

Introduction to Parallel COMPUTING

Luca Tornatore, I.N.A.F.

2025 INAF Course on HPC



September, 22nd - 26th, OACT, Catania



“CRUCIAL PROBLEMS that we can only hope to address computationally REQUIRE US TO DELIVER **EFFECTIVE COMPUTING POWER ORDERS-OF-MAGNITUDE GREATER THAN WE CAN DEPLOY TODAY.**”

DOE's Office of Science, 2012

..even more valid nowadays, though

Outline



Intro to Parallel
Computing



Parallel
Performance

Outline



Intro to Parallel Computing

- Some general definition and introduction to HPC's rationale
- Why parallelism ?
- How do you do parallelism ?
- a definition of parallel computing
- shared- vs distributed- memory paradigm
- HPC components:
 - the hardware, from cores to clusters: where the parallelism actually lives
 - the memory



Some general definition and introduction to HPC's rationale

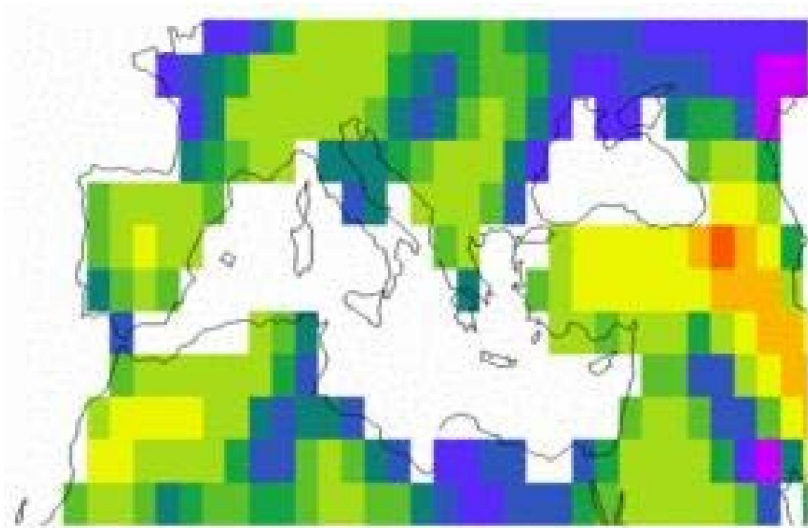
What is High Performance Computing

High performance computing (HPC), also known as supercomputing, refers to computing systems with extremely high computational power that are able to solve hugely complex and demanding problems.

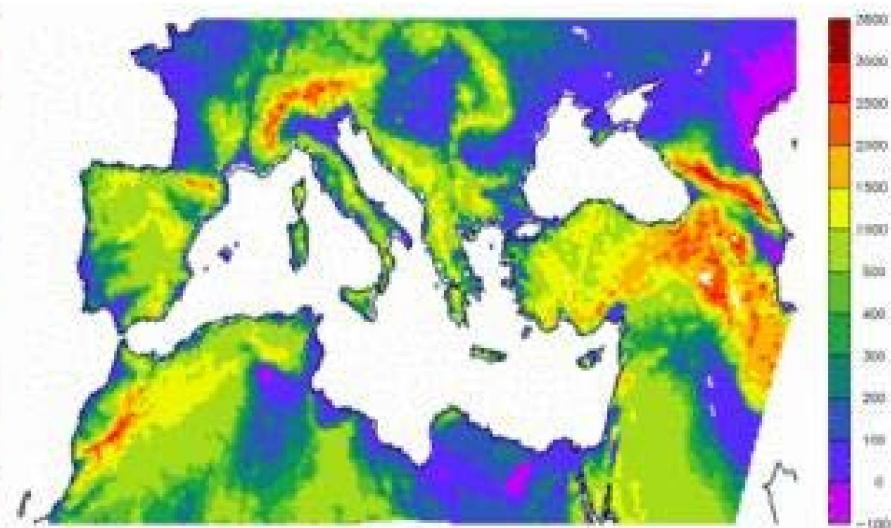
Taken from <https://ec.europa.eu/digital-single-market/en/high-performance-computing>

Why High Performance Computing ?

Example of a complex problem: climate change over the Mediterranean Sea.
How the required memory increases if the resolution passes from 200 km (~1GB) to 2km ?



200 km



25 km

Why High Performance Computing ?

Complex problems solved by simulations

- Simulation has become the way to research and develop new scientific and engineering solutions.
- Used nowadays in leading science domains like aerospace industry, astrophysics, material science, etc.
- Challenges related to the complexity, scalability and data production of the simulators arise.
- Impact on the relaying IT infrastructure.



Why High Performance Computing ?

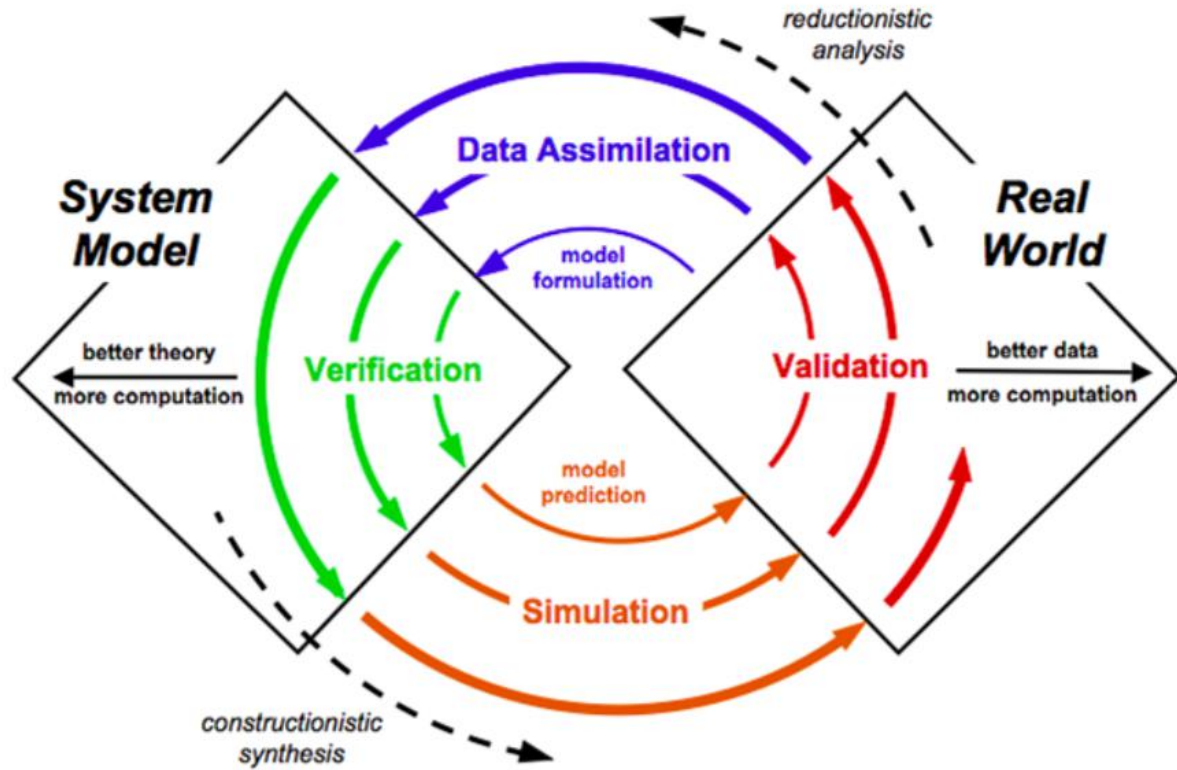
Complex problems solved by simulations

- Simulation has become the way to research and develop new scientific and engineering solutions.
- Used nowadays in leading science domains like aerospace industry, astrophysics, material science, etc.
- Challenges related to the complexity, scalability and data production of the simulators arise.
- Impact on the relaying IT infrastructure.

HPC capabilities are used to solve and address scientific, industrial and societal challenges.

<p>Health</p> <ul style="list-style-type: none">• Development of personalised and precision medicine to provide individual and accurate patient treatment.• Saving time and money on the development of new drugs from the initial idea phase to the final phase of reaching the market. <p>development of new drugs reduced from 17 years to several months</p><p>HPC can help save hundreds of millions up to billions of Euros</p>• Early detection of diseases and quicker diagnosis.	<p>Climate change & weather forecast</p> <ul style="list-style-type: none">• Europe paid severe weather damage costs between 1970 and 2012¹.With HPC technology, climate scientists will be able to predict the size and paths of storms and floods with accuracy.This will allow implementation of measures such as alerting or evacuating people, thus saving human lives. <p>150.000 lives  €270 billion in economic damages</p>	<p>Industry</p> <ul style="list-style-type: none">• Reducing development time, minimising costs, optimising decision processes and producing higher quality goods and services.E.g. automotive industry will save time and money to develop new vehicle platforms, with improved environmental friendliness and passenger comfort and safety². <p>development reduced from 60 months to 24 months  HPC can help save up to € 40 billions</p>
<p>Cybersecurity</p> <ul style="list-style-type: none">• Enabling complex encryption technologies and better reactions to cyberattacks.• Combined with Artificial Intelligence, HPC detects:<ul style="list-style-type: none">◦ strange systems behaviour,◦ insider threats and electronic fraud,◦ very early cyberattack patterns (few hours instead of a few days)This will allow automated and immediate actions even before a cyberattack happens.	<p>Energy</p> <ul style="list-style-type: none">• HPC provides critical tools for example in:<ul style="list-style-type: none">◦ designing renewable energy parks◦ designing high-performance photovoltaic materials,◦ optimising turbines for electricity production.• HPC expenditure in the energy sector is projected to grow by 5% in the next years.	

Why High Performance Computing ?



The inference spiral

As models become more complex and increasingly sophisticated experiment experiments and new data bring in more information, an ever increasing computational power is required.



Why High Performance Computing ?

A flood of data

In today's world, larger and larger amounts of data are constantly being generated, from ~30 zettabytes globally in 2018 to an expected ~200 zettabytes in 2025).

As a result, the nature of computing is changing, with an increasing number of data-intensive critical applications.

Is key to processing and analysing this growing volume of data, and to making the most of it for the benefit of citizens, businesses, researchers and public administrations.

Taken from <https://ec.europa.eu/digital-single-market/en/high-performance-computing>

Name	Symbol	Factor
Yotta	Y	1024
Zetta	Z	1021
Exa	E	1018
Peta	P	1015
Tera	T	1012
Giga	G	109
Mega	M	106
Kilo	K	103

... High Performance Computing ?

We have seen some argument in favour of HPC as a fundamental tool for the scientific and technological research.

However, how it happens in practice ?

Can you do HPC on your laptop ?

If not, what do you need ?



- Why parallelism
- How do you do parallelism ?



Why parallelism ?

For two main reasons:

1. Time-to-solution

- To solve the same problem in a smaller time
- To solve a larger problem in the same time

2. Problem size (~ data size)

To solve a problem that could not fit in the memory addressable by a single computational units (or that could fit in the space around a single computational units without serious performance loss)



Two types of problems

Let's suppose that you have a computational problem.
There are two basic, and distinct, cases:

1. The problem's solution for each data point is completely independent of the solution for any other data point (these are called ***embarrassingly parallel problems***)
2. All the rest (the opposite than the previous case)

Of course, normally you have some blending of computations that are independent of each other and of computations that are not, with a significant imbalance towards the second case.

Truly embarrassingly parallel problems are a tiny fraction, in real world.



Two types of problems

Examples:

- 1) reducing the spectra of a set of photon sources is normally an embarrassingly parallel problem, since the operations on a spectrum are truly independent of the operations on another spectrum
- 2) if you're doing interference, as in radio astronomy, that is no more true because you have to treat the sources together, at least for some operations.



Two types of problems

Examples:

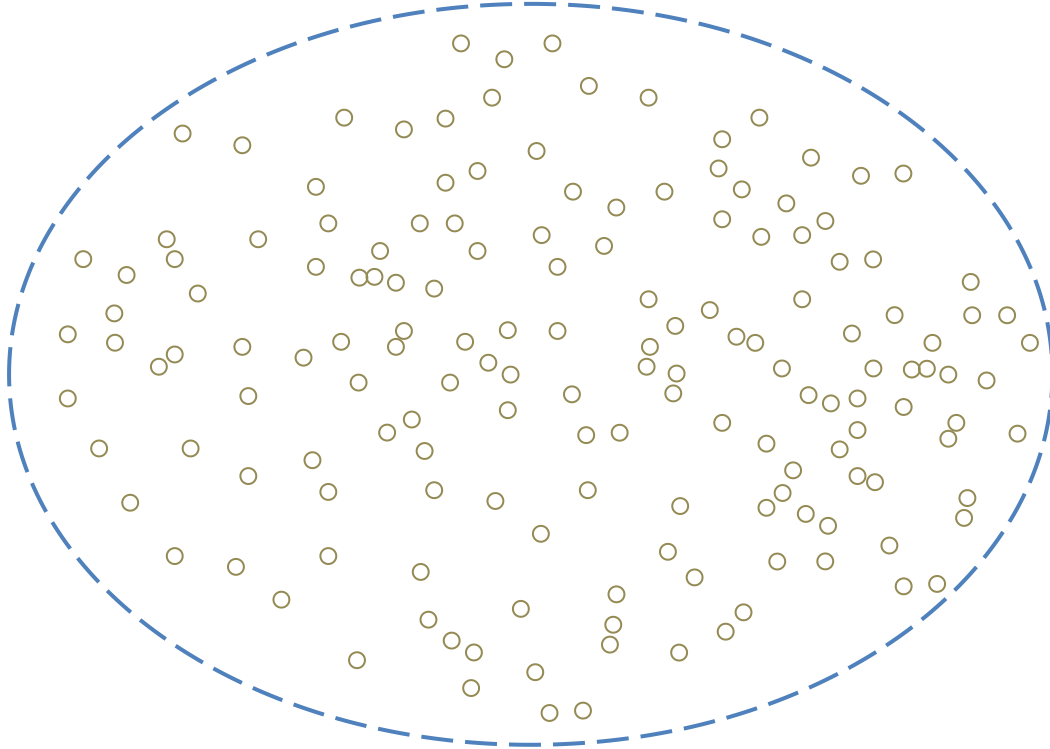
In a simulation,

- 1) estimating local properties (of a single cell or particle) may be a truly embarrassingly parallel problem - for instance: once all the physical quantities are at hands, estimating the cooling function is a task that can be performed for each element independently.

- 2) estimating the dynamics (gravitational force and acceleration) of a mass element is *not* embarrassingly parallel because you need to evolve all the mass elements together.



Embarassingly parallelism

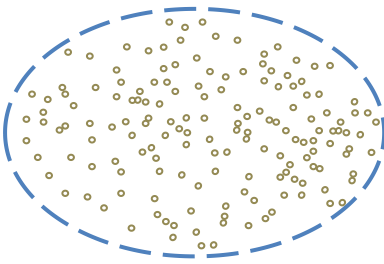


Your data set is made of data points for each of which the problem's solution is independent of other points (for instance: a series of spectra to be reduced and analysed).

Your task is to solve the computational problem for each and all of them.



Embarassing parallelism



Note that it is irrelevant that the solution for every data point amounts to run exactly the same instructions, as in:

```
reduce spectrum;
```

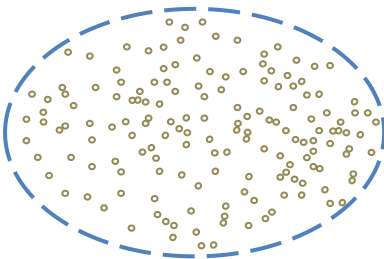
or not, as in:

```
if (is_a_spectrum) reduce spectrum;  
else if (is_a_match) play quidditch.exe;  
else if (is_Darth_Vader) play Jedi.exe;  
...
```

The point of embarrassinglyness is the solutions to be completely independent of every other solution, for each data point.



Embarassing parallelism



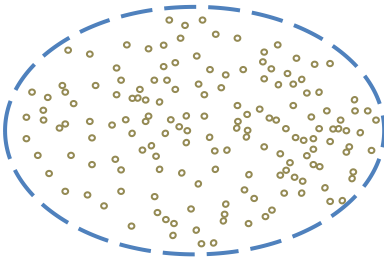
For the sake of simplicity, suppose that the solution is about running the same code for every data point.

If a single processing unit (p.u.) was applied to the problem's solution, performing the task T for each data point subsequently, you would have a **serial** solution that would take $N \times \Delta t$ run-time, where N is the size of your data set and Δt is the time needed for a single data-point (which here we suppose to be constant)





Embarassing parallelism

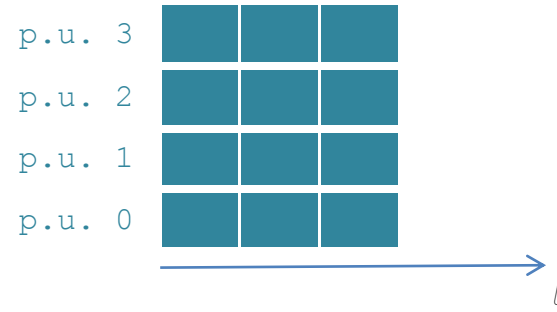


Applying more than one p.u., each p.u. would solve the problem for a data point, then to another one and so on.

If you can evenly distribute the data points among the p.u., the run-time scales obviously as $(N \times \Delta t)/\eta$ where η is the number of p.u.

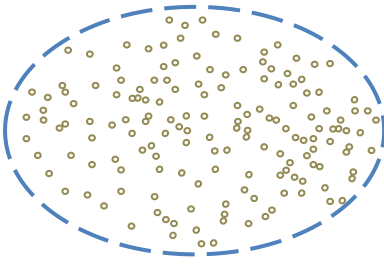
That would be a perfect scaling

NOTE: this naïve estimate does not account for data dispatching, data accessing and so on. Normally, these are significant factors.





Embarassing parallelism



A sad note:

In these cases, what people normally do is, roughly, to buy what they call “a server” with expensive many-cores CPUs and a gargantuesque amount of RAM, and then to launch as many separate instances of their code as data points.

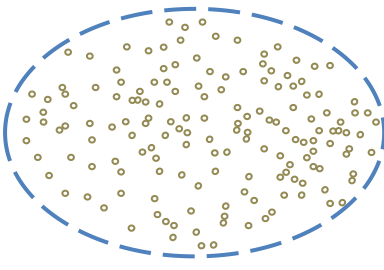
That is *not* parallelism, that is **concurrency**.

In best cases, data sets are sub-divided in bunches and there is some scheduling in the code executions.

In most of the cases (for instance: the same machine serves many users that have the same kind of approach) there is not such thing and the performance (i.e. the run-time) severely degrades.

More over, what happens when your data set grows spectacularly?

What happens if the computational task for each data point has not a negligible run-time and/or requires a non-negligible amount of memory?



Embarassing parallelism

A sad note:

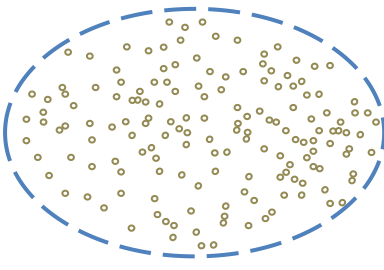
In these cases, what people normally do is, roughly, to buy what they call “a server” with expensive many-cores CPUs and a gargantuesque amount of RAM, and then to launch as many separate instances of their code as data points.

That is *not* parallelism, that is **concurrency**.

You have **parallelism** when the code execution runs on more than one p.u. tackling the entire data set (internally managing the memory limits), and making the p.u. to **cooperatively** solve the problem: for embarrassingly parallel problems, the cooperation may simple consists in the agreement for the domain decomposition and the final synchronization.



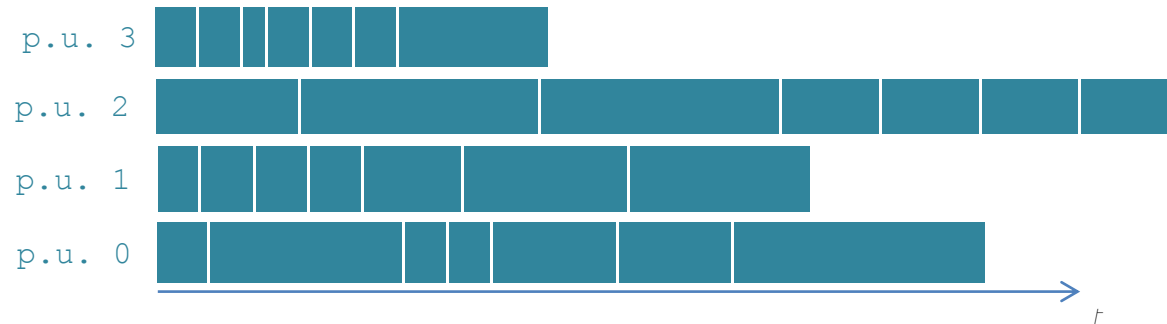
The slowest rules 'em all



Now let's abandon the assumption that the run-time for each data point is constant. That may be due to several factors: for instance, the execution being not exactly the same for each data point, or the data-size of each data point being not constant and so on.

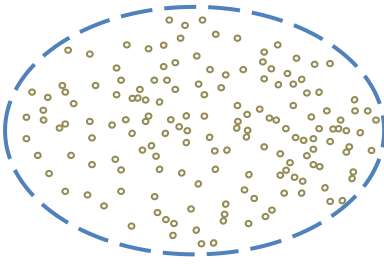
Then, **suddenly how you spread your data points among the p.u. may strongly affect your run-time**, because the following general rule holds

you will never be faster than the slowest of your players





Work imbalance

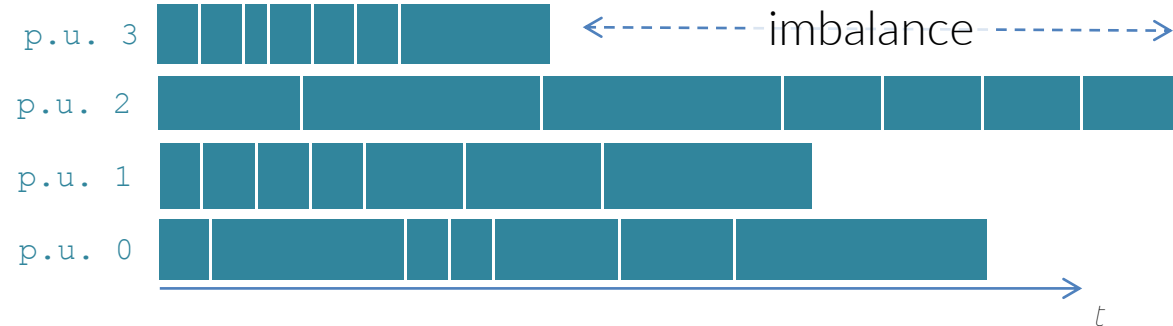


The work-imbalance is a fundamental metrics to characterize your code, or the different sections of it.

tips: always instrument your codes to track the imbalance.

you will never be faster than the slowest of your players

From that it descends that ideally the slowest must progress at the same pace than the fastest, i.e. you do not have any **imbalance**.



The **work-imbalance** is the measure of how different the computational load is among your players (it is zero when the work is perfectly spread - which is achieved virtually in no cases).

There are various possible definition of work-imbalance (the max difference in timing, the std.dev. of timings, ...); once you've got the point, you may apply/find that which fits at best your needs.



Domain decomposition

A severe imbalance may be due to several factors.

How the data are distributed among p.u. may be sub-optimal, which is quite common when the computational work for every data point is strongly dependent on the data properties and is not easily predictable.

How you decide which data point are processed by each p.u. is called **domain decomposition** and depends on the nature of the data and their availability (are they already there, are they arriving in bunches or singularly, ...).

The domain decomposition is a vey big and focal chapter in parallel computation.

There is not a general optimal way to perform the domain decomposition: that is strongly dependent on the nature of both the computational problem and of the data.

Quite often, there are many different algorithms in a same problem, and the optimal decomposition may be different among them.

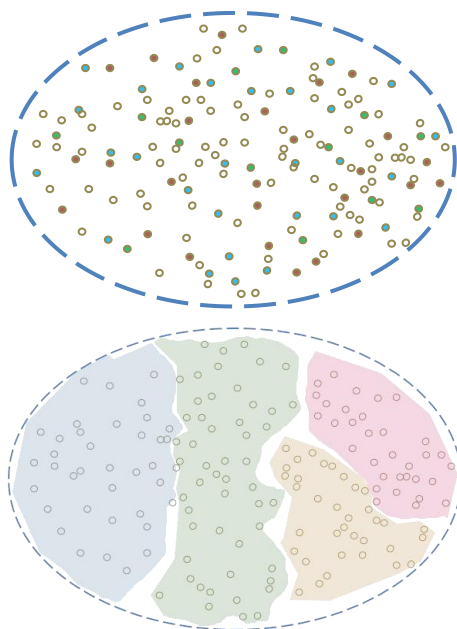
Take into account that almost always, the domain decomposition is itself an expensive task that often requires computation, sorting, searching and lot of data moving.



Domain decomposition

When you are reasonably certain that the work load is reasonably constant for every point, just go for the simplest decomposition possible to minimize the overhead due to the decomposition itself.

Trivial random decomposition may serve quite well in these cases.



for instance: random or first-arrived first-served basis

or very simple “geometrical” decomposition (where the “geometry” depends on the dimensional space of the data)

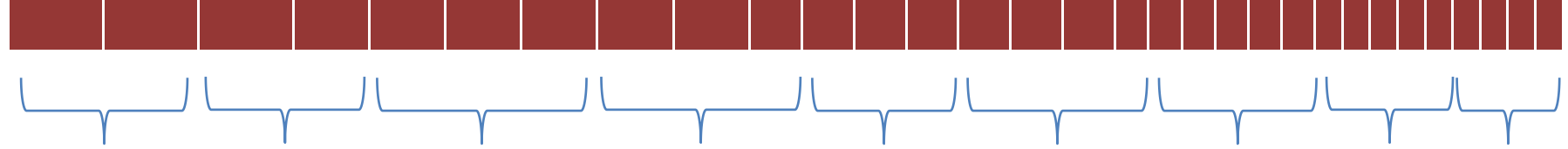


Domain decomposition



When the work-load is strongly dependent on the properties of the data, and moreover those properties change in time during the computation, there is not a general rule.

You may, for instance, sort your data by computational intensity and distribute them to different players so to achieve an even work-load with a minimum imbalance; then, repeating the procedure as often as needed while the data evolves during a multi-step computation.



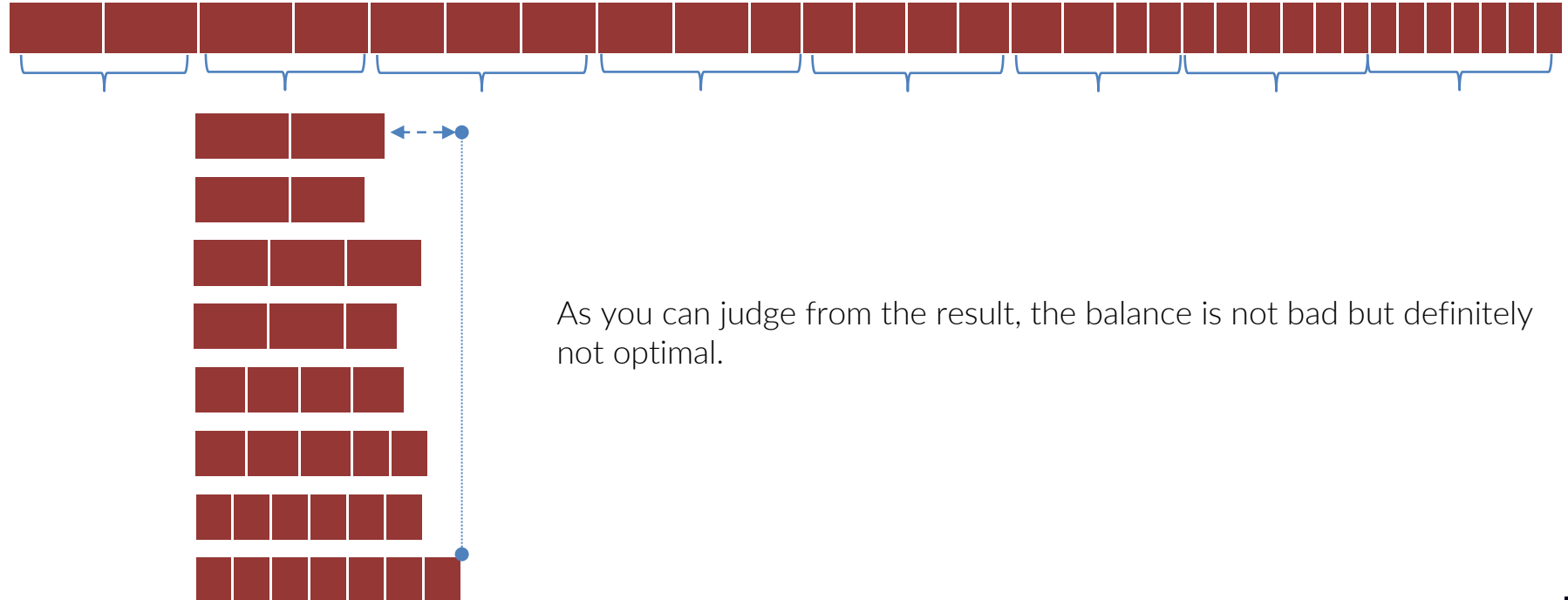


Domain decomposition



example 1:

after sorting the data by computational load (the cells' size in the figure below is supposed to be proportional to the computational cost, you pick up chunks of as even as possible size



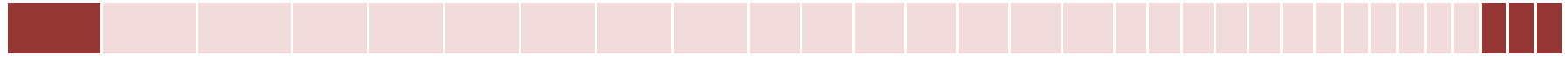


Domain decomposition



example 2:

A slightly different approach could be to have a round-robin assignment, alternatively from the most-expensive and the least expensive queue

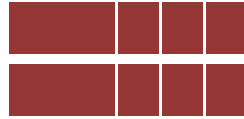
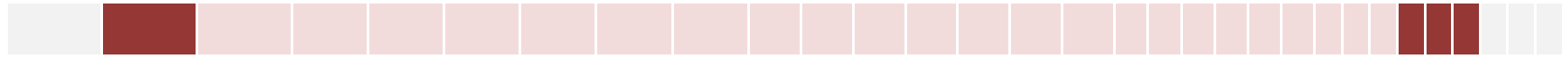




Domain decomposition

example 2:

A slightly different approach could be to have a round-robin assignment, alternatively from the most-expensive and the least expensive queue

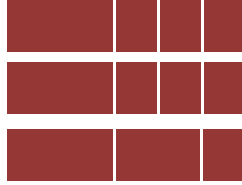
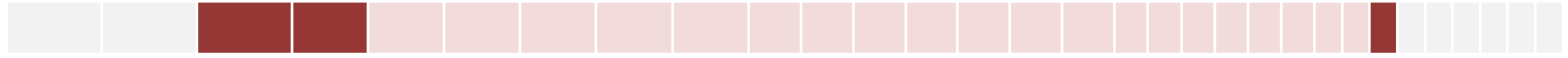




Domain decomposition

example 2:

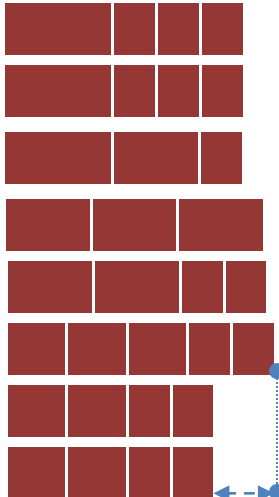
A slightly different approach could be to have a round-robin assignment, alternatively from the most-expensive and the least expensive queue



Domain decomposition

example 2:

A slightly different approach could be to have a round-robin assignment, alternatively from the most-expensive and the least expensive queue



The resulting distribution is slightly better.

An even better result may be achieved by iteratively adjusting the domain decomposition moving the sub-domains at the frontiers of the chunks to minimize the total imbalance (i.e., for instance, the difference between the most and the least computational heavy chunk).

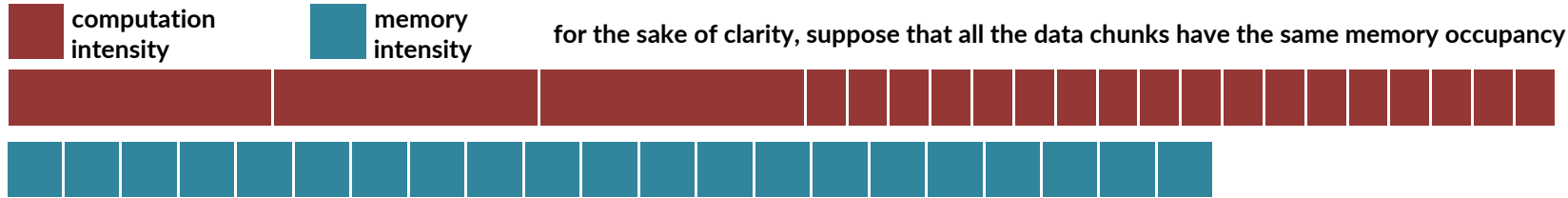
A third option is to define a multiple-chunks domain decomposition.

Still, these were just example to give you an idea of how complex a domain decomposition could be.



Domain decomposition

Achieving a best-possible work-balance is a fundamental goal. Normally, for this purpose you have to accept a **memory-inbalance** cost, which basically amounts to assign a larger number of data (and hence of memory) of lower computational intensity to some players and a smaller number of computationally-intensive data to some other.



work-load distribution that achieve the
best-possible imbalance for 2 players



corresponding memory-load distribution

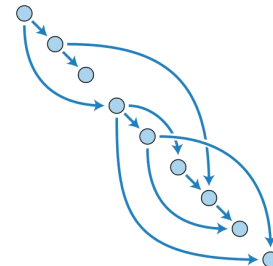




Functional decomposition

Quite often, your computation is made of several different “sections”. Let’s call them **tasks**. There will be, in general, a dependency graph among the tasks: some will be completely independent of any other, some will be dependent on 1, 2 or more “previous” tasks and will feed some “subsequent” tasks.

Then, you may decompose not the data but the tasks among your workers. In general, that requires some **synchronization** to manage the dependencies.



Dependency graph among task;
this is an **acyclic graph**, i.e. there
are no loops.



Example of task decomposition with as
fewer dependencies as possible;
all the task are executed at the same time
with the least possible unbalance.



Functional decomposition



Almost always, an optimal choice is performing both type of decomposition.

At least to fit in the memory constraints, data are distributed among tasks so that to find the best trade-off among the desired work-load and the constraints of memory-load.

Some phases of the computation require a distributed parallelism, i.e. the computation on the data requires a communication that involves all the data

player 1



player 2



Once all the communication among players has been done, every player perform the computation on its data, possibly by distributing tasks among sub-players.

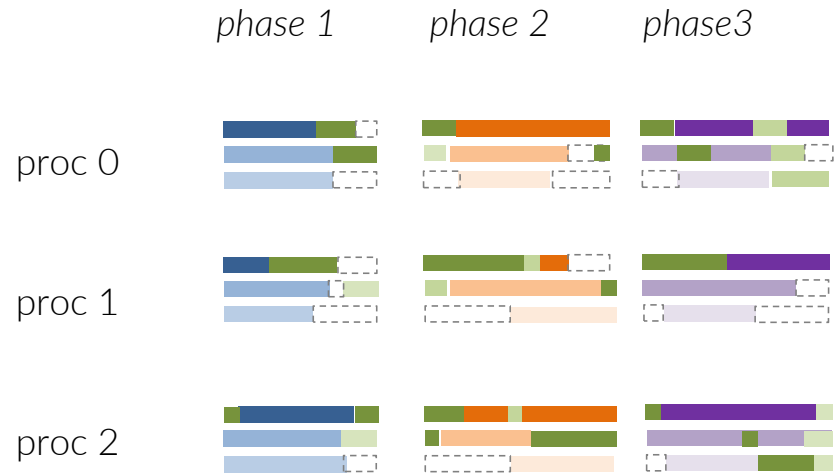
You may think to sub-players as the threads and to players as MPI tasks (that is called *hybrid parallelism*)



True (non-embarassing) parallelism

Almost always, a significant fraction of your problem is non-embarassingly parallel. As you know, that means that the computation needed to solve your problem can not proceed on every data point independently of any other data point (but, of course, for an initial set-up).

Normally you have something that looks like the following:



Let's suppose that the code has 3 main algorithms (the "phases", by color) and that each player has 3 sub-players (the different color intensity) that runs different tasks on the data.

Dark green indicates inter-player communication and pale green indicates inter-task data exchange or synchronization.

White dotted segment indicates idle spinning of a task.



How do you do parallelism ?

A good reference is the Part 1 of “Designing and building parallel programs” by Ian Foster, [publicly available](#) (pdf is easily findable).

It is outdated in some respects, but the concepts about parallelism, parallel design and parallel performance are explained in a clear and clean way.

While the **whole Part 1 is a useful reading**, I also suggest

Chapter 2, 0 → 2.5

Chapter 3, 0 → 3.8

Note that also the case studies in Chap 2 and 3 are interesting, of course.
Chap 4 is a useful re-collection of concepts, applied to case studies.



- a definition of parallel computing
- shared- vs distributed paradigm

What is parallel computing ?

1. A **parallel computer** is a computational system that offers *simultaneous access to many computational units* fed by memory units.
The computational units are required to be able to *co-operate* in some way, meaning *exchanging data and instructions* with the other computational units.

2. **Parallel processing** is the ensemble of techniques and algorithms that makes you able to actually use a parallel computer to successfully and efficiently solve a problem.

What is parallel computing

The parallel processing is expressed by **software entities** that have an increasing level of granularity:

processes, threads, routines, loops, instructions..

The software entities run on underlying **computational hardware entities** as processors, cores, accelerators

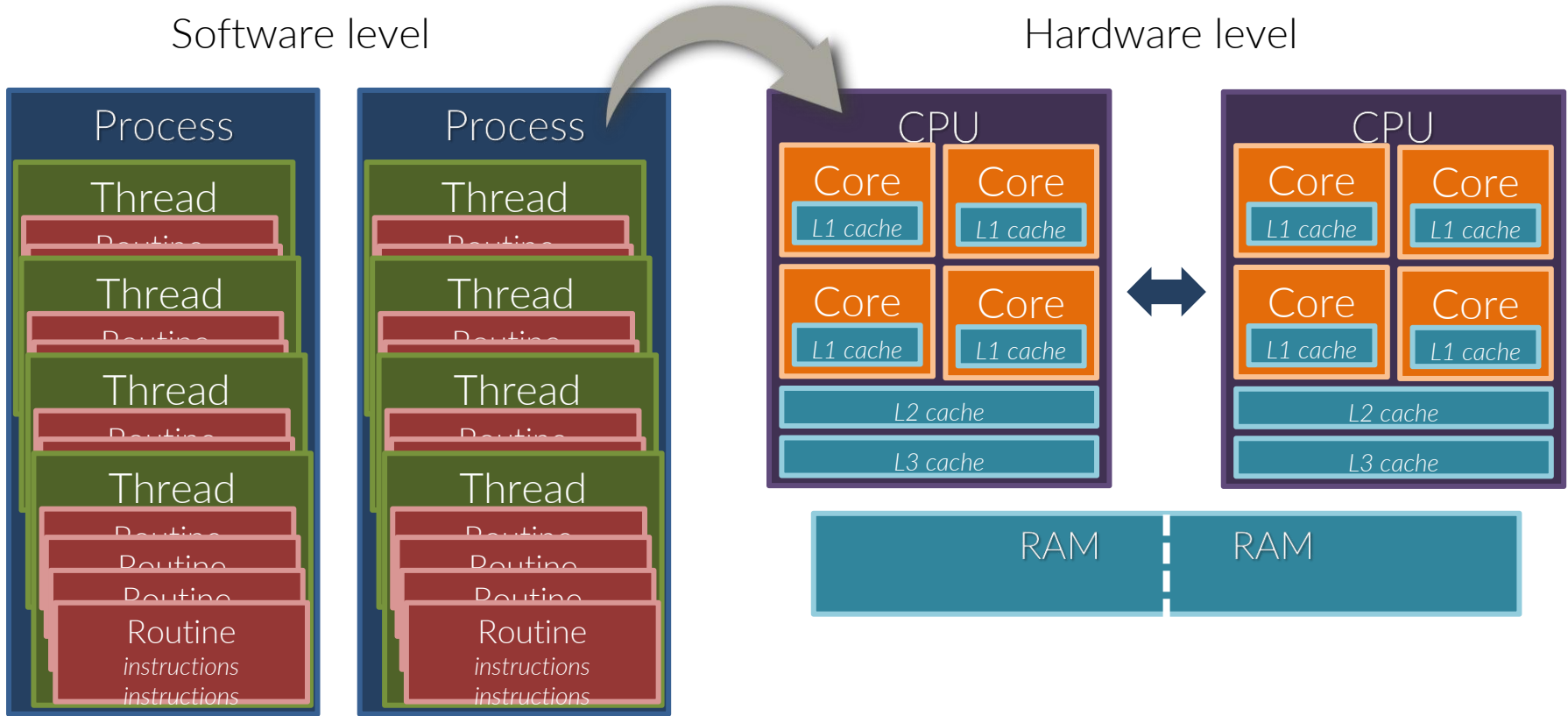
The data to be processed/created live and travel in **storage hardware entities** as

Memory, caches, NVM, networks, DMA

The *exploitation/access* of hardware resources (computational and storage) is **concurrent** among software entities



What is parallel computing





Shared vs Distributed Memory

We have discussed before the shared- and distributed- memory concepts. As we have seen, they refer to the physical placement of memory.

However, they may also refer to two distinct **programming paradigms**, named after them, i.e. exactly the shared-memory and distributed-memory paradigm.

While the two terms are logically derived from the *physical* access to the memory, as programming paradigms they refer to how the players exchange data; in other words, in shared-memory no active collaboration is strictly needed because there is some “common area” that can be accessed in writing and reading by all the players. In distributed memory, at odds, there is no such area: the players must collaborate to ask and provide data to each other.



Shared vs Distributed Memory

DISTRIBUTED MEMORY

A typical programming paradigm is the message-passing, i.e. the processes communicate via “messages” while each process has its own memory space. The actual communication may happen either over the network (different protocols are possible at hardware/middleware level) or via shared memory techniques if the communicating processes can directly access the same memory (the user still does not see it as shared memory but uses messages in its coding; the actual communication is managed by the middle-ware).

A very diffuse standard is **MPI**, **M**essage-**P**assing **I**nterface.

A message-passing standard may expose (as MPI does since version 2.0) interfaces for direct memory access, mimicking shared-memory mechanisms.



Shared vs Distributed Memory

SHARED MEMORY

A typical programming paradigm is multi-threading, i.e. multiple threads access concurrently the same virtual address space.

No such a thing as “messages” exists, communications and synchronizations must be managed in shared-memory.

A very diffuse high-level standard is **OpenMP (openmp.org)**, **Open Multi-Processing**. On all platforms, a very low-level library for threading is present. On POSIX system it is named *pthread*.

On some systems, a software middleware may hide the actual physical details from you and expose the entire machine’s memory of all nodes as a shared memory, even if underneath the remote memory access happens via network.
That is called PGAS, partitioned global address space.



Shared vs Distributed Memory

The two programming paradigms, shared- and distributed-memory, can easily be fused and used together.

That typically happens as the following.

An MPI process running on a node spawns a number of threads that by definition run on shared-memory among themselves. The same MPI process will communicate with other MPI processes via messages, irrespectively of where they run.

Actually, MPI v2.0 and v3.0 offer handy mechanisms to have RDMA (Remote Direct Memory Access) and even shared-memory access among MPI processes.



Shared vs Distributed Memory

If you wonder how you should proceed - opting for either an MPI code or an OpenMP code - here are a couple of practical advice.

Opting for a distributed memory paradigm (i.e. MPI) since the beginning has the big advantage that your code will be able to run on whatever number of nodes that will be needed to host your larger and larger data sets.

Of course, that will cost you some effort at the begin.

As a second step, you can add the OpenMP-ization.

For the cases in which you need a quick impact on your code's performance for runs that fit on single-node, then OpenMP is a good move. Multi-threading for many problems offers an easier parallel point of view and if properly implemented it may deliver very good performance boosts.



MPI & OpenMP

The two (by far) most used implementations of distributed memory and shared memory parallel paradigms are MPI (distributed memory) and OpenMP (shared-memory) that have already been mentioned before.

OpenMP is delivered by the C/C++/FORTRAN compilers themselves by asking them to switch on the OpenMP support; that happens by a command-line option, read the manual of your compiler to discover it (gcc and icc have -fopenmp, pgi and nvc have -mp, ...).

MPI is delivered by many different implementation: one of the most diffuse is OpenMPI (note the possible confusion in names: OpenMPI has nothing to do with OpenMP), others are MPICH, MVAPICH, ...



MPI & OpenMP

standard specifications

implementations

usage

OpenMP

www.openmp.org

the C/C++/FORTRAN compilers implement themselves the standard up to some version (check the compiler manual; usually there is complete support up to v4.5).

You switch on the OpenMP with a command-line option: `gcc` and `icc` use `-fopenmp`, `pgcc` and `nvc` use `-mp`, and so on.

You compile your code as

`cc -fopenmp -o executable my_code.c`

and you get an executable that is linked against the O.S. threading library.

You then execute it as every standard executable handed out by the compiler.

MPI

www.mpi-forum.org

There are several implementation of the MPI, which is basically a compiler wrapper and a library that interacts with lower-level library that manage the network.

Among the major implementations: OpenMPI, MPICH, MVAPICH, IntelMPI

You compile the code as

`mpicc -o executable my_code.c`

and afterwards you execute it by

`mpirun -np`

`number_of_MPI_processes`
`executable`



What is parallel computing ?

The Flynn's taxonomy (~60s)

Instructions		
single	multiple	
Data	multiple	SISD
	single	MISD
Data	single	SIMD
	multiple	MIMD

The Flynn's taxonomy helps in understanding the logical subdivision of parallel systems, but it is no more up-to date; it mainly refers to HW capabilities but in 60yrs the HW evolved a lot and today we can imagine it refers to a mix of HW and SW

SISD - Single Instruction on Single Data

MISD - Multiple Instructions on Single Data

SIMD - Single Instruction on Multiple Data

MIMD - Multiple Instructions on Multiple Data



What is parallel computing ?



	HW level	SW level
SISD	A Von Neumann CPU	no parallelism at all
MISD	On a superscalar CPU, different ports executing different <i>read</i> on the same data	<ul style="list-style-type: none">• ILP on same data;• Multiple tasks or threads operating on the same data
SIMD	Any vector-capable hardware, the vector registers on a core, a GPU, a vector processor, an FPGA, ...	data parallelism through vector instructions and operations
MIMD	Every multi-core/processor system; on a superscalar CPUs, different ports executing different ops on different data	<ul style="list-style-type: none">• ILP on different data;• Multiple tasks or threads executing different code on different data.



HPC components

- from cores to clusters
- the memory

Refining the definition

High Performance Computing (HPC) is the use of servers, clusters, and supercomputers – plus associated software, tools, components, storage, and services – for scientific, engineering, or analytical tasks that are particularly intensive in computation, memory usage, or data management.

HPC is used by scientists and engineers both in research and in production across industry, government and academia.

What is High Performance Computing

Refining the definition

High Performance Computing (HPC) is the use of servers, clusters, and supercomputers – plus associated software, tools, components, storage, and services – for scientific, engineering, or analytical tasks that are particularly intensive in computation, memory usage, or data management.

HPC is used by scientists and engineers both in research and in production across industry, government and academia.

The HPC ecosystem

1. HARDWARE LEVEL

servers, cluster, supercomputers, performant I/O and networks

2. SOFTWARE LEVEL

Software, tools, programming paradigms

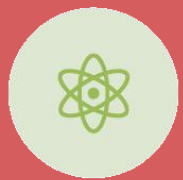
3. PROBLEMS to be solved, and theoretically sounded basis

scientific, engineering, analytical tasks

What is High Performance Computing



COMPUTATIONAL
SERVERS



ACCELERATORS



HIGH SPEED NETWORKS



HIGH END PARALLEL
STORAGE



MIDDLEWARE



SCIENTIFIC/TECHNICAL/
DATA ANALYSIS
SOFTWARE



RESEARCH/TECHNICAL
DATA



PROBLEMS TO BE
SOLVED

The HPC ecosystem

- 1. HARDWARE LEVEL**
servers, cluster, supercomputers,
performant I/O and networks
- 2. SOFTWARE LEVEL**
Software, tools, programming paradigms
- 3. PROBLEMS to be solved, and
theoretically sounded basis**
scientific, engineering, analytical tasks

The hardware behind HPC

The essential elements of a supercomputer are the following:

- a (large) number of computational “nodes” (i.e. “objects” that have processors, ram, accelerators, network ports, possibly non-volatile memories for fast storage)
- one or more switch-based networks that interconnect the nodes together; there is large variety of topologies for the networks;
- storage: usually you have 3 storage areas: home, a huge long-term resident storage (usually RAID-based) and a fast parallel file-system for production
- some login nodes to connect
- some parallel scheduler

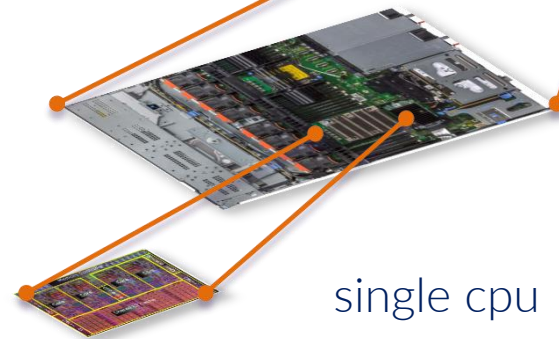




The hardware behind HPC

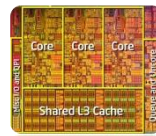
Let's now inspect in some detail the complex architecture that is behind the login node of a supercomputer, starting from the most basic element, the single CPU, whose elemental component, "the *core*", we analyzed in detail in the single-core part of the course.

interconnected
racks of connected
nodes



single nodes

single cpu



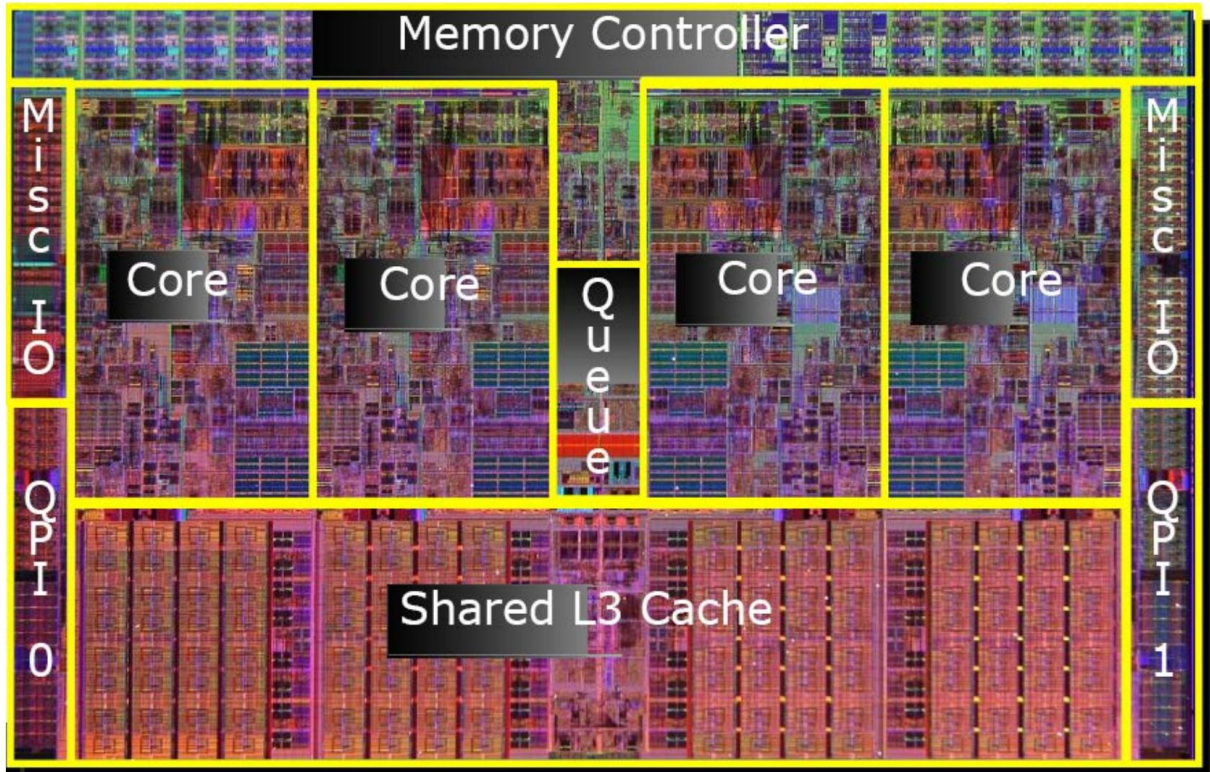
Race to
Multicore

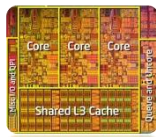
hardware
level

Single-CPU topology

Introduction

Modern
CPUs
are multi-
(or many-)
cores





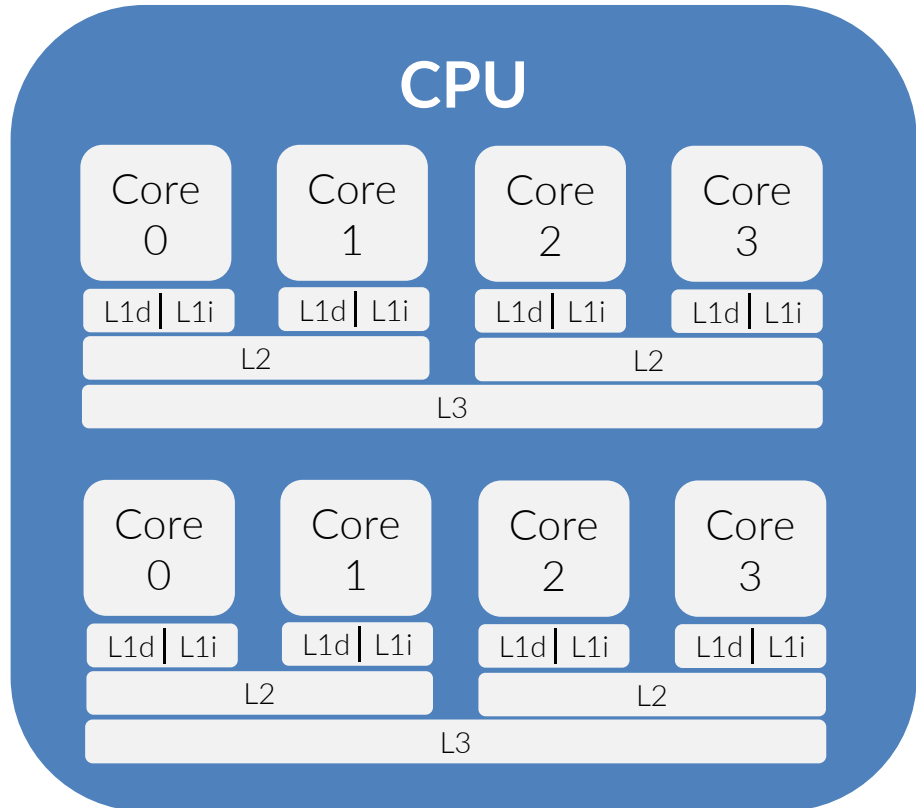
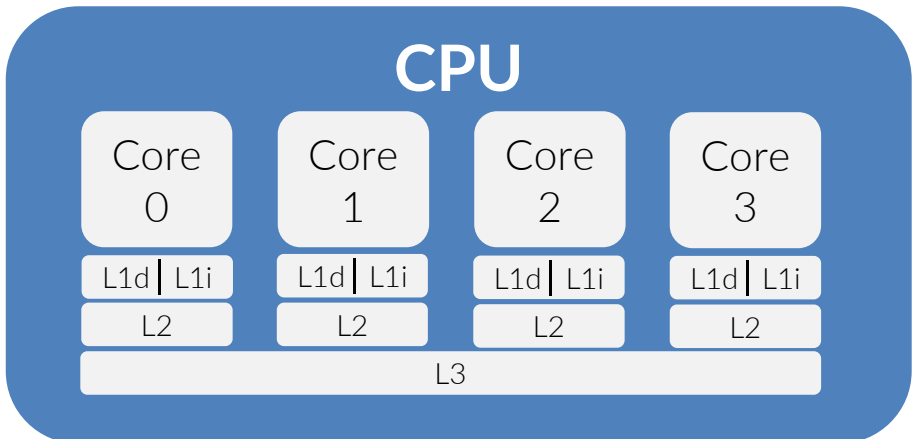
Race to
Multicore

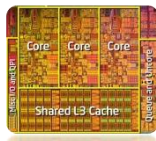
hardware
level

Single-CPU topology



Cache hierarchy can have
different topologies





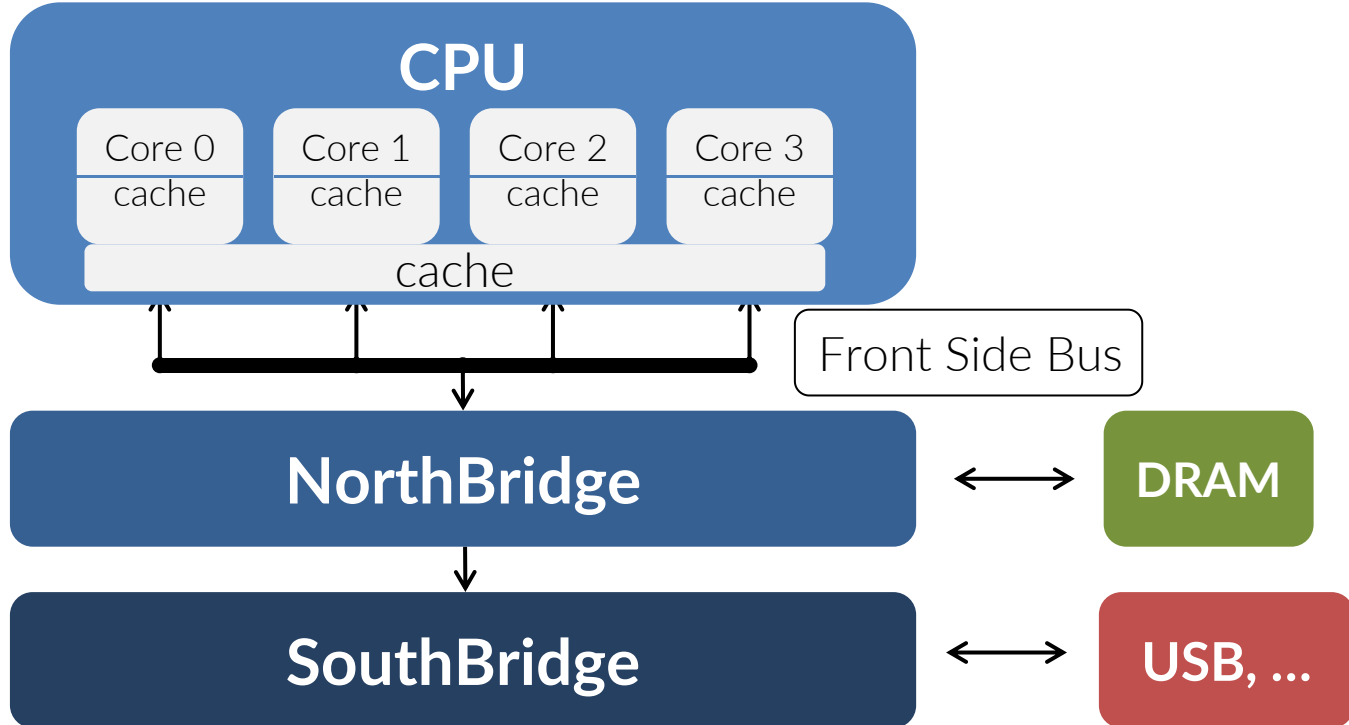
Race to
Multicore

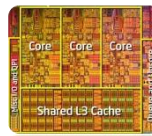
hardware
level



Single-CPU topology

In old times, this was the typical way in which a cpu connected to the outer world.
The FSB managed all communications, quickly becoming a bottleneck when an increasing nr. of cores shared its bandwidth.



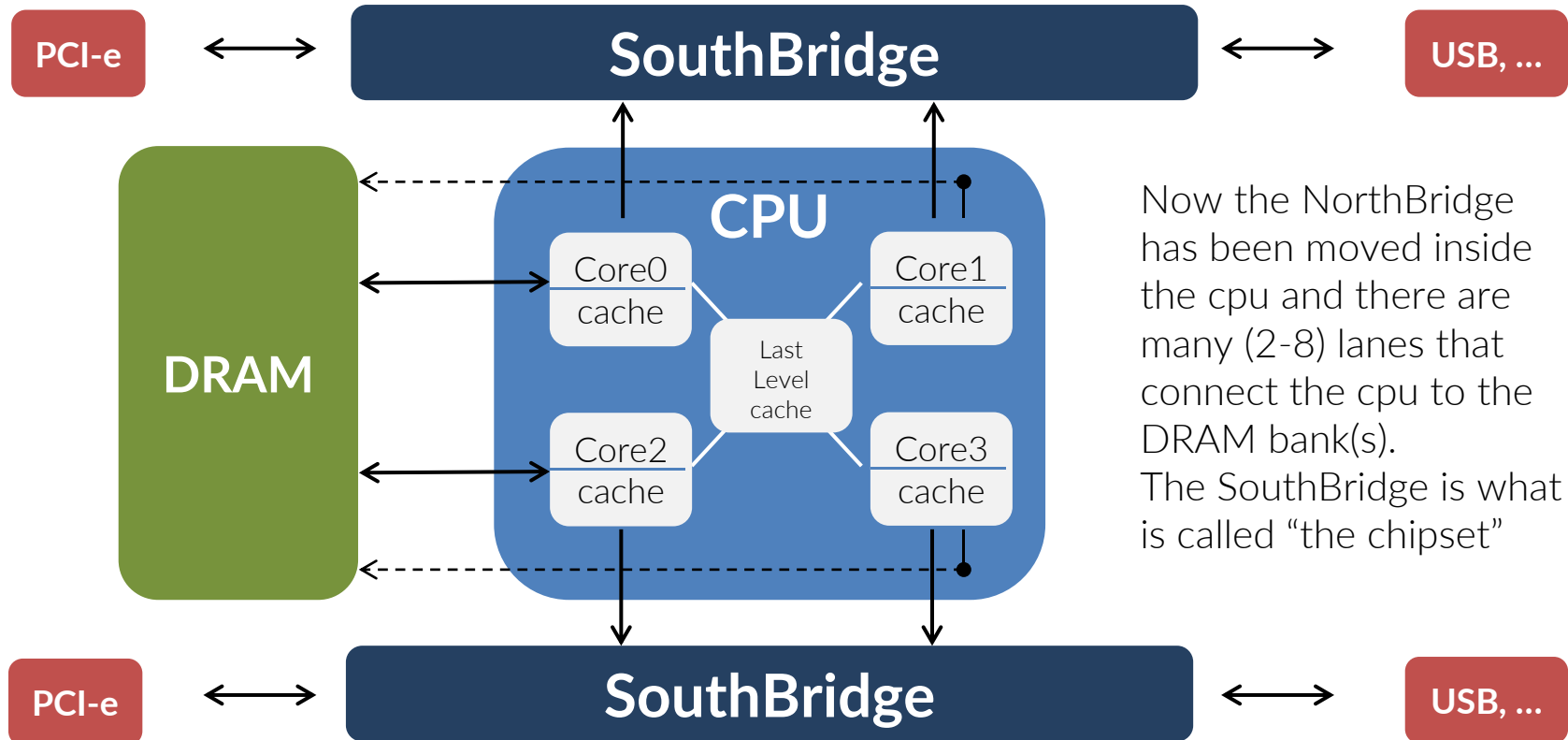


Race to
Multicore

hardware
level



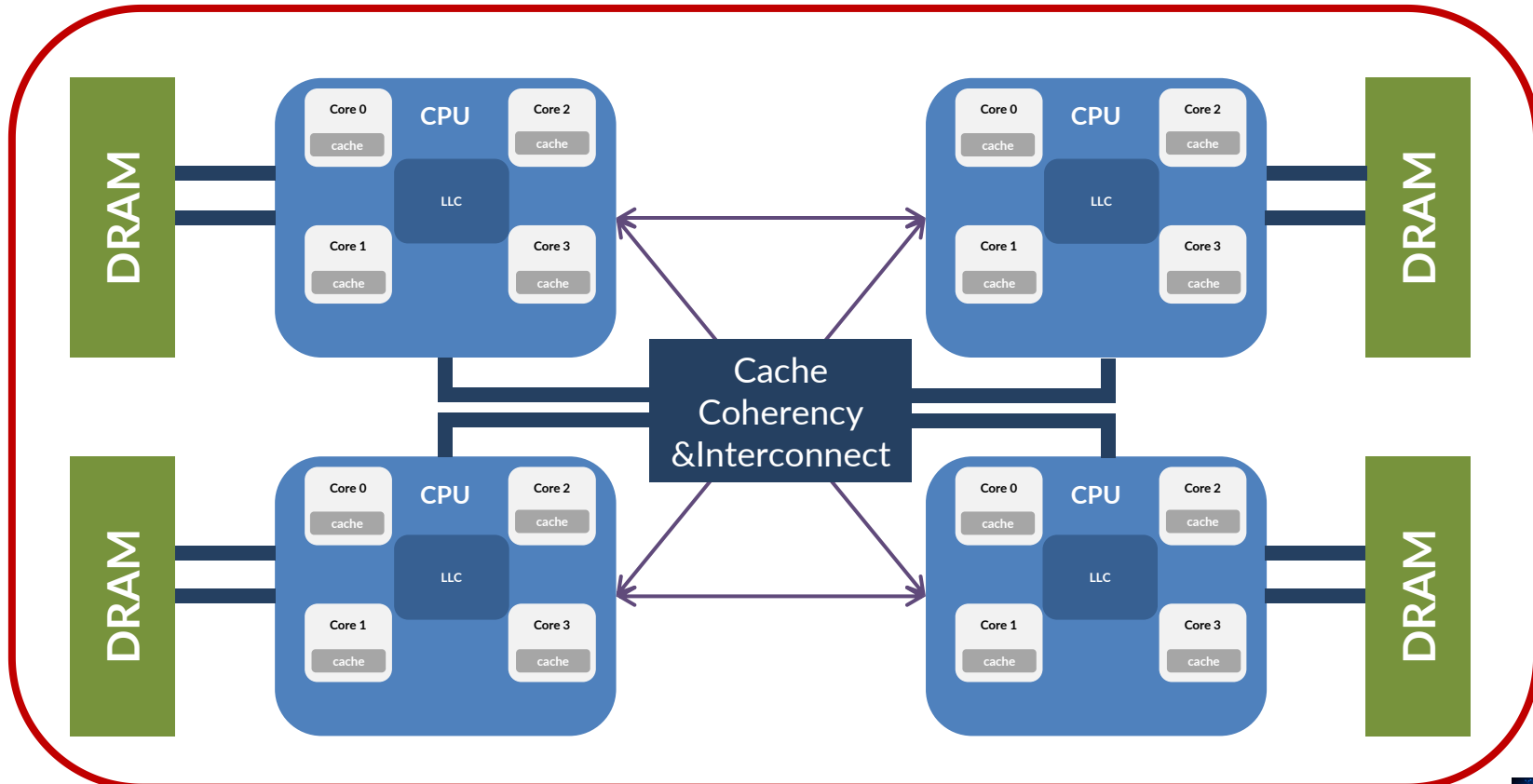
Single-CPU topology





The NODE topology

COMPUTING NODE
there may be more or less sockets
(usually an even number and ≥ 2)

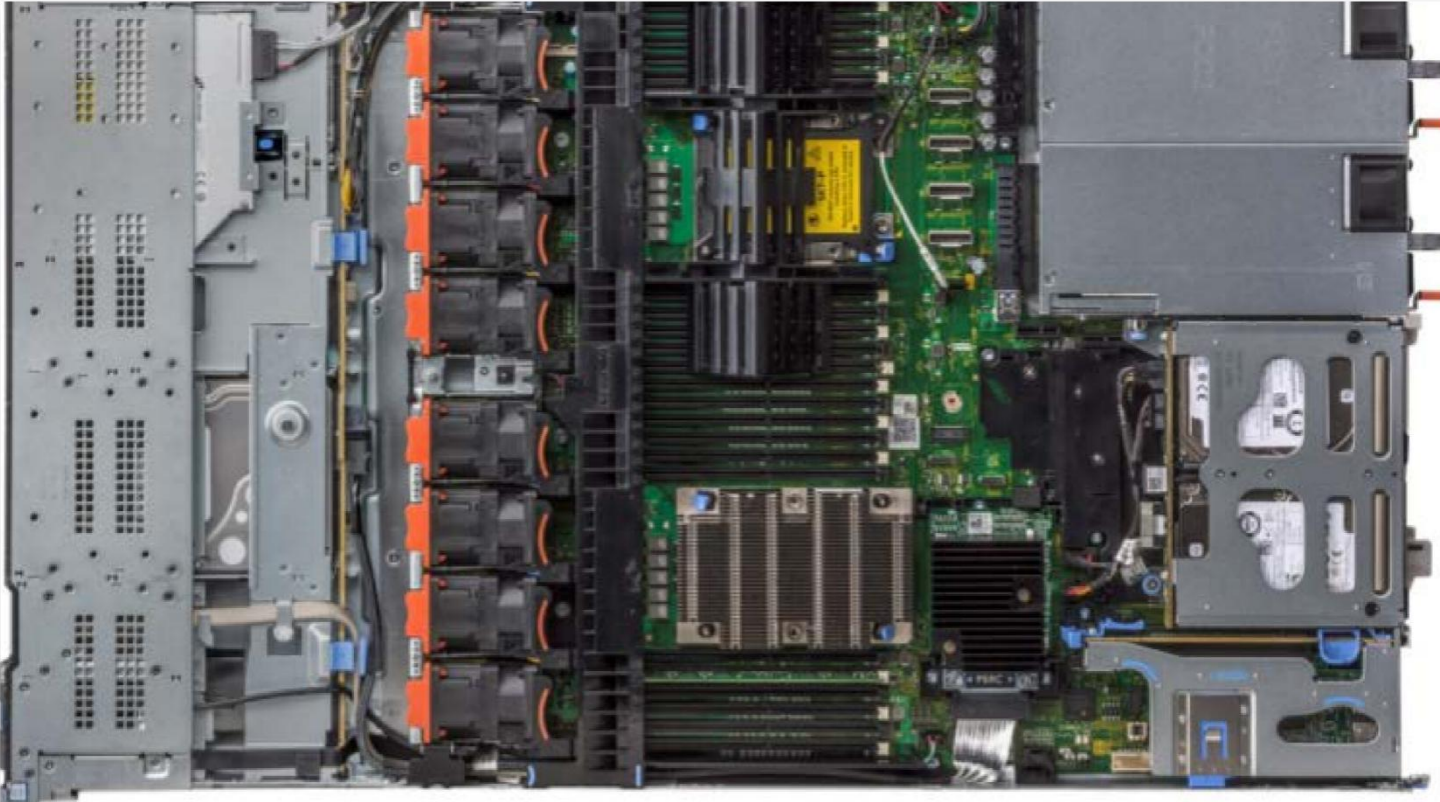




COMPUTING NODE

there may be more or less sockets
(usually an even number and ≥ 1)

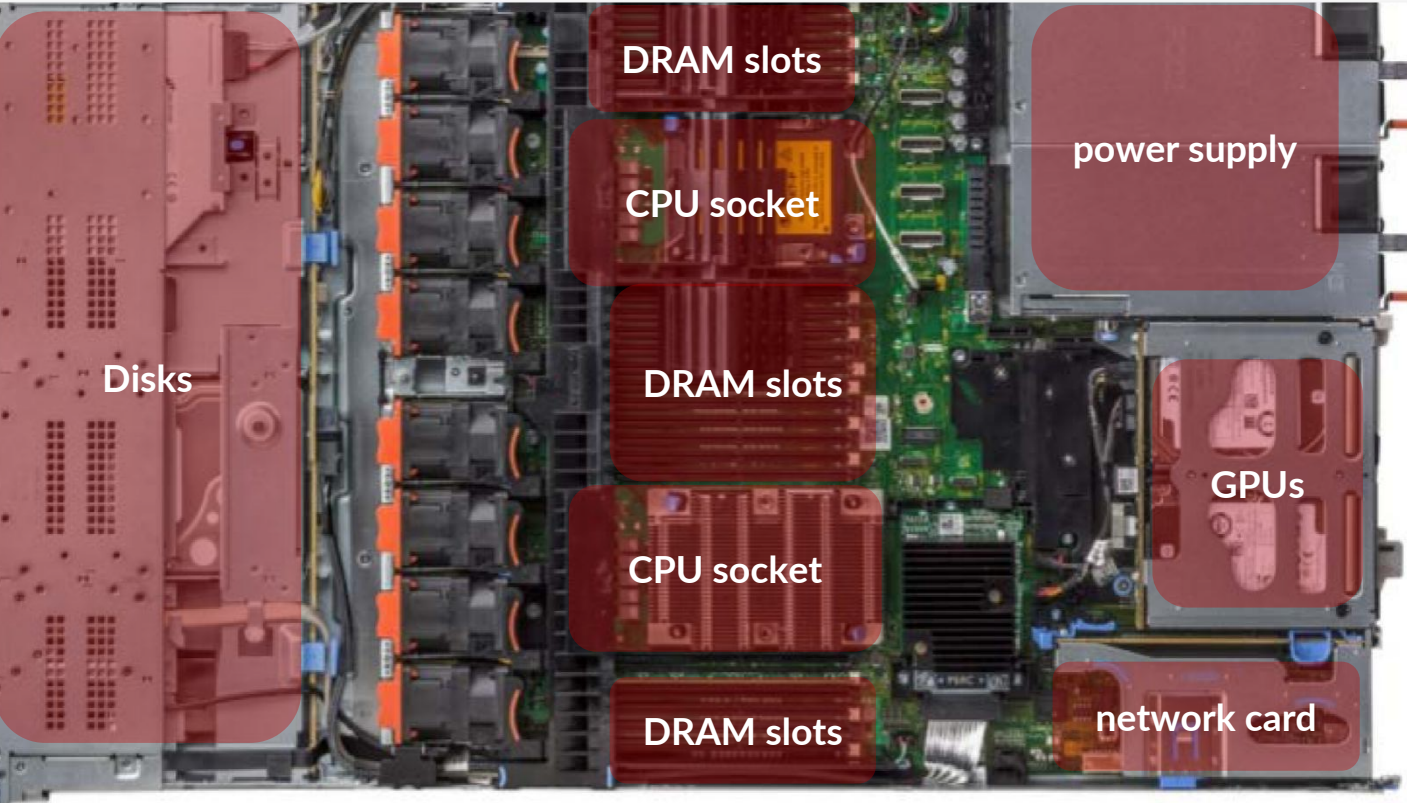
The NODE topology





COMPUTING NODE

there may be more or less sockets
(usually an even number and ≥ 1)

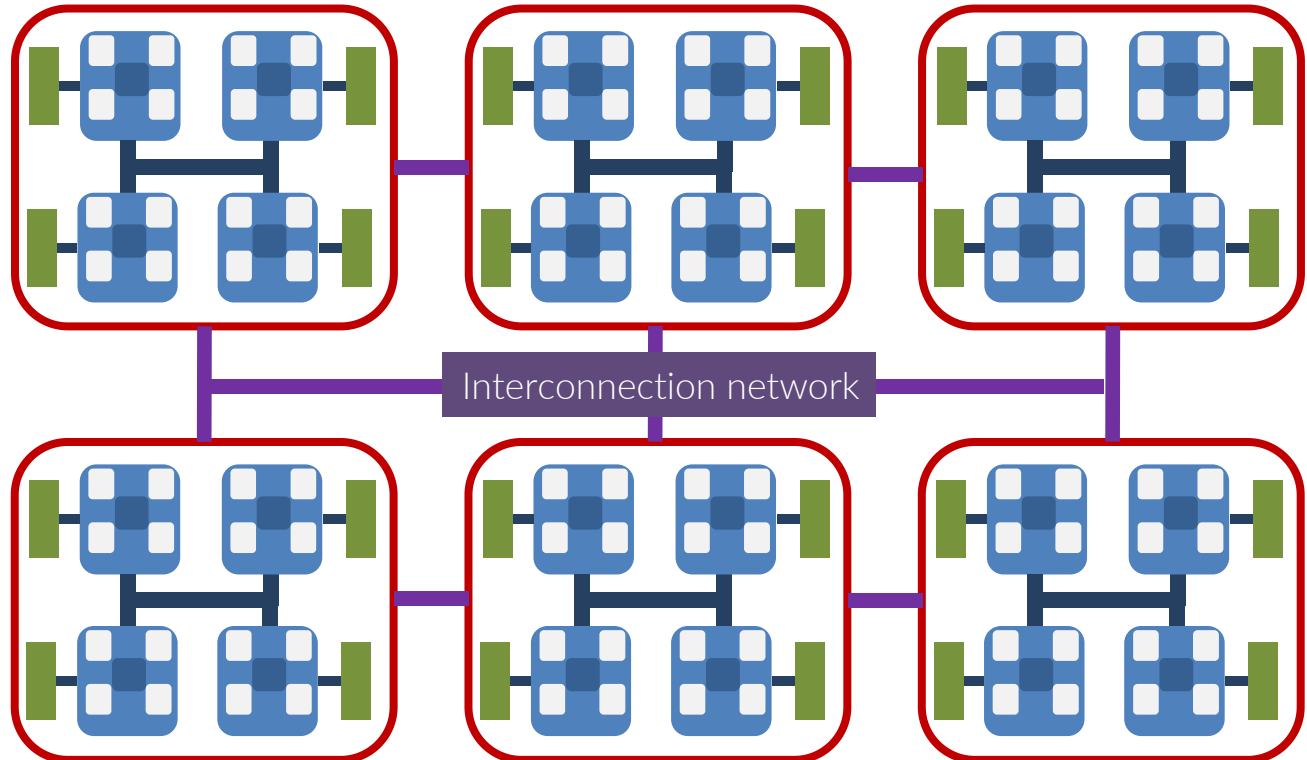


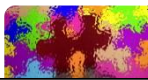


CLUSTER OF COMPUTING NODES

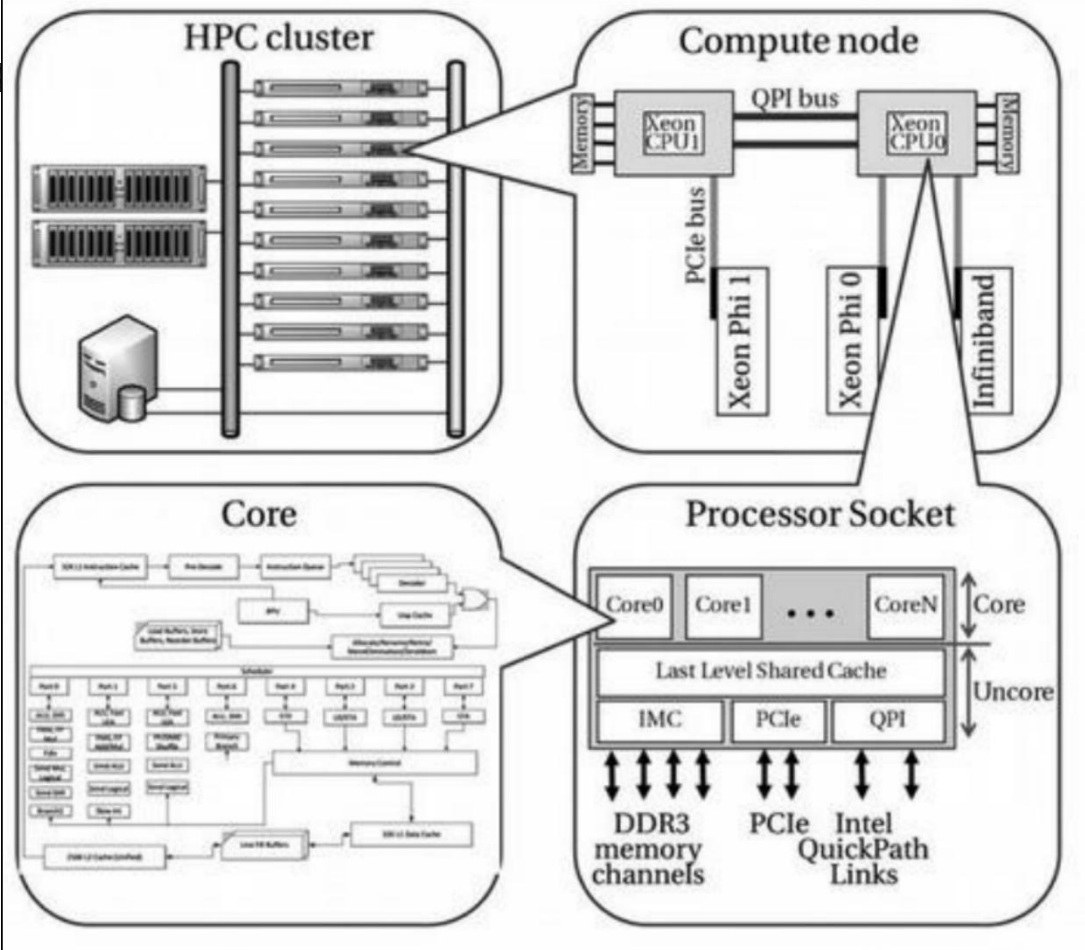
Note: there are many different topologies for the interconnection network.

The overall topology





The overall topology



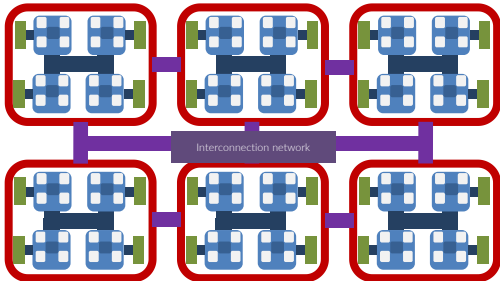
Many ($\sim 10\text{-}10^5$) nodes are connected by a switch-based network, whose topology may vary a lot. The details of that may severely affect the overall performance.

Typical figures for latency and bandwidth are $\sim 1\mu\text{s}$ and $\sim 100\text{Gbit/s}$ respectively.

The most common standard are InfiniBand and OmniPath.

The overall topology

Note that on a supercomputer there is a **hybrid approach as for the memory placement:**



- the memory on a single node can be accessed directly by all the cores on that node, meaning that memory access is a “read/write” instruction irrespectively of what exact memory bank it refers to. This is called **shared-memory**.
- when you use many nodes at a time, a process can not directly access the memory on a different node. It needs to issue a request for that, not a read/write instruction. That is named **distributed memory**.

These are *hardware concepts*, i.e. they describe how the memory is physically accessible. However, they do also refer to *programming paradigms*, as we'll see in a while.

Shared vs Distributed Memory

Then, in a multiprocessor system (i.e. a one having multiple cpus)

When some cpus can directly access the same DRAM memory, they are in **shared-memory**:

- the processes running on those cpu share a unique physical address space that is mapped on memory physically distributed on the memory banks
- read and write are simple memory accesses, and the process communicate using the shared memory

When some cpus can not access each other's DRAM memory because they are on different nodes, they are in **distributed-memory**

- the physical address space is separated and private to the processes that run on a given cpu
- the processes running on different cpu communicate through *messages* that travel on a linking network.



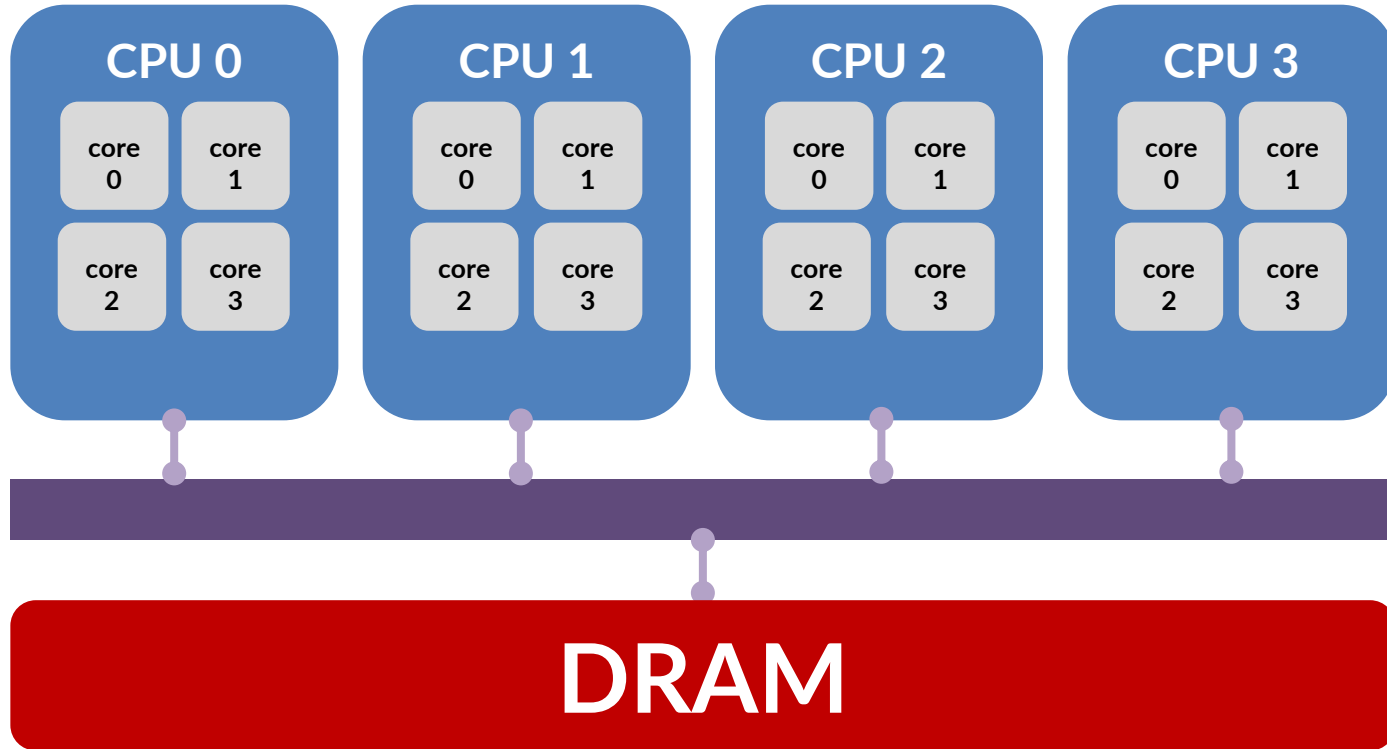
Shared Memory

In turn, there are two fundamental types of shared-memory:

- if the access to every RAM location is equally costly for all cpus, the system is **UMA, Uniform Memory Access**
- if the cost of RAM memory access depends on the location, then the system is said to be **NUMA, Non-Uniform Memory Access**



Shared Memory : UMA

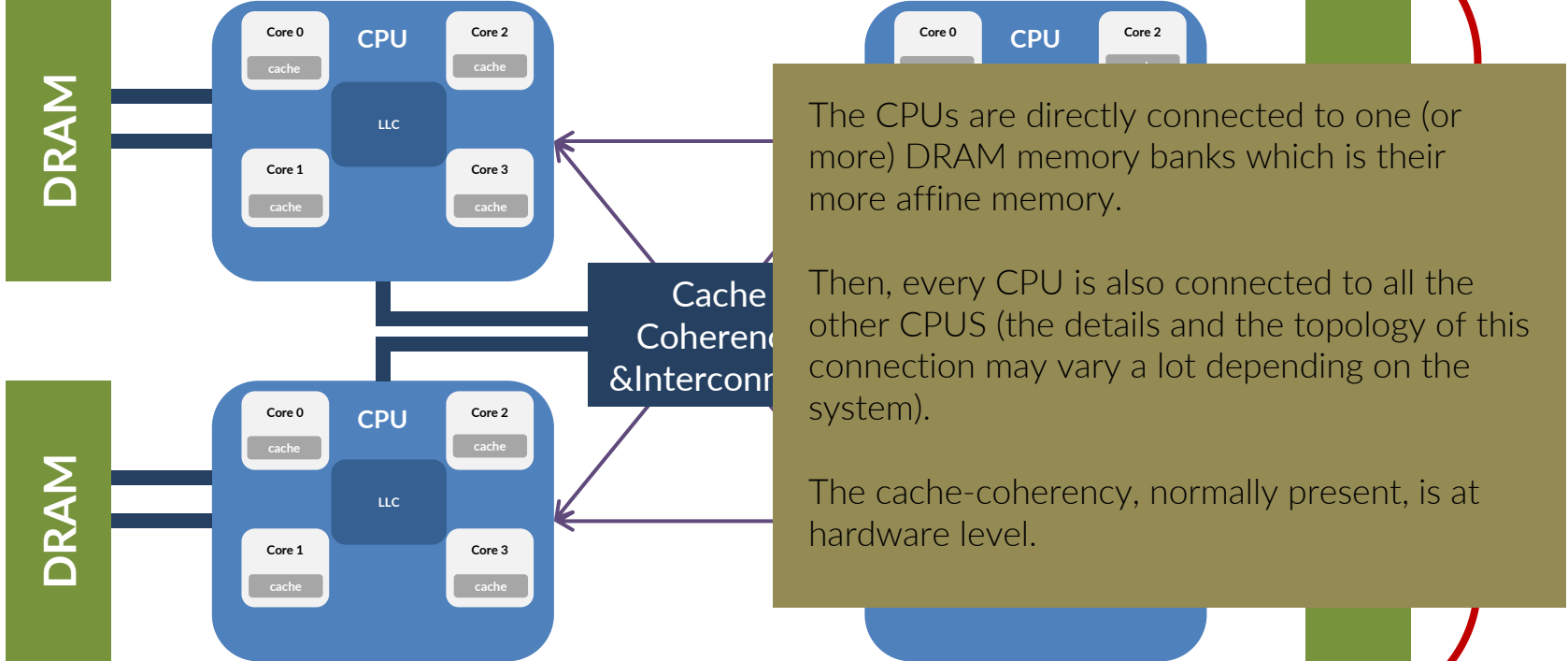


In Uniform Memory Access systems, all the cpus are connected to the DRAM banks through a common connection, and for each CPU the cost for accessing any DRAM memory location is independent on the memory location itself.



Shared Memory : NUMA

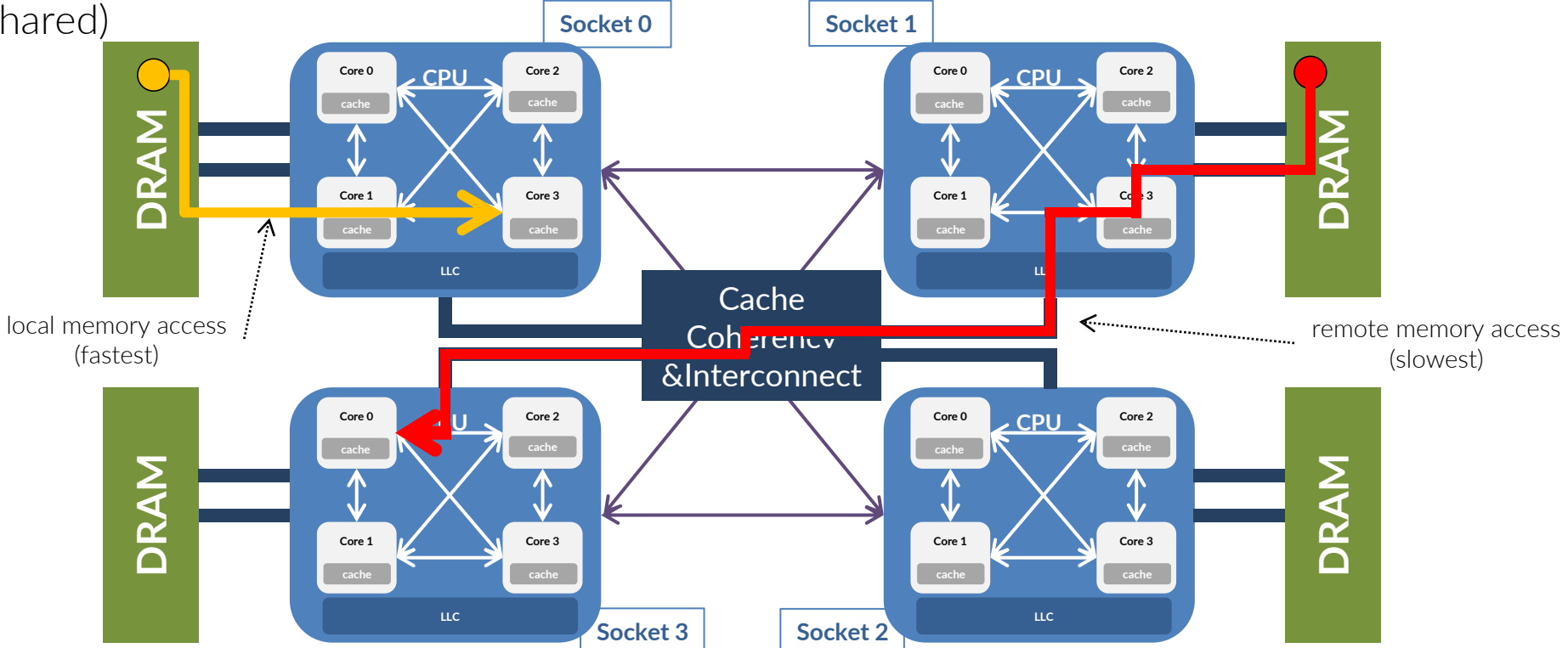
A zoom-out to a multi-socket node





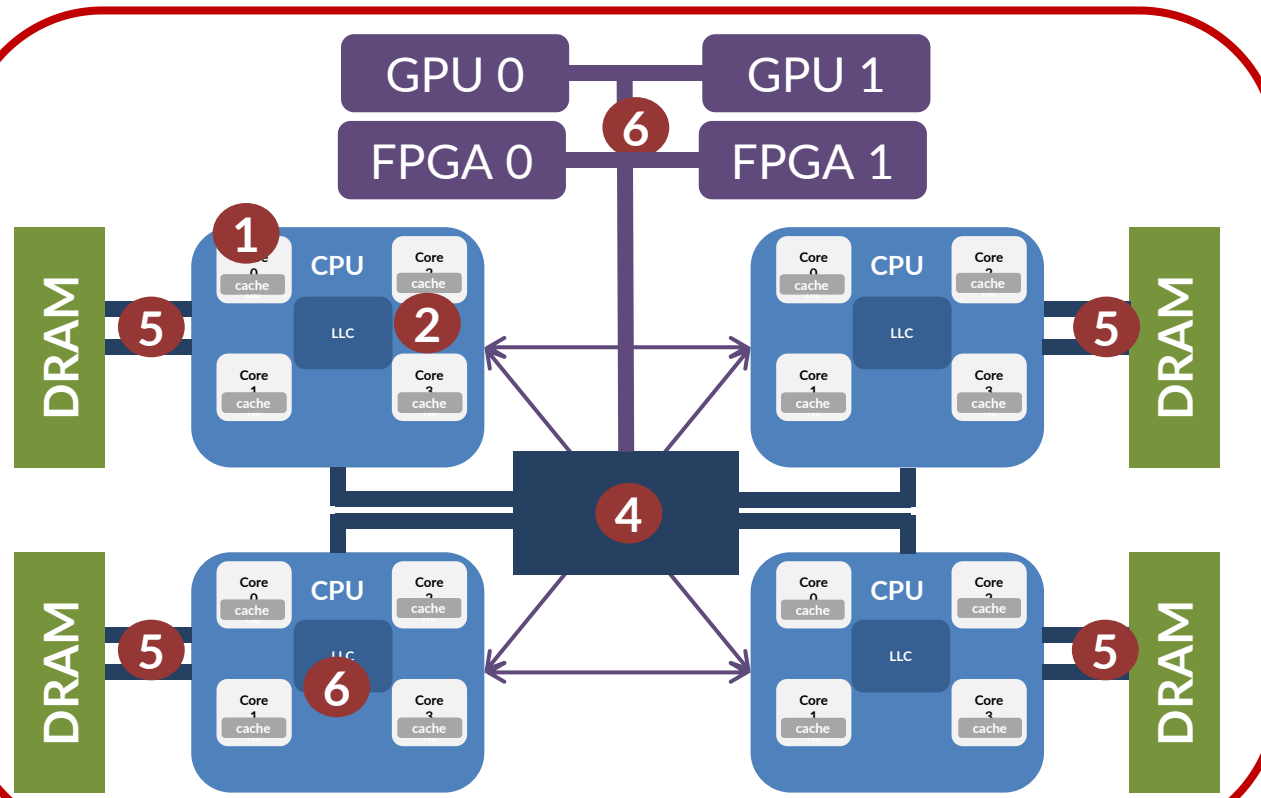
Shared Memory : NUMA

Zoom-out to a multi-socket node (all the RAM is accessible from every core, i.e. it is shared)





Summary: node-level parallelism



1. ILP/SIMD at core level
2. many cores per each socket
3. last-level cache per socket
4. inter-socket links
5. multiple memory links
6. heterogeneous accelerators



Shared Memory : NUMA

Two examples, both of nodes with 4 sockets each

```
[ltornatore@hp10 ~]$ lscpu
Architecture:          x86_64
CPU op-mode(s):        32-bit, 64-bit
Byte Order:            Little Endian
CPU(s):                40
On-line CPU(s) list:  0-39
Thread(s) per core:   1
Core(s) per socket:   10
Socket(s):             4
NUMA node(s):          4
Vendor ID:             GenuineIntel
CPU family:            6
Model:                 63
Model name:            Intel(R) Xeon(R) CPU E5-4627 v3 @ 2.60GHz
Stepping:               2
CPU MHz:               1200.000
CPU max MHz:           2600.000
CPU min MHz:           1200.000
BogoMIPS:              5194.05
Virtualization:        VT-x
L1d cache:              32K
L1i cache:              32K
L2 cache:               256K
L3 cache:               25600K
NUMA node0 CPU(s):      0-4,20-24
NUMA node1 CPU(s):      5-9,25-29
NUMA node2 CPU(s):      10-14,30-34
NUMA node3 CPU(s):      15-19,35-39
```

hyperthreading off

node distances:

node	0	1	2	3
0:	10	21	21	21
1:	21	10	21	21
2:	21	21	10	21
3:	21	21	21	10

non-uniform access to memory

```
[ltornatore@gen10-01 ~]$ lscpu
Architecture:          x86_64
CPU op-mode(s):        32-bit, 64-bit
Byte Order:            Little Endian
CPU(s):                96
On-line CPU(s) list:  0-95
Thread(s) per core:   2
Core(s) per socket:   12
Socket(s):             4
NUMA node(s):          4
Vendor ID:             GenuineIntel
CPU family:            6
Model:                 85
Model name:            Intel(R) Xeon(R) Gold 5118 CPU @ 2.30GHz
Stepping:               4
CPU MHz:               2663.305
CPU max MHz:           3200.0000
CPU min MHz:           1000.0000
BogoMIPS:              4600.00
Virtualization:        VT-x
L1d cache:              32K
L1i cache:              32K
L2 cache:               1024K
L3 cache:               16896K
NUMA node0 CPU(s):      0-11,48-59
NUMA node1 CPU(s):      12-23,60-71
NUMA node2 CPU(s):      24-35,72-83
NUMA node3 CPU(s):      36-47,84-95
```

hyperthreading on

node distances:

node	0	1	2	3
0:	10	21	21	21
1:	21	10	21	21
2:	21	21	10	21
3:	21	21	21	10

Discovering a NUMA topology

How to discover the topology of your node:

- **numactl** tool
it also controls the Linux NUMA policy
- **cpuinfo** tool (by Intel)
- **hwloc** (by OpenMPI)



Shared Memory : NUMA

How to discover the topology of your node (examples):

- **numactl** tool

numactl -H

- **cpuinfo** tool (by Intel)

- **hwloc** (by OpenMPI)

```
[ltornatore@hp08 ~]$ numactl -H
available: 4 nodes (0-3)
node 0 cpus: 0 1 2 3 4 20 21 22 23 24
node 0 size: 65411 MB
node 0 free: 40998 MB
node 1 cpus: 5 6 7 8 9 25 26 27 28 29
node 1 size: 65536 MB
node 1 free: 58475 MB
node 2 cpus: 10 11 12 13 14 30 31 32 33 34
node 2 size: 65536 MB
node 2 free: 59344 MB
node 3 cpus: 15 16 17 18 19 35 36 37 38 39
node 3 size: 65536 MB
node 3 free: 59641 MB
node distances:
node   0   1   2   3
 0: 10 21 21 21
 1: 21 10 21 21
 2: 21 21 10 21
 3: 21 21 21 10
```



Shared Memory : NUMA

How to discover the topology of your node (examples):

- **numactl** tool
- **cpuinfo** tool (by Intel)
- **hwloc** (by OpenMPI)

```
cpuinfo -d
=====
Placement on packages
Package Id. Core Id. Processors
0          0,2,4,9,11,1,3,8,10,12      0,1,2,3,4,20,21,22,23,24
1          0,2,4,9,11,1,3,8,10,12      5,6,7,8,9,25,26,27,28,29
2          0,2,4,9,11,1,3,8,10,12      10,11,12,13,14,30,31,32,33,34
3          0,2,4,9,11,1,3,8,10,12      15,16,17,18,19,35,36,37,38,39
```

```
cpuinfo -g
=====
Processor composition
Processor name   : Intel(R) Xeon(R) E5-4627 v3
Packages(sockets) : 4
Cores           : 40
Processors(CPUs) : 40
Cores per package : 10
Threads per core  : 1
```

```
cpuinfo -c
=====
Cache sharing
Cache  Size      Processors
L1    32 KB     no sharing
L2    256 KB    no sharing
L3    25 MB      (0,1,2,3,4,20,21,22,23,24)(5,6,7,8,9,25,26,27,28,29)(10,11,12,13,14,30,31,32,33,34)(15,16,17,18,19,35,36,37,38,39)
```

Shared Memory : NUMA

How to discover the topology of a system

- **numactl** tool
- **cpuinfo** tool (by Intel)
- **hwloc-ls** (by OpenMPI)
hwloc-info

Machine (256GB total)

NUMANode L#0 (P#0 64GB)

Package L#0 + L3 L#0 (25MB)
 L2 L#0 (256KB) + L1d L#0 (32KB) + L1i L#0 (32KB) + Core L#0 + PU L#0 (P#0)
 L2 L#1 (256KB) + L1d L#1 (32KB) + L1i L#1 (32KB) + Core L#1 + PU L#1 (P#1)
 L2 L#2 (256KB) + L1d L#2 (32KB) + L1i L#2 (32KB) + Core L#2 + PU L#2 (P#2)
 L2 L#3 (256KB) + L1d L#3 (32KB) + L1i L#3 (32KB) + Core L#3 + PU L#3 (P#3)
 L2 L#4 (256KB) + L1d L#4 (32KB) + L1i L#4 (32KB) + Core L#4 + PU L#4 (P#4)
 L2 L#5 (256KB) + L1d L#5 (32KB) + L1i L#5 (32KB) + Core L#5 + PU L#5 (P#20)
 L2 L#6 (256KB) + L1d L#6 (32KB) + L1i L#6 (32KB) + Core L#6 + PU L#6 (P#21)
 L2 L#7 (256KB) + L1d L#7 (32KB) + L1i L#7 (32KB) + Core L#7 + PU L#7 (P#22)
 L2 L#8 (256KB) + L1d L#8 (32KB) + L1i L#8 (32KB) + Core L#8 + PU L#8 (P#23)
 L2 L#9 (256KB) + L1d L#9 (32KB) + L1i L#9 (32KB) + Core L#9 + PU L#9 (P#24)

NUMANode L#1 (P#1 64GB) + Package L#1 + L3 L#1 (25MB)

L2 L#10 (256KB) + L1d L#10 (32KB) + L1i L#10 (32KB) + Core L#10 + PU L#10 (P#5)
 L2 L#11 (256KB) + L1d L#11 (32KB) + L1i L#11 (32KB) + Core L#11 + PU L#11 (P#6)
 L2 L#12 (256KB) + L1d L#12 (32KB) + L1i L#12 (32KB) + Core L#12 + PU L#12 (P#7)
 L2 L#13 (256KB) + L1d L#13 (32KB) + L1i L#13 (32KB) + Core L#13 + PU L#13 (P#8)
 L2 L#14 (256KB) + L1d L#14 (32KB) + L1i L#14 (32KB) + Core L#14 + PU L#14 (P#9)
 L2 L#15 (256KB) + L1d L#15 (32KB) + L1i L#15 (32KB) + Core L#15 + PU L#15 (P#25)
 L2 L#16 (256KB) + L1d L#16 (32KB) + L1i L#16 (32KB) + Core L#16 + PU L#16 (P#26)
 L2 L#17 (256KB) + L1d L#17 (32KB) + L1i L#17 (32KB) + Core L#17 + PU L#17 (P#27)
 L2 L#18 (256KB) + L1d L#18 (32KB) + L1i L#18 (32KB) + Core L#18 + PU L#18 (P#28)
 L2 L#19 (256KB) + L1d L#19 (32KB) + L1i L#19 (32KB) + Core L#19 + PU L#19 (P#29)

NUMANode L#2 (P#2 64GB) + Package L#2 + L3 L#2 (25MB)

L2 L#20 (256KB) + L1d L#20 (32KB) + L1i L#20 (32KB) + Core L#20 + PU L#20 (P#10)
 L2 L#21 (256KB) + L1d L#21 (32KB) + L1i L#21 (32KB) + Core L#21 + PU L#21 (P#11)
 L2 L#22 (256KB) + L1d L#22 (32KB) + L1i L#22 (32KB) + Core L#22 + PU L#22 (P#12)
 L2 L#23 (256KB) + L1d L#23 (32KB) + L1i L#23 (32KB) + Core L#23 + PU L#23 (P#13)
 L2 L#24 (256KB) + L1d L#24 (32KB) + L1i L#24 (32KB) + Core L#24 + PU L#24 (P#14)
 L2 L#25 (256KB) + L1d L#25 (32KB) + L1i L#25 (32KB) + Core L#25 + PU L#25 (P#30)
 L2 L#26 (256KB) + L1d L#26 (32KB) + L1i L#26 (32KB) + Core L#26 + PU L#26 (P#31)
 L2 L#27 (256KB) + L1d L#27 (32KB) + L1i L#27 (32KB) + Core L#27 + PU L#27 (P#32)
 L2 L#28 (256KB) + L1d L#28 (32KB) + L1i L#28 (32KB) + Core L#28 + PU L#28 (P#33)
 L2 L#29 (256KB) + L1d L#29 (32KB) + L1i L#29 (32KB) + Core L#29 + PU L#29 (P#34)

NUMANode L#3 (P#3 64GB) + Package L#3 + L3 L#3 (25MB)

L2 L#30 (256KB) + L1d L#30 (32KB) + L1i L#30 (32KB) + Core L#30 + PU L#30 (P#15)
 L2 L#31 (256KB) + L1d L#31 (32KB) + L1i L#31 (32KB) + Core L#31 + PU L#31 (P#16)
 L2 L#32 (256KB) + L1d L#32 (32KB) + L1i L#32 (32KB) + Core L#32 + PU L#32 (P#17)
 L2 L#33 (256KB) + L1d L#33 (32KB) + L1i L#33 (32KB) + Core L#33 + PU L#33 (P#18)
 L2 L#34 (256KB) + L1d L#34 (32KB) + L1i L#34 (32KB) + Core L#34 + PU L#34 (P#19)
 L2 L#35 (256KB) + L1d L#35 (32KB) + L1i L#35 (32KB) + Core L#35 + PU L#35 (P#35)
 L2 L#36 (256KB) + L1d L#36 (32KB) + L1i L#36 (32KB) + Core L#36 + PU L#36 (P#36)
 L2 L#37 (256KB) + L1d L#37 (32KB) + L1i L#37 (32KB) + Core L#37 + PU L#37 (P#37)
 L2 L#38 (256KB) + L1d L#38 (32KB) + L1i L#38 (32KB) + Core L#38 + PU L#38 (P#38)
 L2 L#39 (256KB) + L1d L#39 (32KB) + L1i L#39 (32KB) + Core L#39 + PU L#39 (P#39)



Shared Memory : NUMA

How to discover the topology of your node:

- `numactl` tool
- `cpuinfo` tool (by Intel)
- `hwloc` (by OpenMPI)
- directly exploring `/sys/devices/system/`

Cache coherence

Data synchronization is one of the main performance killers for multi-core applications.

- when a memory region is accessed by two cores (i.e. by two different threads running on two different cores), it must be present in both L1/L2, and when one core updates the value stored in the region, the change must be propagated.
- when a thread migrates, the data will still resides on another's core memory.

Memory consistency for the whole system is guaranteed at hardware level, resulting in huge wasting of time if data are not properly handled.
For instance, concurrent access in writing is a main sink of cpu cycles.



Cache coherence: MESI

Data consistency is maintained by the **MESI** standard.

It is the successor of the MSI protocol and the ancestor of MESOI one

MODIFIED

X's values has been modified by this core, and then this is the only valid copy in the system

EXCLUSIVE

X is used by this core only; changes do not need to be signalled

SHARED

X is used by multiple cores; changes need to be signalled

INVALID

X's value has been modified by another core (or X is not used)

see:
MD64 Architecture Programmer's Manual
Volume 2: System Programming

In the above, "X" stands for any given memory location

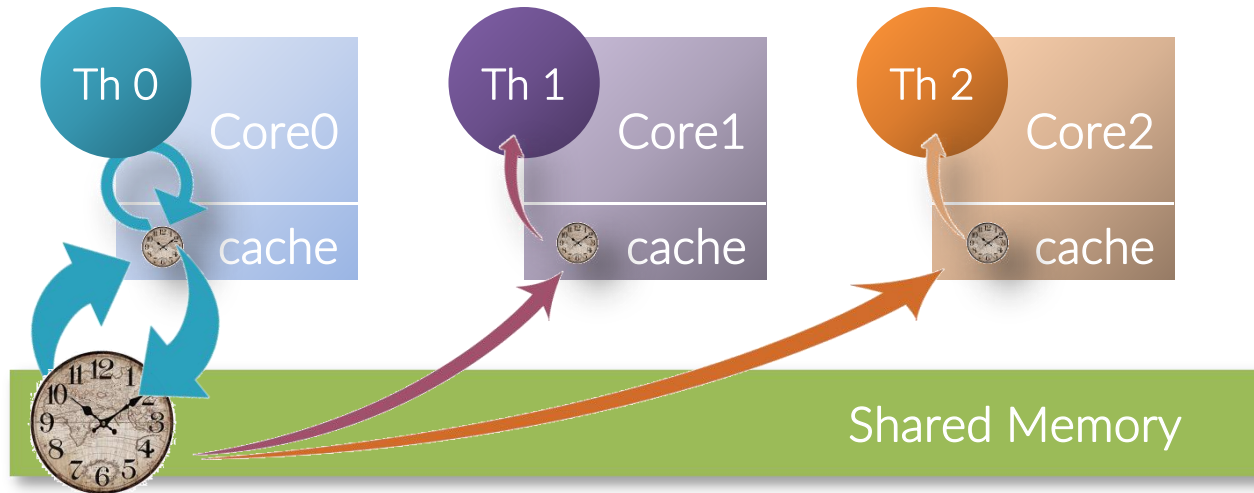


MESI – an example



Let's clarify with an example. Let's say that there are 3 threads, running on separate cores, accessing some shared-memory.

Thread0 is running the application `clock()`, which ticks a shared-memory variable that contains the wall-clock time. In time to time, both Thread1 and Thread2 want to know what time it is.





MESI – an example

		Time (in secs)	Action	cache status		
M	Core0	Core1	Core2			
M	Modified					
E	Exclusive	0	-	I	I	I
S	Shared	1	Th0 reads	E	I	I
I	Invalid	2	Th0 writes ⁰	E	I	I
		2.3	Th1 reads ¹	S	S	I
		2.7	Th2 reads	S	S	S
		3	Th0 writes ²	M	I	I
		4	Th0 writes	M	I	I
		4.4	Th2 reads ¹	S	I	S
		5	Th0 writes	M	I	I
	

0 Core0 is the only one using the value, that is then “Exclusive”. No signal needs to be sent around.

1 A signal is issued to “the memory”, which recognizes that the only valid copy is in the Core0 cache.

Hence, that value is copied back into the shared memory, and from there it is copied in the cache. At that point, everybody has a valid copy, which is then “Shared” (*).

2 A signal about the change is issued to all the interested actors (those who have a copy) because their values are now “Invalid”. O’s copy is instead “Modified”.

(*) In the MESOI protocol, in this case the copy can be sent directly to the other caches, without having to transit by the DRAM.

Great powers, great responsibility

A

Variables used by a single core

They should resides in a single cache

B

Read-only variables

No issues in being shared among many cores

C

Modified variables

Variables modified by many cores, or read by many cores

If possibler, a cache line should contain only one type of data (A, B or C on the left).

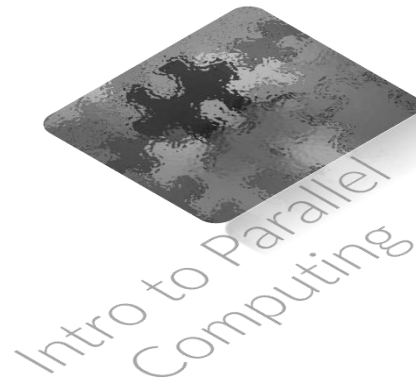
Put variables in the order they will be used.

Type C, modified variables, should stay together, they are the bottlenecks.

The **false sharing** happens when variables of type A or B resides in the same cache line of a type C. Or when two type C variables, modified by two different cores, reside in the same cache line.



Introduction Outline



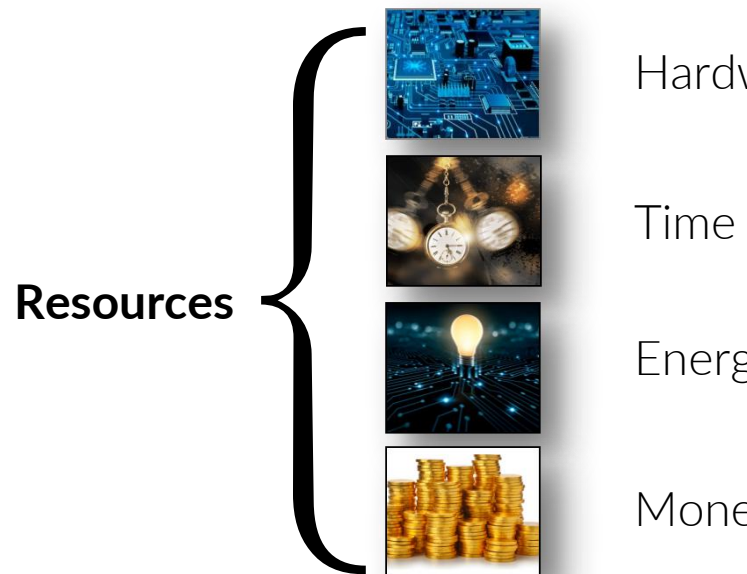
Parallel
Performance



What is parallel performance

Has we have seen, «performance» is a tag that can stand for many things.

In this frame, with «performance» we mean the relation between the computational requirements and the computational resources needed to meet those requirements.



$$\text{Performance} \approx \frac{1}{\text{resources}}$$

$$\text{Performance ratios} \approx \frac{\text{resources}_1}{\text{resources}_2}$$



What is parallel performance



Performance is a measure of how well the computational requirements are met and, at the same time, of how well the computational resources are exploited.

“The most constant difficulty in contriving the engine has arisen from the desire to reduce the time in which the calculations were executed to the shortest which is possible.”

Charles Babbage, 1791 – 1871



Key factors



n	Problem size
$T_s(n)$	Serial run-time for a problem of size n
$T_p(n)$	Parallel run-time with p processes for a problem of size n
p	Number of computing units
f_n	Intrinsic sequential fraction of the problem of size n
$k(n, t)$	Parallel overhead
Speedup	$Sp(n, t) = \frac{T_s(n)}{T_p(n)}$
Efficiency	$Eff(n, t) = \frac{T_s(n)}{p \times T_p(n)} = \frac{Sp(n, t)}{p}$



Naïve expectations



- If single processor has $\sim m$ Mflops, parallel flops performance with p tasks is $p \times m$ Mflops.
- If sequential run-time is T , parallel run-time with p tasks is $\propto T/p$.
- If parallel run-time with p tasks is T , and the run-time with p_1 tasks is T_1 , then $T_1/T_2 \propto p_2/p_1$
- If parallel run-time with p tasks and problem size Z is T , the run-time with size Z_1 is $T_1 \propto T \times Z_1/Z$.

Is that correct ?



Parallel performance



The sequential execution time for a problem of size n is

$$T_S(n) = T_S(n) \times f_n + T_S(n) \times (1-f_n)$$

Assuming that the parallel fraction of the computation is *perfectly parallel*, meaning that its run time scales as $1/p$, then we can express the parallel execution time as

$$T_P(n) = T_S(n, 1) \times f_n + T_S(n) \times (1-f_n)/p$$

And then

$$\textbf{speedup} : Sp(n, p) = T_S(n) / T_P(n) = \frac{1}{f + \frac{1-f}{p}} \quad \lim_{p \gg 1} Sp(n, p) = \frac{1}{f}$$

$$\textbf{efficiency} : Eff(n, p) = Sp(n, p) / p = \frac{1}{f(p-1)+1} \quad \lim_{p \gg 1} Eff(n, p) = 0$$



Parallel performance



If the parallel fraction ($1-f_n$) is not perfectly parallelizable, we must take into account some *parallel* overhead, and then we can express the parallel execution time as

$$T_P(n) = T(n,p) = T_S \times f_n + T_S \times (1-f_n)/p + k(n,t)$$

And then

$$\text{speedup} = Sp(n,p) \leq T_S / T_P$$

$$\text{efficiency} = Eff(n,p) \leq Sp(n,p) / p$$



Amdahl's law



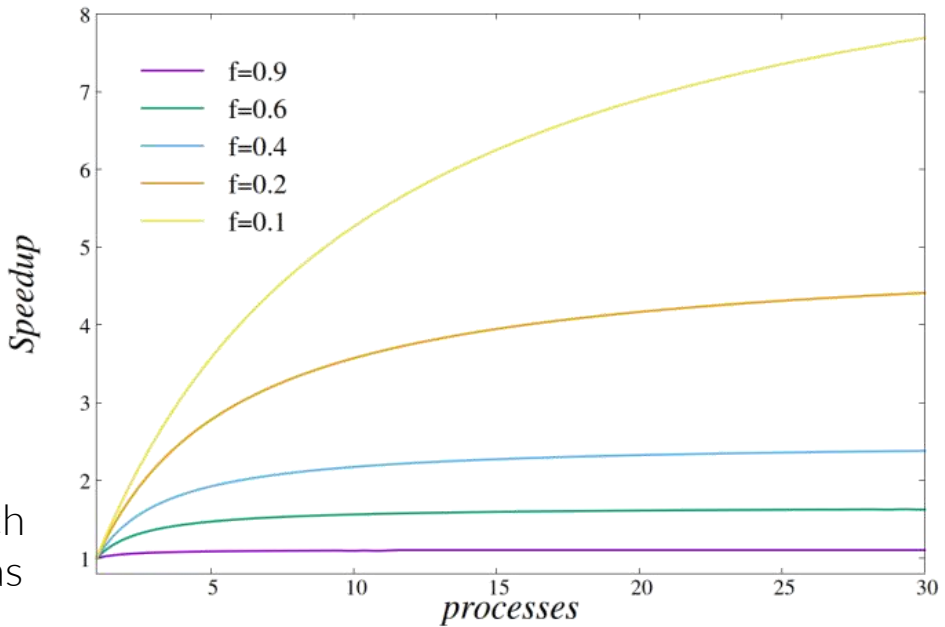
If f is the fraction of the code which is intrinsically sequential, as we have seen the speedup is

$$Sp(n, t) \leq \frac{1}{f + \frac{1-f}{p}} \leq \frac{1}{f}$$

i.e. the serial fraction is severely limiting the achievable speedup.

Is that a problem ?

Actually tens of years ago there was a huge debate about whether or not a huge effort towards developing massive parallelism (which means developing hw and ws technology) was in order or not.





Amdahl's law



There are some significant issues in the Amdhal's law shown in the previous slide:

- No matter of how many processes p are used, the maximum achievable speedup is determined by f (and is quite low for ordinary problems).
- The problem size n is kept fixed when estimating the possible speedup while the number of processes increases (*strong scaling*). However, most often the problem size increases as well.
- The parallel overhead $k(n, t)$ is ignored, which leads to an optimistic estimate of the speedup, and usually, $p(n)/t > k(n,t)$
- The fraction of sequential part may decrease when the problem size increases

Then, usually the speedup increases with problem size



Gustafson's law



However, normally when you increase the problem's size, the parallelizable part increases way more than the sequential part.

If we consider the workload as the sum

$$w = a+b$$

where a and b being the serial and the parallel work, and we assign the same amount of workload to every process, that would amount to a serial run-time

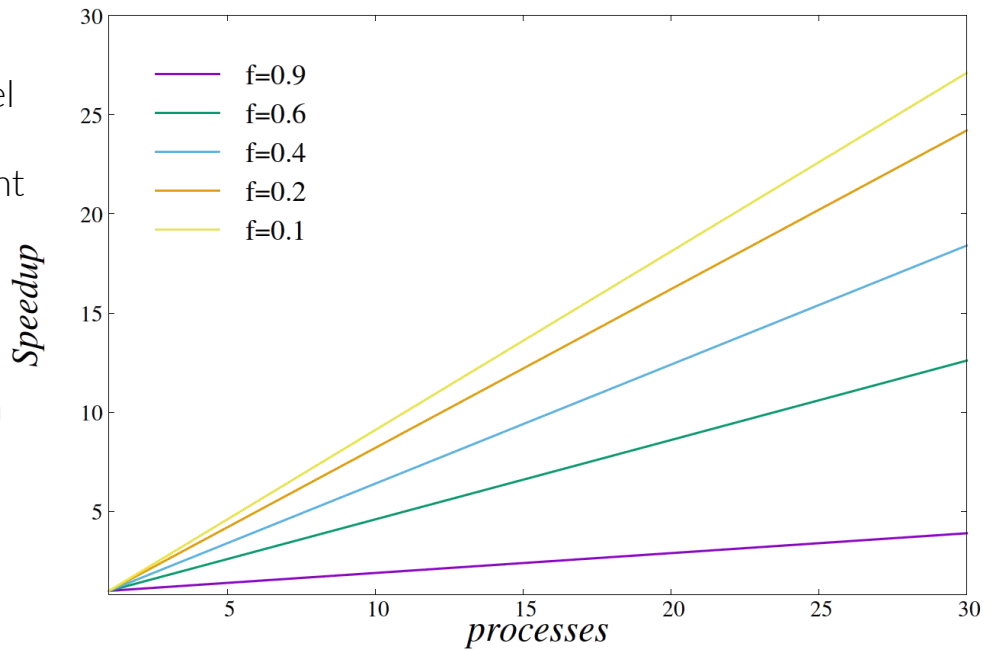
$$T_s \propto a + p \times b$$

while it still takes

$$T_p \propto a+b \text{ using } p \text{ processes}$$

Hence the speedup is $[a + p \times b] / [a+b]$, which if $f_n = a/(a+b)$ we can rewrite as the Gustafson's law for the speedup:

$$Sp_G(n, t) = p - (p - 1)f_n \leq p$$





Scalability



The two lines of reasoning, the former by Amdhal and the latter by Gustafson, lead us to two different concepts for the *scalability*, which is the ability of a parallel system to increase its efficiency when the number of processes and/or the size of the problem get larger.

- 1. STRONG SCALABILITY:** the problem size is fixed, p increases
- 2. WEAK SCALABILITY:** the workload is fixed, the problem size and p increase



Parallel overhead



In parallel computing there may be several sources of overhead due to the parallelization itself:

- Communication overhead
- Algorithmic overhead
- Synchronization
 - Critical paths - Dependencies across different processes
 - Bottlenecks (some processes are stuck and make all the others wait)
 - Work-load imbalance
- Thread/processes creation

Hence, if $k(n,t)$ is the overhead of some kind, t_S and t_P the run-time for the serial and the parallel part, the parallel run-time can be written as

$$T_P(n, p) = t_S + \frac{t_P}{p} + k(n, t)$$



Parallel overhead



A simple, quick-and-dirty way to measure your parallel overhead is to consider the “distance” between your code’s scaling and the perfect scaling.

$$\begin{aligned} T_s(n) - T_p(n, p) &= \\ t_s + t_p - t_s - \frac{t_p}{p} - k(n, p) &= \\ t_p \times \frac{p - 1}{p} - k(n, p) &= \end{aligned}$$

which becomes for ”large” p
 $\approx t_p - k(n, p)$



The Jargon behind HPC



- CPU = Socket = processor
- A CPU contains multi cores; “many” cores are multi-cores with large number of cores.
- sometimes (for instance on some software tools that explore the machine topology), the hardware cores are referred as “cpus”
- NUMA nodes (in tools like numactl, ...) = sockets
- MPI processes are also called “tasks”, whereas OpenMP threads are always threads. However, a “task” is also a concept in functional decomposition, as discussed in the OpenMP part that introduces the, in fact, tasks.

that's all, have fun

"So long
and thanks
for all the fish"