

Traditional Use of Parallel Computing: Large-Scale HPC Applications

- **High Performance Computing (HPC)**
 - Much computational work (in FLOPs, floatingpoint operations)
 - Often, large data sets
 - E.g. climate simulations, particle physics, engineering, sequence matching or proteome docking in bioinformatics, ...
- Single-CPU computers and even today's multicore processors cannot provide such massive computation power
- Aggregate LOTS of computers → **Clusters**
 - Need scalable parallel algorithms
 - Need exploit multiple levels of parallelism



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More Recent Use of Parallel Computing: Big-Data Analytics Applications

- **Big Data Analytics**
 - Data access intensive (disk I/O, memory accesses)
 - ▶ Typically, very large data sets (GB ... TB ... PB ... EB ...)
 - Also some computational work for combining/aggregating data
 - E.g. data center applications, business analytics, click stream analysis, scientific data analysis, machine learning, ...
 - Soft real-time requirements on interactive queries
- Single-CPU and multicore processors cannot provide such massive computation power and I/O bandwidth+capacity
- Aggregate LOTS of computers → **Clusters**
 - Need scalable parallel algorithms
 - Need exploit multiple levels of parallelism
 - Fault tolerance



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HPC vs Big-Data Computing

- Both need **parallel computing**
- **Same kind of hardware** – Clusters of (multicore) servers
- Same OS family (Linux)
- **Different programming models**, languages, and tools

HPC application	Big-Data application
HPC prog. languages: Fortran, C/C++ (Python)	Big-Data prog. languages: Java, Scala, Python, ...
Par. programming models: MPI, OpenMP, ...	Par. programming models: MapReduce, Spark, ...
Scientific computing libraries: BLAS, ...	Big-data storage/access: HDFS, ...
OS: Linux	OS: Linux
HW: Cluster	HW: Cluster

→ Let us start with the common basis: Parallel computer architecture

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Parallel Computer

A **parallel computer** is a computer consisting of

- + two or more **processors** that can cooperate and communicate to solve a **large** problem faster,
- + one or more **memory modules**,
- + an **interconnection network** that connects processors with each other and/or with the memory modules.

Multiprocessor: tightly connected processors, e.g. shared memory

Multicomputer: more loosely connected, e.g. distributed memory

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Parallel Computer Architecture Concepts

Classification of parallel computer architectures:

- by control structure
 - SISD, SIMD, MIMD
- by memory organization
 - in particular, Distributed memory vs. Shared memory
- by interconnection network topology

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Classification by Control Structure [Flynn'72]

SISD single instruction stream, single data stream
+ sequential. OK where performance is not an issue.

SIMD single instruction stream, multiple data streams
Common clock, common program memory, common program counter.
+ VLIW processors
+ traditional vector processors
+ traditional array computers
+ SIMD instructions on wide data words (e.g. AltiVec, SSE, ...)

MIMD multiple instruction streams, multiple data streams
Each processor has its own program counter.

Hybrid forms

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Classification by Memory Organization

Distributed memory system
e.g. (traditional) HPC cluster

Shared memory system
e.g. multiprocessor (SMP) or computer with a standard multicore CPU

Most common today in HPC and Data centers:

Hybrid Memory System

- Cluster (distributed memory) of hundreds, thousands of shared-memory servers each containing one or several multi-core CPUs

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Hybrid (Distributed + Shared) Memory

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Interconnection Networks (1)

■ **Network**
= physical interconnection medium (wires, switches)
+ communication protocol

(a) connecting cluster nodes with each other (DMS)
(b) connecting processors with memory modules (SMS)

Classification

■ **Direct / static interconnection networks**

- connecting nodes directly to each other
- Hardware routers (communication coprocessors) can be used to offload processors from most communication work

■ **Switched / dynamic interconnection networks**

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Interconnection Networks (2): Simple Topologies

bus 1 wire – bus saturation with many processors e.g. Ethernet

linear array **ring** e.g. Token Ring

2D grid **torus**

3D grid **3D torus**

tree root processor is bottleneck

fully connected

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Interconnection Networks (3): Fat-Tree Network

■ **Tree network** extended for higher bandwidth (more switches, more links) closer to the root

- avoids bandwidth bottleneck

Example: Infiniband network (www.mellanox.com)

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More about Interconnection Networks

- Hypercube, Crossbar, Butterfly, Hybrid networks... → TDDC78
- Switching and routing algorithms
- Discussion of interconnection network properties
 - Cost (#switches, #lines)
 - Scalability (asymptotically, cost grows not much faster than #nodes)
 - Node degree
 - Longest path (→ latency)
 - Accumulated bandwidth
 - Fault tolerance (worst-case impact of node or switch failure)

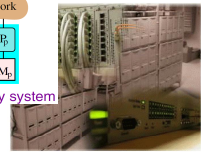
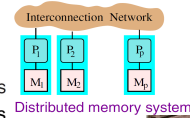
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Example: Beowulf-class PC Clusters

Characteristics:

- off-the-shelf (PC) nodes with off-the-shelf CPUs (Xeon, Opteron, ...)
- commodity interconnect G-Ethernet, Myrinet, Infiniband, SCI
- Open Source Unix Linux, BSD
- Message passing computing MPI, PVM



Advantages:

- + best price-performance ratio
- + low entry-level cost
- + vendor independent
- + scalable
- + rapid technology tracking

T. Sterling: The scientific workstation of the future may be a pile of PCs.

Communications of the ACM 39(9), Sep. 1996

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Cluster Example: Triolith (NSC, 2012 / 2013)

A so-called **Capability cluster** (fast network for *parallel* applications, not for just lots of independent sequential jobs)

1200 HP SL230 servers (compute nodes), each equipped with 2 Intel E5-2660 (2.2 GHz Sandybridge) processors with 8 cores each

→ 19200 cores in total

→ Theoretical peak performance of 338 Tflops/s

Mellanox Infiniband network (Fat-tree topology)



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NSC Triolith

The Challenge

■ Today, basically *all* computers are parallel computers!

- Single-thread performance stagnating
- Dozens of cores and hundreds of HW threads available per server
- May even be heterogeneous (core types, accelerators)
- Data locality matters
- Large clusters for HPC and Data centers, require message passing
- Utilizing more than one CPU core requires thread-level parallelism
- One of the biggest *software* challenges: **Exploiting parallelism**
 - Need LOTS of (mostly, independent) tasks to keep cores/HW threads busy and overlap waiting times (cache misses, I/O accesses)
 - All application areas, not only traditional HPC
 - ▶ General-purpose, data mining, graphics, games, embedded, DSP, ...
 - Affects HW/SW system architecture, programming languages, algorithms, data structures ...
 - Parallel programming is more error-prone (deadlocks, races, further sources of inefficiencies)
 - ▶ And thus more expensive and time-consuming

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Can't the compiler fix it for us?

■ Automatic parallelization?

- at compile time:
 - ▶ Requires static analysis – not effective for pointer-based languages
 - inherently limited – missing runtime information
 - ▶ needs programmer hints / rewriting ...
 - ▶ ok only for few benign special cases:
 - loop vectorization
 - extraction of instruction-level parallelism
- at run time (e.g. speculative multithreading)
 - ▶ High overheads, not scalable

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Insight

- Design of efficient / scalable parallel algorithms is, *in general*, a creative task that is not automatizable
- But some good recipes exist ...
 - Parallel algorithmic design patterns →

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The remaining solution ...

■ Manual parallelization!

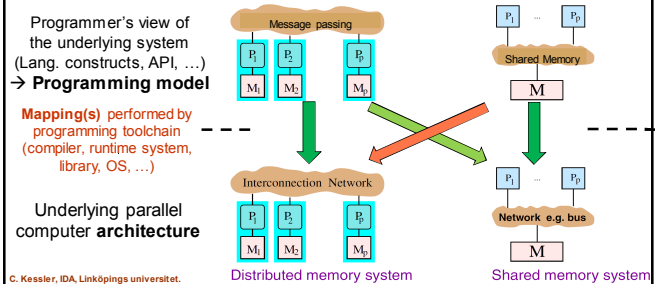
- using a parallel programming language / framework,
 - ▶ e.g. MPI message passing interface for distributed memory;
 - ▶ Pthreads, OpenMP, TBB, ... for shared-memory
- Generally harder, more error-prone than sequential programming,
 - ▶ requires special programming expertise to exploit the HW resources effectively
- Promising approach:
 - **Domain-specific languages/frameworks,**
 - ▶ Restricted set of predefined constructs doing most of the low-level stuff under the hood
 - ▶ e.g. MapReduce, Spark, ... for big-data computing

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Parallel Programming Model

- System-software-enabled **programmer's view** of the underlying hardware
 - **Abstracts** from details of the underlying architecture, e.g. network topology
 - Focuses on a **few characteristic properties**, e.g. memory model
- **Portability** of algorithms/programs across a family of parallel architectures



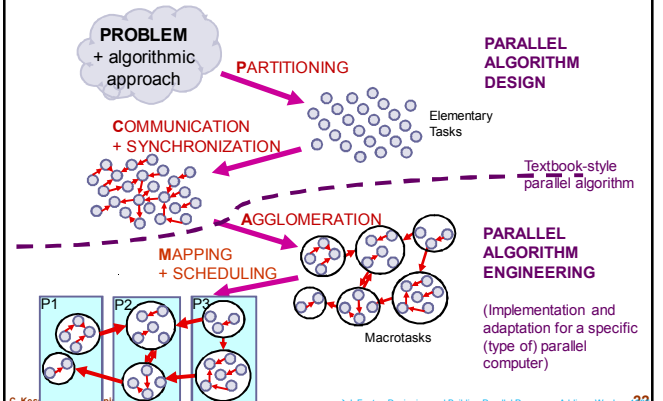
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Design and Analysis of Parallel Algorithms

Introduction

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Foster's Generic Method for the Design of Parallel Programs ("PCAM")



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→ 1. Foster, Designing and Building Parallel Programs, Addison-Wesley, 2002

Parallel Computation Model = Programming Model + Cost Model

- + abstract from hardware and technology
 - + specify basic operations, when applicable
 - + specify how data can be stored
- analyze algorithms **before** implementation $\rightarrow T = f(n, p, \dots)$
independent of a particular parallel computer
- focus on **most characteristic** (w.r.t. influence on exec. time) features of a broader class of parallel machines

Programming model

- shared memory / message passing,
- degree of synchronous execution

Cost model

- key parameters
- cost functions for basic operations
- constraints

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Parallel Cost Models

A Quantitative Basis for the Design of Parallel Algorithms

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Cost Model

Cost model: should

- + explain available observations
- + predict future behaviour
- + abstract from unimportant details → generalization

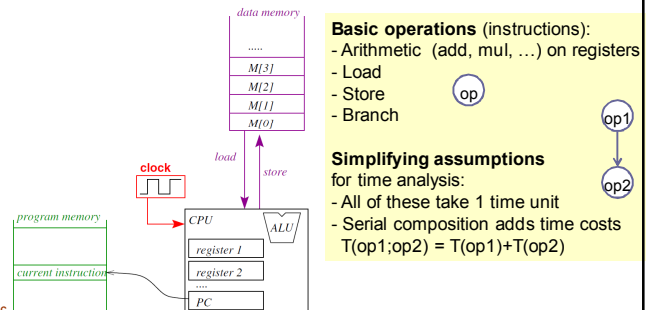
Simplifications to reduce model complexity:

- use idealized multicomputer model
ignore hardware details: memory hierarchies, network topology, ...
- use scale analysis
drop insignificant effects
- use empirical studies
calibrate simple models with empirical data
rather than developing more complex models

How to analyze sequential algorithms: The RAM (von Neumann) model for sequential computing

RAM (Random Access Machine)

programming and cost model for the analysis of sequential algorithms



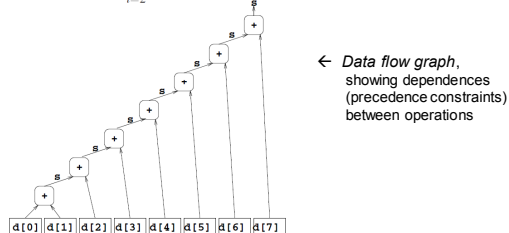
Analysis of sequential algorithms: RAM model (Random Access Machine)

Algorithm analysis: Counting instructions

Example: Computing the global sum of N elements

```
s = d[0]
for (i=1; i<N; i++)
    s = s + d[i]
```

$$t = t_{load} + t_{store} + \sum_{i=2}^N (2t_{load} + t_{add} + t_{store} + t_{branch}) = 5N - 3 \in \Theta(N)$$



→ arithmetic circuit model, directed acyclic graph (DAG) model

The PRAM Model – a Parallel RAM

Parallel Random Access Machine

[Fortune/Wyllie'78]

p processors

MIMD

common clock signal

arithm./jump: 1 clock cycle

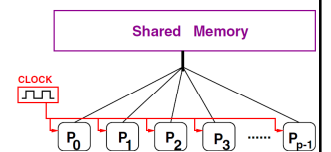
shared memory

uniform memory access time

latency: 1 clock cycle (!)

concurrent memory accesses

sequential consistency



PRAM variants → TDD56, TDDC78

Remark

PRAM model is very idealized, extremely simplifying / abstracting from real parallel architectures:

unbounded number of processors:

abstracts from scheduling overhead

local operations cost 1 unit of time

every processor has unit time memory access

to any shared memory location:

abstracts from communication time, bandwidth limitation, memory latency, memory hierarchy, and locality

→ focus on pure, fine-grained parallelism

→ Good for **early analysis** of parallel algorithm designs:

A parallel algorithm that does not scale under the PRAM model does not scale well anywhere else!

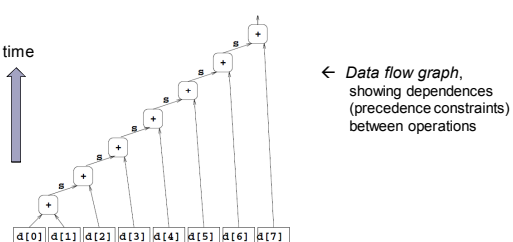
The PRAM cost model has only 1 machine-specific parameter: the number of processors

A first parallel sum algorithm ...

Keep the sequential sum algorithm's structure / data flow graph.

Giving each processor one task (load, add) does not help much

- All n loads could be done in parallel, but
- Processor i needs to wait for partial result from processor $i-1$, for $i=1, \dots, n-1$



→ Still $O(n)$ time steps!

Divide&Conquer Parallel Sum Algorithm in the PRAM / Circuit (DAG) cost model

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Given n numbers x_0, x_1, \dots, x_{n-1} stored in an array.

The global sum $\sum_{i=0}^{n-1} x_i$ can be computed in $\lceil \log_2 n \rceil$ time steps on an EREW PRAM with n processors.

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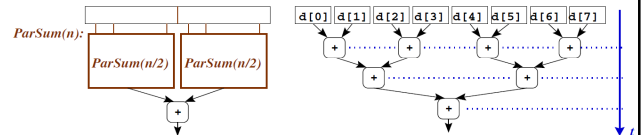
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Parallel algorithmic paradigm used: **Parallel Divide-and-Conquer**



Divide phase: trivial, time $O(1)$

Recursive calls: parallel time $T(n/2)$

with base case: load operation, time $O(1)$

Combine phase: addition, time $O(1)$

$$\begin{cases} T(n) = T(n/2) + O(1) \\ T(1) = O(1) \end{cases}$$

Use induction or the master theorem [Cormen+'90 Ch.4] $\rightarrow T(n) \in O(\log n)$

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Recursive formulation of DC parallel sum algorithm in some programming model

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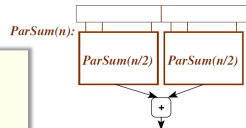
Implementation e.g. in Cilk: (shared memory)

```
cilk int parsum ( int *d, int from, int to )
{
    int mid, sumleft, sumright;
    if (from == to) return d[from]; // base case
    else {
        mid = (from + to) / 2;
        sumleft = spawn parsum ( d, from, mid );
        sumright = parsum ( d, mid+1, to );
        sync;
        return sumleft + sumright;
    }
}
```

// The main program:

```
main()
{
    ...
    parsum ( data, 0, n-1 );
    ...
}
```

Fork-Join execution style:
single task starts,
tasks spawn child tasks for
independent subtasks, and
synchronize with them



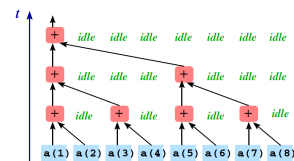
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Circuit / DAG model

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Independent of how the parallel computation is expressed, the resulting (unfolded) task graph looks the same.



■ **Task graph** is a directed acyclic graph (DAG) $G=(V,E)$

• Set V of vertices: elementary tasks (taking time 1 resp. $O(1)$ each)

• Set E of directed edges: dependencies (partial order on tasks)

• $(v_1, v_2) \in E \rightarrow v_1$ must be finished before v_2 can start

■ **Critical path** = longest path from an entry to an exit node

• Length of critical path is a lower bound for parallel time complexity

■ **Parallel time** can be longer if number of processors is limited

\rightarrow **schedule tasks** to processors such that dependences are preserved

(by programmer (SPMD execution) or run-time system (fork-join exec.))

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For a fixed number of processors ... ?

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- Usually, $p \ll n$
- Requires scheduling the work to p processors

(A) manually, at algorithm design time:

- Requires **algorithm engineering**
 - E.g. stop the parallel divide-and-conquer e.g. at subproblem size n/p and switch to sequential divide-and-conquer (= task agglomeration)
- For parallel sum:
- Step 0. Partition the array of n elements in p slices of n/p elements each (= domain decomposition)
 - Step 1. Each processor calculates a local sum for one slice, using the sequential sum algorithm, resulting in p partial sums (intermediate values)
 - Step 2. The p processors run the parallel algorithm to sum up the intermediate values to the global sum.

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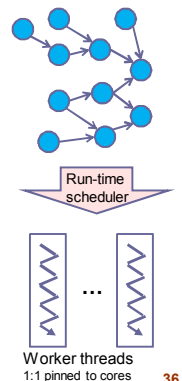
For a fixed number of processors ... ?

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- Usually, $p \ll n$
- Requires scheduling the work to p processors

(B) automatically, at run time:

- Requires a **task-based runtime system** with dynamic scheduler
 - Each newly created task is dispatched at runtime to an available worker processor.
 - Load balancing (\rightarrow runtime overhead)
 - Central task queue where idle workers fetch next task to execute
 - Local task queues + Work stealing – idle workers steal a task from some other processor



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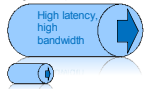
Analysis of Parallel Algorithms

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Analysis of Parallel Algorithms

Performance metrics of parallel programs

- **Parallel execution time**
 - Counted from the start time of the earliest task to the finishing time of the latest task
- **Work** – the total number of performed elementary operations
- **Cost** – the product of parallel execution time and #processors
- **Speed-up**
 - the factor by how much faster we can solve a problem with p processors than with 1 processor, usually in range $(0 \dots p)$
- **Parallel efficiency** = Speed-up / #processors, usually in $(0 \dots 1)$
- **Throughput** = #operations finished per second
- **Scalability**
 - does speedup keep growing well also when #processors grows large?



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Analysis of Parallel Algorithms

Asymptotic Analysis

- Estimation based on a cost model and algorithm idea (pseudocode operations)
- Discuss behavior for large problem sizes, large #processors

Empirical Analysis

- Implement in a concrete parallel programming language
- Measure time on a concrete parallel computer
 - Vary number of processors used, as far as possible
- More precise
- More work, and fixing bad designs at this stage is expensive

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Parallel Time, Work, Cost

problem size n

processors p

time $t(p, n)$

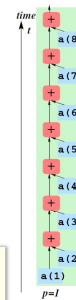
work $w(p, n)$

cost $c(p, n) = t \cdot p$

Example:
seq. sum algorithm

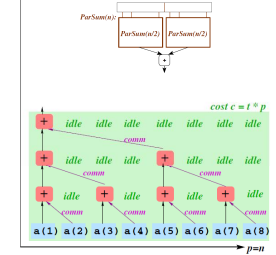
```
s = d[0]
for (i=1; i<N; i++)
    s = s + d[i]
```

$t(1, n) = t_{seq}(n) = O(n)$
 $w(1, n) = O(n)$
 $n-1$ additions
 n loads
 $O(n)$ other
 $c(1, n) = t(1, n) \cdot 1 = O(n)$



$p=1$

parallel sum algorithm



$t(n, n) = O(\log n)$

$w(n, n) = O(n)$

$c(n, n) = O(n \log n)$

par. sum alg. *not* cost-effective!

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Parallel work, time, cost

parallel work $w_A(n)$ of algorithm A on an input of size n
 = max. number of instructions performed by all procs during execution of A , where in each (parallel) time step as many processors are available as needed to execute the step in constant time.

parallel time $t_A(n)$ of algorithm A on input of size n
 = max. number of parallel time steps required under the same circumstances

parallel cost $c_A(n) = t_A(n) * p_A(n) \rightarrow c_A(n) \geq w_A(n)$
 where $p_A(n) = \max_i p_i(n) = \max.$ number of processors used in a step of A

Work, time, cost are thus *worst-case* measures.

$t_A(n)$ is sometimes called the **depth** of A
 (cf. **circuit model** of (parallel) computation)

$p_i(n)$ = number of processors needed in time step i , $0 \leq i < t_A(n)$, to execute the step in constant time. Then, $w_A(n) = \sum_{i=0}^{t_A(n)} p_i(n)$

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Speedup

Consider problem \mathcal{P} , parallel algorithm A for \mathcal{P}

T_s = time to execute the best serial algorithm for \mathcal{P}
 on one processor of the parallel machine

$T(1)$ = time to execute parallel algorithm A on 1 processor

$T(p)$ = time to execute parallel algorithm A on p processors

Absolute speedup $S_{abs} = \frac{T_s}{T(p)}$

Relative speedup $S_{rel} = \frac{T(1)}{T(p)}$

$S_{abs} \leq S_{rel}$

Speedup $S(p)$ with p processors is usually in the range $(0 \dots p)$

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Amdahl's Law: Upper bound on Speedup

Consider execution (trace) of parallel algorithm A:

sequential part A^s where only 1 processor is active

parallel part A^p that can be sped up perfectly by p processors

→ total work $w_A(n) = w_{A^s}(n) + w_{A^p}(n)$, time $T = T_{A^s} + \frac{T_{A^p}}{p}$,

Amdahl's Law

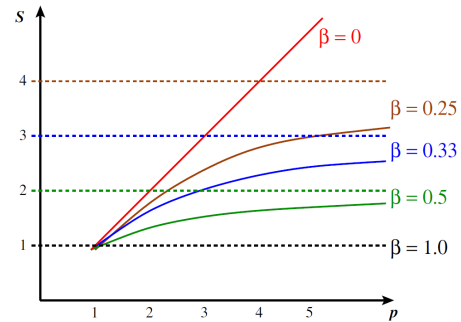
If the sequential part of A is a **fixed** fraction of the total work irrespective of the problem size n , that is, if there is a constant β with

$$\beta = \frac{w_{A^s}(n)}{w_A(n)} \leq 1$$

the relative speedup of A with p processors is limited by

$$\frac{p}{\beta p + (1 - \beta)} < 1/\beta$$

Amdahl's Law

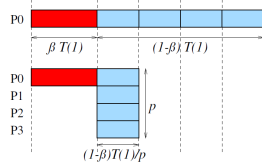


Proof of Amdahl's Law

$$S_{rel} = \frac{T(1)}{T(p)} = \frac{T(1)}{T_{A^s} + T_{A^p}(p)}$$

Assume perfect parallelizability of the parallel part A^p , that is, $T_{A^p}(p) = (1 - \beta)T(1)/p$:

$$S_{rel} = \frac{T(1)}{\beta T(1) + (1 - \beta)T(1)/p} = \frac{p}{\beta p + 1 - \beta} \leq 1/\beta$$



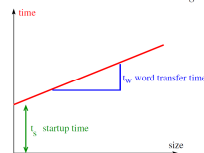
Towards More Realistic Cost Models

Modeling the cost of communication and data access

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Modeling Communication Cost: Delay Model

Idealized multicomputer: point-to-point communication costs overhead t_{msg} .



Cost of communicating a larger block of n bytes:

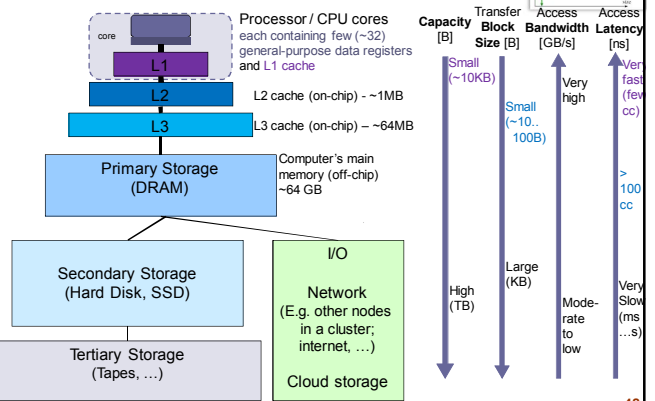
$$\text{time } t_{msg}(n) = \text{sender overhead} + \text{latency} + \text{receiver overhead} + n/\text{bandwidth} \\ =: t_{startup} + n \cdot t_{transfer}$$

Assumption: network not overloaded; no conflicts occur at routing

$t_{startup}$ = startup time (time to send a 0-byte message)
accounts for hardware and software overhead.

$t_{transfer}$ = transfer rate, send time per word sent.
depends on the network bandwidth.

Memory Hierarchy And The Real Cost of Data Access



Data Locality

- **Memory hierarchy rationale:** Try to amortize the high access cost of lower levels (DRAM, disk, ...) by caching data in higher levels for faster subsequent accesses
 - **Cache miss** – stall the computation. fetch the block of data containing the accessed address from next lower level, then resume
 - More reuse of cached data (**cache hits**) → better performance
- **Working set** = the set of memory addresses accessed together in a period of computation
- **Data locality** = property of a computation: keeping the working set small during a computation
 - **Temporal locality** – re-access same data element multiple times within a short time interval
 - **Spatial locality** – re-access neighbored memory addresses multiple times within a short time interval
- High latency favors larger transfer block sizes (cache lines, memory pages, file blocks, messages) for amortization over many subsequent accesses

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Memory-bound vs. CPU-bound computation

- **Arithmetic intensity** of a computation
= #arithmetic instructions (computational work) executed per accessed element of data in memory (after cache miss)
- A computation is **CPU-bound** if its arithmetic intensity is $\gg 1$.
 - The performance bottleneck is the CPU's arithmetic throughput
- A computation is **memory-access bound** otherwise.
 - The performance bottleneck is memory accesses, CPU is not fully utilized
- Examples:
 - Matrix-matrix-multiply (if properly implemented) is CPU-bound.
 - Array global sum is memory-bound on most architectures.

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Some Parallel Algorithmic Design Patterns

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Data Parallelism

Given:

- One (or several) data containers $\mathbf{x}, \mathbf{z}, \dots$ with n elements each, e.g. array(s) $\mathbf{x} = (x_1, \dots, x_n), \mathbf{z} = (z_1, \dots, z_n), \dots$
- An operation f on individual elements of $\mathbf{x}, \mathbf{z}, \dots$ (e.g. *incr*, *sqrt*, *mult*, ...)

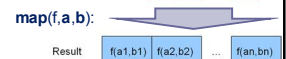
Compute: $\mathbf{y} = f(\mathbf{x}) = (f(x_1), \dots, f(x_n))$

Parallelizability: Each data element defines a task

- Fine grained parallelism
- Easily partitioned into independent tasks, fits very well on all parallel architectures

Notation with higher-order function:

- $\mathbf{y} = \text{map}(f, \mathbf{x})$



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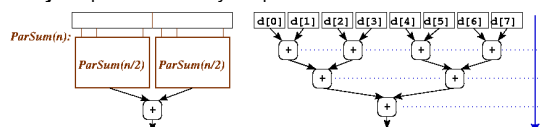
Data-parallel Reduction

Given:

- A data container \mathbf{x} with n elements, e.g. array $\mathbf{x} = (x_1, \dots, x_n)$
- A **binary, associative** operation op on individual elements of \mathbf{x} (e.g. *add*, *max*, *bitwise-or*, ...)

Compute: $\mathbf{y} = OP_{i=1 \dots n} \mathbf{x} = x_1 op x_2 op \dots op x_n$

Parallelizability: Exploit *associativity* of op



Notation with higher-order function:

- $\mathbf{y} = \text{reduce}(op, \mathbf{x})$

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MapReduce (pattern)

- A **Map** operation with operation f on one or several input data containers \mathbf{x}, \dots , producing a temporary output data container \mathbf{w} , directly followed by a **Reduce** with operation g on \mathbf{w} producing result \mathbf{y}

■ $\mathbf{y} = \text{MapReduce}(f, g, \mathbf{x}, \dots)$

■ Example:

Dot product of two vectors \mathbf{x}, \mathbf{z} : $\mathbf{y} = \sum_i x_i * z_i$

f = scalar multiplication,

g = scalar addition

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Task Farming

Independent subcomputations f_1, f_2, \dots, f_m could be done in parallel and/or in arbitrary order, e.g.

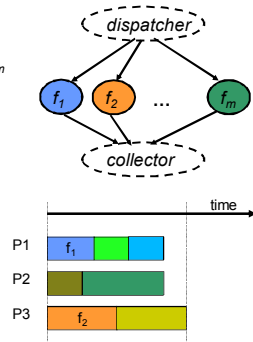
- independent loop iterations
- independent function calls

Scheduling (mapping) problem

- m tasks onto p processors
- static (before running) or dynamic
- **Load balancing** is important: most loaded processor determines the parallel execution time

Notation with higher-order function:

- **farm** (f_1, \dots, f_m) (x_1, \dots, x_n)



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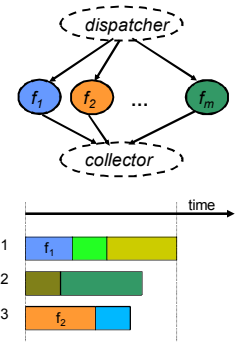
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Parallel Divide-and-Conquer

(Sequential) Divide-and-conquer:

- If given problem instance P is *trivial*, solve it *directly*. Otherwise:
- **Divide**: Decompose problem instance P in one or several smaller independent instances of the same problem, P_1, \dots, P_k
- For each i : solve P_i by recursion.
- **Combine** the solutions of the P_i into an overall solution for P

Parallel Divide-and-Conquer:

- Recursive calls can be done in parallel.
- Parallelize, if possible, also the divide and combine phase.
- Switch to sequential divide-and-conquer when enough parallel tasks have been created.

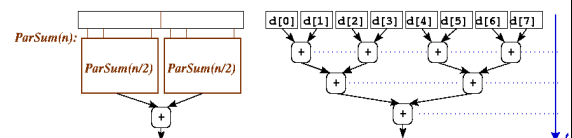
Notation with higher-order function:

- **solution** = **DC** (*divide, combine, istrivial, solvedirectly, n, P*)

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Example: Parallel Divide-and-Conquer



Example: Parallel Sum over integer-array x

Exploit associativity:

$$\text{Sum}(x_1, \dots, x_n) = \text{Sum}(x_1, \dots, x_{n/2}) + \text{Sum}(x_{n/2+1}, \dots, x_n)$$

Divide: trivial, split array x in place

Combine is just an addition.

$$y = \text{DC}(\text{split}, \text{add}, \text{nilsSmall}, \text{addFewInSeq}, n, x)$$

→ Data parallel reductions are an important special case of DC.

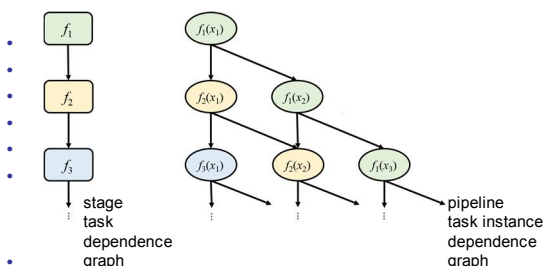
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Pipelining

applies a sequence of dependent computations/tasks (f_1, f_2, \dots, f_k) elementwise to data sequence $x = (x_1, x_2, x_3, \dots, x_n)$

- For fixed x_j , must compute $f_i(x_j)$ before $f_{i+1}(x_j)$
- ... and $f_i(x_j)$ before $f_i(x_{j+1})$ if the tasks f_i have a *run-time state*



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- ... and $f_i(x_j)$ before $f_i(x_{j+1})$ if the tasks f_i have a *run-time state*

Parallelizability: Overlap execution of all f_i for k subsequent x_j

- $\text{time}=1$: compute $f_1(x_1)$
- $\text{time}=2$: compute $f_1(x_2)$ and $f_2(x_1)$
- $\text{time}=3$: compute $f_1(x_3)$ and $f_2(x_2)$ and $f_3(x_1)$
- ...
- Total time: $O((n+k) \max_i(\text{time}(f_i)))$ with k processors
- Still, requires good mapping of the tasks f_i to the processors for even load balancing – often, static mapping (done before running)

Notation with higher-order function:

- $(y_1, \dots, y_n) = \text{pipe}((f_1, \dots, f_k), (x_1, \dots, x_n))$

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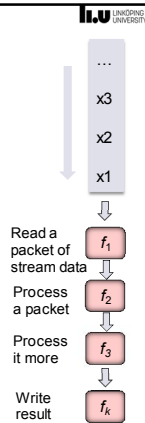
Streaming

- **Streaming** applies pipelining to processing of large (possibly, infinite) data *streams* from or to memory, network or devices, usually partitioned in fixed-sized data packets,

- in order to **overlap** the processing of each packet of data **in time** with access of *subsequent* units of data and/or processing of preceding packets of data.

Examples

- Video streaming from network to display
- Surveillance camera, face recognition
- Network data processing e.g. deep packet inspection



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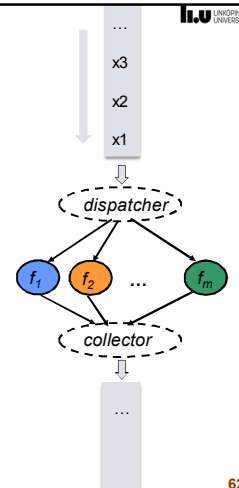
Stream Farming

Combining streaming and task farming patterns

Independent streaming subcomputations f_1, f_2, \dots, f_m on each data packet

Speed up the pipeline by parallel processing of subsequent data packets

In most cases, the original order of packets must be kept after processing



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(Algorithmic) Skeletons

Skeletons are reusable, parameterizable SW components with well defined semantics for which efficient parallel implementations may be available.

Inspired by higher-order functions in functional programming

One or very few skeletons per parallel algorithmic paradigm

- map, farm, DC, reduce, pipe, scan ...

Parameterised in user code

- Customization by instantiating a **skeleton template** in a **user-provided function**

Composition of skeleton instances in program code normally by sequencing+data flow

- e.g. `squaresum(x)` can be defined by

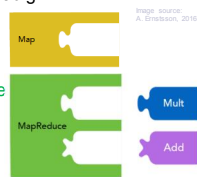
```

{
    tmp = map( sqr, x );
    return reduce( add, tmp );
}
    
```

For frequent combinations, may define advanced skeletons, e.g.:

```

{
    mapreduce( sqr, add, x )
}
    
```



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SkePU [Enmyren, K. 2010]

- Skeleton programming library for heterogeneous multicore systems, based on C++

- Example: Vector addition in SkePU-2 [Ernstsson 2016]

```

int add(int a, int b)
{
    return a + b;
}
    
```

```

auto vec_add = Map<2>(add);
    
```

```

vec_add(result, v1, v2);
    
```



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SkePU [Enmyren, K. 2010]

- Skeleton programming library for heterogeneous multicore systems, based on C++

- Example: Vector sum in SkePU-2 [Ernstsson 2016]

```

int add(int a, int b)
{
    return a + b;
}
    
```

```

auto vec_sum = Reduce(add);
    
```

```

vec_sum(result, v1, v2);
    
```



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High-Level Parallel Programming with Skeletons

- **Skeletons** (constructs) *implement* (parallel) **algorithmic design patterns**
 - ☺ Abstraction, hiding complexity (parallelism and low-level programming)
 - ☹ Enforces structuring, restricted set of constructs
 - ☺ Parallelization for free
 - ☺ Easier to analyze and transform
 - ☹ Requires complete understanding and rewriting of a computation
 - ☹ Available skeleton set does not always fit
 - ☹ May lose some efficiency compared to manual parallelization
- Idea developed in HPC (mostly in Europe) since the late 1980s.
- Many (esp., academic) frameworks exist, mostly as libraries
- Industry (also beyond HPC domain) has adopted skeletons
 - map, reduce, scan in many modern parallel programming APIs
 - e.g., Intel *Threading Building Blocks (TBB)*: par. for, par. reduce, pipe
 - NVIDIA *Thrust*
 - Google *MapReduce* (for distributed data mining applications)

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Further Reading

C. Kessler: **Design and Analysis of Parallel Algorithms – An Introduction**. Compendium for TDDC78 and TDDD56, Edition Spring 2019. PDF, 123 pages.
<http://www.ida.liu.se/~TDDC78/handouts> (login: parallel, password see whiteboard)
 • Chapter 2 on analysis of parallel algorithms as background reading

On PRAM model and Design and Analysis of Parallel Algorithms

- J. Keller, C. Kessler, J. Träff: **Practical PRAM Programming**. Wiley Interscience, New York, 2001.
- J. JaJa: **An introduction to parallel algorithms**. Addison-Wesley, 1992.
- D. Cormen, C. Leiserson, R. Rivest: **Introduction to Algorithms**, Chapter 30. MIT press, 1989, or a later edition.
- H. Jordan, G. Alaghband: **Fundamentals of Parallel Processing**. Prentice Hall, 2003.
- A. Grama, G. Karypis, V. Kumar, A. Gupta: **Introduction to Parallel Computing**, 2nd Edition. Addison-Wesley, 2003.

On skeleton programming, see e.g. our publications on SkePU:

- <http://www.ida.liu.se/labs/pelab/skepu>

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Questions for Reflection

- Model the overall cost of a streaming computation with a very large number N of input data elements on a *single* processor
 - (a) if implemented as a loop over the data elements running on an ordinary memory hierarchy with hardware caches (see above)
 - (b) if overlapping computation for a data packet with transfer/access of the next data packet
 - (b1) if the computation is CPU-bound
 - (b2) if the computation is memory-bound
- Which property of streaming computations makes it possible to overlap computation with data transfer?
- Can each dataparallel computation be streamed?
- What are the performance advantages and disadvantages of large vs. small packet sizes in streaming?
- Why should servers in datacenters running I/O-intensive tasks (such as disk/DB accesses) get many more tasks to run than they have cores?
- How would you extend the skeleton programming approach for computations that operate on secondary storage (file/DB accesses)?

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