Darshan
0	2	Calls (nb)	Total number of IO calls	Darshan
	3	Throughput (MB/s)	IO.1 / Global.2	Computed
	4	Individual IO Access (kB)	IO.1 / IO.2	Computed
	1	P2P Calls (nb)	Average number of peer to peer communi-	Scalasca
			cations per MPI rank	
	2	P2P Calls (s)	Average time spent in peer to peer com-	Scalasca
IPI			munications per MPI rank	
	3	Collective Calls (nb)	Average number of collective communica-	Scalasca
			tions per MPI rank	
	4	Collective Calls (s)	Average time spent in collective communi-	Scalasca
			cations per MPI rank	
	5	Synchro / Wait MPI (s)	Average time spent in synchronization per	Scalasca
			MPI rank	
	6	Ratio Synchro / Wait MPI	MPI.5 / Global.3	Computed
	7	Message Size (kB)	Average message size sent	Scalasca
	8	Load Imbalance MPI	MPI load imbalance	Scalasca
le	1	Time OpenMP (s)	Time spent in OpenMP parallel region	Scalasca
Noc	2	Ratio OpenMP	Node.1 / (Global.1 - Global.2 - Global.3)	Computed
	3	Time Synchro / Wait OpenMP	Average time spent in synchronization per	Scalasca
			thread	
	4	Ratio Synchro / Wait OpenMP	Node.4 / Node.1	Computed
	5	Load Imbalance OpenMP	OpenMP load imbalance	Scalasca
В	1	Memory Footprint	Average memory footprint of an MPI pro-	IdrMem/
Me			cess	Slurm
	2	Cache Usage Intensity	Cache Hit / (Cache Hit + miss) in Last	PAPI
			Level Cache	
	3	RAM Avg Throughput (GB/s)	Average memory throughput per socket	Vtune
	1	IPC	Total number of instructions executed /	PAPI
e			Total number of cycles	
Cor	2	Runtime without vectorization	Total application wall time compiled with	time
			vectorization disabled	
	3	Vectorisation efficiency	Global.1 / Core.2	Computed
	4	Runtime without FMA	Total application wall time when compiled	time
			with FMA disabled	
	5	FMA efficiency	Global.1 / Core.4	Computed

Table 2: Global performance metrics definition



improve the performance of the code by 20%.

3.4 Towards an automated metrics extraction process

To make it easier for all code teams to carry out performance evaluation of their application themselves, without the need for detailed familiarisation of the tools, it was decided to strive for an automatic generation of as many metrics in table 2 as possible. Two codes, out of the first workshop - Metalwalls and Esias - use already a very extended automatisation process. Also the other codes already included several profiling tools within a automated JUBE script. Section 4 gives an overview about the status of all codes involved in the first workshop.

So far, the following tools have been integrated into an automated process:

- The UNIX time command is used to measure total application wall time
- Darshan provides all metrics concerning IO
- Scalasca provides all metrics concerning MPI, OpenMP and load balancing
- PAPI is used through Scalasca and provides all performance counters
- VTune is the only tool able to extract the measure of the memory throughput on recent hardware
- SLURM scheduling system is able to retrieve the memory footprint of the first MPI rank of the application.
- IdrMem library is used to retrieve the memory footprint on systems where Slurm is not available.

There is the option to add more tools along the same lines. This is work in progress.

The code team of Metalwalls has dedicated themselves to set up a comprehensive and well documented example of how this can be done. Thanks to very intensive collaborative efforts, such a process has been successfully implemented and has proven its value. The code team created scripts to extract the relevant metrics out of the different profiling tools result files and allow the integration of these metrics into the JUBE environment. Thus, it has the potential to serve as a best practice anchor for other code teams and can thereby strongly leverage the overall work within EoCoE, even more so since this achievement was reached very early, not even six months into the project.

Specifically, for the purpose of automation, four separate code binaries are initially needed:

- Normal (bin)
- scalasca instrumented (scalasca)
- Normal plus "no-vectorization" (bin-no-vec)
- Normal plus "no-fma" (bin-no-fma)

Next, 7 runs are performed:



- 1. bin \Rightarrow reference run, only time and mem footprint is taken
- 2. $bin + Darshan \Rightarrow IO$ metrics
- 3. scalasca \Rightarrow Global, MPI, OMP, CPU counters
- 4. (bin-no-vec) \Rightarrow Core, vectorization efficiency
- 5. (bin-no-fma) \Rightarrow Core, FMA efficiency
- 6. bin compact run \Rightarrow mem vs comp. bound
- 7. bin scatter run \Rightarrow mem vs comp. bound

The generation of the binaries as well as the execution of all necessary runs has been automated by using the JUBE environment. Specific metrics as well as a full metric overview can be created with a single JUBE execution by extracting relevant information from the seven runs performed.

To proof the automation process, designed by the Metallwall code team, the Esias code team used the provided scripts and configuration techniques to automate their code in a similar manner on a different HPC system (JUQUEEN). The metrics provided by Scalasca, Darshan and the reference values could be easily included into the automated process and analyzed in a very short amount of time with the help of the Metallwalls configuration examples.

This procedure can now serve as a blueprint for other code teams and eventually of course also by the general public, via dissemination through WP 6. Within the project the relevant code examples were distributed via the Gitlab infrastructure. Table 3 and 4 in appendix shows the results of fully automated runs for Metalwalls and ESIAS.



4. Codes evaluated on the period Oct 2015 - March 2016

All codes mentioned in table 1 have established a close cooperation between HPCexperts and code developers following the above mentioned underlying lean management structure. They regularly update and report on their progress by means of the Code Diaries which are maintained on the Git structure along with code changes, automation processes and metrics.

Figure 2 shows the status of all codes regarding the implementation and analysation of the different profiling tools and of the benchmark automatisation process.

Code	WP	JUBE integration	Benchmarks defined in JUBE	Tools integrated in JUBE	Allinea report	Score-P profile	Score-P trace	Scalasca analysis	Vampir analysis	Extrae measurement	Paraver analysis	Darshan results	VTune analysis	Advisor analysis	Total Progress
ALYA	WP 2	1	1	0	1	1	1	1	1	1	1	0	1	1	11
ESIAS	WP 2	2	1	1	0	2	1	1	0	0	0	1	0	0	9
Metalwalls	WP 3	2	2	2	2	2	2	2	0	2	2	2	0	0	20
PVnegf	WP 3	1	0	0	0	0	0	0	0	0	0	0	0	0	1
SHEMAT	WP 4	2	2	1	2	1	1	1	0	2	2	2	0	0	16
ParFlow	WP 4	2	0	1	1	1	1	1	1	1	1	1	1	0	12
GYSELA	WP 5	2	2	2	1	1	1	1	0	2	2	2	0	0	16

EoCoE code benchmarking and analysis progress sheet - checkpoint April 4th 2016

Legend 0 not started 1 in progress

2 established

Figure 2: Code benchmarking and analysis progress sheet



A. Performance evaluation reports

A.1 Metalwalls

Code ID card

<u>Code name:</u> Metalwalls <u>Scientific domain:</u> WP3 Molecular dynamic Description:

Metalwalls is a classical molecular dynamics code that simulates energy storage devices: supercapacitors. These devices could replace in the future the batteries used in nowadays hybrid vehicles.

Languages: Fortran90 (20k lines) Library dependencies: MPI, OpenMP is in project. Programing models: MPI, OpenMP is in project. Platforms:

- PRACE Tier0 Mare Nostrum (20 MCPUh in 2016)
- French Tier1 Occigen (5 MCPUh in 2015)

Scalability results: It has been ported on X86 architectures, scaling results are good up to $\overline{1000}$ cores.

Typical production run: 24h on 64 - 512 cores Input / Output requirement:

- Size: 10 GB / 24 h run
- Single post-processing output: 50MB
- Single restart output: 50MB

Application references:

Merlet, C.; Rotenberg, B.; Madden, P. A.; Taberna, P.-L.; Simon, P.; Gogotsi, Y.; Salanne, M. Nature Materials. 2012, 11, 306–310 Contact:

- Mathieu Salanne (mathieu.salanne@upmc.fr)
- Matthieu Haefele (matthieu.haefele@maisondelasimulation.fr)

Metrics and performance report

<u>Code team</u>:

- Matthieu Haefele (MdlS) for WP1
- Mathieu Salanne (MdlS) for WP3

Case1 characteristics:

• Domain size: 3776 ions (walls + melt)



- Resources: 1 node on Jureca (24 cores)
- IO details: Checkpoint written every 10 steps instead of $1000 \Rightarrow$ much larger than production
- Type of run: both a development and small production run

	Metric name	03/01/2016			
	Test-case	case1			
1	Total Time (s)	43.2			
lba	Time IO (s)	0.3			
Go	Time MPI (s)	12.4			
	Memory vs Compute Bound	1.1			
	IO Volume (MB)	35.8			
0	Calls (nb)	384000			
Ī	Throughput (MB/s)	105.0			
	Individual IO Access (kB)	0.1			
	P2P Calls (nb)	0			
	P2P Calls (s)	0.0			
	Collective Calls (nb)	2721			
Ы	Collective Calls (s)	0.1			
Μ	Synchro / Wait MPI (s)	11.7			
	Ratio Synchro / Wait MPI	94.8			
	Message Size (kB)	908.4			
	Load Imbalance MPI	24.8			
le	Ratio OpenMP	0.0			
Noc	Load Imbalance OpenMP	0.0			
~	Ratio Synchro / Wait OpenMP	0.0			
n	Memory Footprint (B)	66 mB			
Чeı	Cache Usage Intensity	N.A.			
I	RAM Avg Throughput (GB/s)	N.A.			
	IPC	N.A.			
e	Runtime without vectorisation (s)	46.5			
Cor	Vectorisation efficiency	1.1			
	Runtime without FMA (s)	44.6			
	FMA efficiency	1.0			

Table 3: Performance metrics for Metalwalls on the JURECA HPC system

According to Table 3, Metalwalls does not seem to need support on IO as less than 1% of execution time is spent in IO on a case that produces much more data than a production run. However, the IO metrics show a very large number of calls compared to the amount data written on disk and this is typical for such ASCII based outputs. The implementation of binary based outputs would help here but it is not a priority.

The 30% time spent in MPI is mostly due to load imbalance. The root of this imbalance could be spot thanks to the analysis of the scalasca trace. It resides in the *cgwallrealE* subroutine. The uniform distribution of atom pairs leads here to a load imbalance because some pairs require more computations than others. The implementation of an ad hoc load balancing scheme that would distribute the load between the MPI processes rather than the pairs could solve the issue and let the code scale much better.

Table 3 shows a poor vectorization efficiency. The trace obtained with scalasca allowed us to identify the most intensive parts of the code. A careful examination of these code regions on top of a very good compute bound indicator of 1.1 gives the feeling that



the vectorization efficiency could be improved.

During this code investigation, we also noticed a discrepancy between the size of the data structures manipulated in the intensive regions and the global memory footprint measured on Table 3. This memory footprint is much larger than expected, some progress can certainly be made in this area.

Finally, the fact that Metalwalls is a pure MPI code can be a limitation on nowadays multi-core architectures and will definitely be one with the upcoming many-core architectures. An OpenMP implementation that could extract a fine grain parallelism could alleviate this limitation.

As a conclusion, in order to improve Metalwalls, we would recommend the following roadmap:

- 1. Single core optimizations would cure the memory footprint issue as well as the vectorization one.
- 2. An ad hoc load balancing scheme would allow the code to scale better in its pure MPI form.
- 3. An OpenMP implementation would prepare the code for the upcoming architectures.

A.2 Esias

Code ID card

<u>Code name:</u> ESIAS (Ensemble for Stochastic Integration of Atmopheric Simulations) <u>Scientific domain:</u> WP2: Meteo4Energy Description:

Coupled Ensemble implementation of Weather Research and Forecasting Model (WRF) and European Air Pollution and Dispersion Inverse Model (EURAD-IM) for short to medium range probabilistic forecasts and emission parameter estimation using Monte Carlo and Variational Data assimilation techniques. WRF is a state-of-the-art mesoscale numerical weather prediction system which is used extensively for research and operational real-time forecasting at numerous public research organizations and the private sector throughout the world and is open to the public. It offers various sophisticated physics and dynamics options. EURAD-IM is a fully adjoint chemistry transport model on the regional scale for chemical species and aerosols which is used for both, operational air quality forecasts and research applications. A main feature is the joint intital value and emission factor optimization using four dimensional variational data assimilation. Languages: Fortran90 and C (500k lines)

Library dependencies: MPI, OpenMP, NetCDF, zlib, libpng, JasPer Programing models: MPI, OpenMP Platforms:

• IBM Blue Gene/Q JUQUEEN

Scalability results: It has been ported on X86 architectures, scaling results are good up to 524288 cores (512 each ensemble member).

Typical production run: 2h on 16384 - 32768 cores



Input / Output requirement:

- Size: 1 TB / 24h run (1000 ensemble members, 1 GB each)
- Single post-processing output: 10 GB (1000 ensemble members, 1 GB each)
- Single restart output: 100 TB (1000 ensemble members, 1 GB each)

Relevant kernel algorithms: Particle Filtering, 4DVAR, Quasi-Newton Minimization (LBFGS), FFT Software licence: None Application references:

W. C. Skamarock, J. B. Klemp, J. Dudhia et al., "A Description of the Advanced Research WRF Version 3". NCAR Technical Note, NCAR, Boulder, Colo, USA, 2008. <u>Contact:</u>

- Hendrik Elbern (h.elbern@fz-juelich.de)
- Jonas Berndt (j.berndt@fz-juelich.de)

Metrics and performance report

<u>Code team</u>:

- Sebastian Lührs (FZJ) for WP1
- Jonas Berndt (FZJ) for WP2

Case characteristics:

The benchmark setup contains a random simulation period of 6 hours with 240x240x24 gridpoints as a typical size. For benchmarking, solely 2 ensemble members run in parallel (instead of the order 1000 for production runs, would be too computational intensive for benchmarking). No particle filtering is performed due to the small ensemble size. 1024 Processors are used. Parallel NetCDF is used. The metrics results by using the Darshan and the Scalasca instrumentation are given in Table 4.

I/O and metadata handling can be a bottleneck when using larger numbers of ensemble members. This will be tested in additional benchmarks by using a higher number of ensemble members. Also the usage of the NetCDF4 instead of the pNetCDF library will be tested.

The single core performance can still be improved by using a higher compiler optimization level but options create stability problems, or will change the final result and has to be checked. Especially vectorization wasn't successfully tested so far.

OpenMP can be used in WRF underneath the Esias ensemble creation, but currently the feature isn't used. The performance benefit towards a full MPI parallelization will be tested.



	Metric name	out.json
	Total Time (s)	259.7
lba	Time IO (s)	27.2
B	Time MPI (s)	178.5
	Memory vs Compute Bound	N.A.
	IO Volume (MB)	3570.9
0	Calls (nb)	63594
Ī	Throughput (MB/s)	131.3
	Individual IO Access (kB)	118.4
	P2P Calls (nb)	135267
	P2P Calls (s)	8.1
	Collective Calls (nb)	6170
Ы	Collective Calls (s)	1.1
Z	Synchro / Wait MPI (s)	98.4
	Ratio Synchro / Wait MPI	55.1
	Message Size (kB)	16.0
	Load Imbalance MPI	38.3
le	Ratio OpenMP	N.A.
Noc	Load Imbalance OpenMP	N.A.
	Ratio Synchro / Wait OpenMP	N.A.
п	Memory Footprint (B)	N.A.
Mer	Cache Usage Intensity	N.A.
	RAM Avg Throughput (GB/s)	N.A.
	IPC	N.A.
e	Runtime without vectorisation (s)	N.A.
Cor	Vectorisation efficiency	N.A.
	Runtime without FMA (s)	N.A.
	FMA efficiency	N.A.

Table 4: Performance metrics for Esias on the JUQUEEN HPC system