

UL HPC School 2014

PS7: Bioinformatics software on the UL HPC Platform

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1 / 17

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Latest versions available on Github:

UL HPC tutorials:

https://github.com/ULHPC/tutorials

UL HPC School:

http://hpc.uni.lu/hpc-school/

PS7tutorial sources:

https://github.com/ULHPC/tutorials/tree/devel/advanced/Bioinformatics



2 / 17

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Summary

- Objectives
- 2 Bioinformatics packages
- **3** Notes
- 4 Practical session
- **5** Conclusion







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Better understand the usage of Bioinformatics packages on the UL HPC Platform.



5 / 17

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Why Bioinformatics? 3Vs:



5 / 17





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Why Bioinformatics? 3Vs:

very relevant in the context of the UL/LCSB



5 / 17

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Why Bioinformatics? 3Vs:

- very relevant in the context of the UL/LCSB
- very fast growing domain



5 / 17

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Better understand the usage of Bioinformatics packages on the UL HPC Platform.

- very relevant in the context of the UL/LCSB
- very fast growing domain
- very many associated workflows, thus excellent examples





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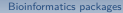














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ABySS

ABySS: Assembly By Short Sequences a de novo, parallel, paired-end sequence assembler designed for short reads



7 / 17

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Bioinformatics packages



ABySS

ABySS: Assembly By Short Sequences a de novo, parallel, paired-end sequence assembler designed for short reads

- several applications in the ABySS package
- only ABYSS-P is parallelized using MPI
- workflow (pipeline) of abyss-pe also includes:
 - → OpenMP-parallel applications
 - \hookrightarrow serial applications
- Note: compared with other de novo assemblers, the per-node memory requirements are smaller due to ABySS' task distribution model





Gromacs

GROMACS: GROningen MAchine for Chemical Simulations versatile package for molecular dynamics, primarily designed for biochemical molecules



8 / 17

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- very large codebase: 1.836.917 SLOC
- many applications in the package, several parallelization modes
- mdrun: computational chemistry engine, performing:
 - \hookrightarrow molecular dynamics simulations
 - ⇔ Brownian Dynamics, Langevin Dynamics

 - $\hookrightarrow L\text{-BFGS}$

 - → Normal Mode Analysis
- mdrun parallelized using MPI, OpenMP, pthreads and with support for GPU acceleration





Bowtie2/TopHat

Bowtie2: Fast and sensitive read alignment

ultrafast & memory-efficient alignment of sequencing reads to long ref. sequences

TopHat: A fast spliced read mapper for RNA-Seq

alignment of RNA-Seq reads to a genome, to identify exon-exon splice junctions





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- TopHat aligns reads to mammalian-sized genomes using Bowtie
- then analyzes the mapping results to identify splice junctions between exons
- bowtie2 is OpenMP-parallel
- rest of workflow is sequential





mpiBLAST

mpiBLAST: Open-Source Parallel BLAST

parallel implementation of NCBI BLAST, scaling to hundreds of processors



10 / 17

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- two main applications: mpiblast mpiformatdb
- requires (NCBI) substitution matrices and formatted BLAST databases
- the databases can be segmented

 - → or a multiple, in order to avoid load imbalance
- mpiblast requires >= 3 processes, 2 used for internal tasks
 - \hookrightarrow mpirun -np 3 mpiblast [...] only gives you one searcher process!







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Notes..

- .. on real world applications (bioinfo or others):
 - make sure you understand the parallel capabilities of your software

 ⇒ pthreads/OpenMP vs MPI vs hybrid
 - \hookrightarrow use of GPU acceleration



12 / 17





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- .. on real world applications (bioinfo or others):
 - make sure you understand the parallel capabilities of your software \hookrightarrow pthreads/OpenMP vs MPI vs hybrid
 - \hookrightarrow use of GPU acceleration
 - make sure you request the appropriate resources for the processing needs of your workflow
 - → Does the software always take advantage of more than 1 core or node?
 - → How does it scale? Many obstacles to perfect scalability!



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- .. on real world applications (bioinfo or others):
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 → pthreads/OpenMP vs MPI vs hybrid
 - \hookrightarrow use of GPU acceleration
 - make sure you request the appropriate resources for the processing needs of your workflow
 - \hookrightarrow Does the software always take advantage of more than 1 core or node?
 - → How does it scale? Many obstacles to perfect scalability!
- .. on data management:
 - make sure you use the appropriate storage place
 - \hookrightarrow \$HOME vs \$WORK vs \$SCRATCH
 - stage data in/out, archive your (many & unused) 'small' files





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Exercises

- Read and understand the Bioinformatics tutorial
 - https://github.com/ULHPC/tutorials/tree/devel/advanced/Bioinformatics
- Run the examples
 - \hookrightarrow all calculations should be fast
 - \hookrightarrow you should attempt the exercises proposed in each section
- Try even more tests, e.g.:
 - → on different node classes
 - \hookrightarrow with one core per node on >= 2 nodes
 - \hookrightarrow vs >= 2 cores on single node







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Conclusion

- Bioinformatics applications execution on the UL HPC Platform
- Outlined:
 - → different workflows
 - $\,\hookrightarrow\,$ some of the concepts you should care about when running complex software

Perspectives

• Personalize the UL HPC launchers with the specific commands for ABySS, Gromacs, TopHat, Bowtie, mpiBLAST...





Questions?



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