3 Introduction

Parallel Computing

A parallel computer is a collection of processors usually of the same type, interconnected to allow coordination and exchange of data.

The processors are primarily used to jointly solve a given problem.

Distributed Systems

A set of possibly many different types of processors are distributed over a larger geographic area.

Processors do not work on a single problem.

Some processors may act in a malicous way.



Cost measures

How do we evaluate sequential algorithms?

- time efficiency
- space utilization
- energy consumption
- programmability

▶ ...

Asymptotic bounds (e.g., for running time) often give a good indication on the algorithms performance on a wide variety of machines.



Cost measures

How do we evaluate parallel algorithms?

- time efficiency
- space utilization
- energy consumption
- programmability
- communication requirement
- ▶ ...

Problems

- performance (e.g. runtime) depends on problem size n and on number of processors p
- statements usually only hold for restricted types of parallel machine as parallel computers may have vastly different characteristics (in particular w.r.t. communication)

Speedup

Suppose a problem *P* has sequential complexity $T^*(n)$, i.e., there is no algorithm that solves *P* in time $o(T^*(n))$.

Definition 1

The speedup $S_p(n)$ of a parallel algorithm A that requires time $T_p(n)$ for solving P with p processors is defined as

$$S_p(n) = \frac{T^*(n)}{T_p(n)}$$

Clearly, $S_p(n) \le p$. Goal: obtain $S_p(n) \approx p$.

It is common to replace $T^*(n)$ by the time bound of the best **known** sequential algorithm for P!



Efficiency

Definition 2

The efficiency of a parallel algorithm A that requires time $T_p(n)$ when using p processors on a problem of size n is

$$E_p(n) = \frac{T_1(n)}{pT_p(n)}$$

 $E_p(n) \approx 1$ indicates that the algorithm is running roughly p times faster with p processors than with one processor.

Note that $E_p(n) \leq \frac{T_1(n)}{pT_{\infty}(n)}$. Hence, the efficiency goes down rapidly if $p \geq T_1(n)/T_{\infty}(n)$.

Disadvantage: cost-measure does not relate to the optimum sequential algorithm.

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Parallel Models — Requirements

Simplicity

A model should allow to easily analyze various performance measures (speed, communication, memory utilization etc.).

Results should be as hardware-independent as possible.

Implementability

Parallel algorithms developed in a model should be easily implementable on a parallel machine.

Theoretical analysis should carry over and give meaningful performance estimates.

A real satisfactory model does not exist!



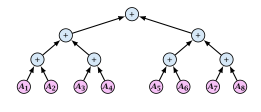
DAG model — computation graph

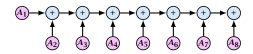
- nodes represent operations (single instructions or larger blocks)
- edges represent dependencies (precedence constraints)
- closely related to circuits; however there exist many different variants
- branching instructions cannot be modelled
- completely hardware independent
- scheduling is not defined

Often used for automatically parallelizing numerical computations.



Example: Addition





Here, vertices without incoming edges correspond to input data. The graph can be viewed as a data flow graph.



DAG model — computation graph

The DAG itself is not a complete algorithm. A scheduling implements the algorithm on a parallel machine, by assigning a time-step t_v and a processor p_v to every node.

Definition 3

A scheduling of a DAG G = (V, E) on p processors is an assignment of pairs (t_v, p_v) to every internal node $v \in V$, s.t.,

• $p_{v} \in \{1, \dots, p\}; t_{v} \in \{1, \dots, T\}$

•
$$t_u = t_v \Rightarrow p_u \neq p_v$$

•
$$(u, v) \in E \Rightarrow t_v \ge t_u + 1$$

where a non-internal node x (an input node) has $t_x = 0$. T is the length of the schedule.



DAG model — computation graph

The parallel complexity of a DAG is defined as

 $T_p(n) = \min_{\text{schedule } S} \{T(S)\}$.

 $T_1(n)$: #internal nodes in DAG $T_{\infty}(n)$: diameter of DAG

Clearly,

 $T_p(n) \ge T_{\infty}(n)$ $T_p(n) \ge T_1(n)/p$

Lemma 4

A schedule with length $O(T_1(n)/p + T_{\infty}(n))$ can be found easily.

Lemma 5

Finding an optimal schedule is in general NP-complete.

Note that the DAG model as defined is a non-uniform model of computation.

In principle, there could be a different DAG for every input size n.

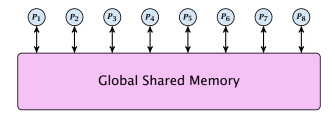
An algorithm (e.g. for a RAM) must work for every input size and must be of finite description length.

Hence, specifying a different DAG for every n has more expressive power.

Also, this is not really a complete model, as the operations allowed in a DAG node are not clearly defined.



PRAM Model



All processors are synchronized.

In every round a processor can:

- read a register from global memory into local memory
- do a local computation à la RAM
- write a local register into global memory



PRAM Model

Every processor executes the same program.

However, the program has access to two special variables:

- p: total number of processors
- $id \in \{1, \dots, p\}$: the id of the current processor

The following (stupid) program copies the content of the global register x[1] to registers $x[2] \dots x[p]$.

Algorithm 1 copy							
1:	if $id = 1$ then $round \leftarrow 1$						
2: while round $\leq p$ and $id = round$ do							
3:	$x[id+1] \leftarrow x[id]$						
4:	$round \leftarrow round + 1$						



PRAM Model

- processors can effectively execute different code because of branching according to *id*
- however, not arbitrarily; still uniform model of computation

Often it is easier to explicitly define which parts of a program are executed in parallel:

```
Algorithm 2 sum
1: // computes sum of x[1] \dots x[p]
2: // red part is executed only by processor 1
3: \gamma \leftarrow 1
4: while 2^{\gamma} \leq p do
5: for id mod 2^r = 1 pardo
6: // only executed by processors whose id matches
7:
             x[id] = x[id] + x[id + 2^{r-1}]
   \gamma \leftarrow \gamma + 1
 8:
 9: return x[1]
```

Different Types of PRAMs

Simultaneous Access to Shared Memory:

EREW PRAM:

simultaneous access is not allowed

CREW PRAM:

concurrent read accesses to the same location are allowed; write accesses have to be exclusive

CRCW PRAM:

concurrent read and write accesses allowed

- commom CRCW PRAM all processors writing to x[i] must write same value
- arbitrary CRCW PRAM values may be different; an arbitrary processor succeeds
- priority CRCW PRAM values may be different; processor with smallest id succeeds



Algorithm 3 sum

```
1: // computes sum of x[1]...x[p]

2: r \leftarrow 1

3: while 2^r \le p do

4: for id mod 2^r = 1 pardo

5: x[id] = x[id] + x[id + 2^{r-1}]

6: r \leftarrow r + 1

7: return x[1]
```

The above is an EREW PRAM algorithm.

On a CREW PRAM we could replace Line 4 by for $1 \le id \le p$ pardo



PRAM Model — remarks

- similar to a RAM we either need to restrict the size of values that can be stored in registers, or we need to have a non-uniform cost model for doing a register manipulation (cost for manipulating x[i] is proportional to the bit-length of the largest number that is ever being stored in x[i])
 - in this lecture: uniform cost model but we are not exploiting the model
- global shared memory is very unrealistic in practise as uniform access to all memory locations does not exist
- global synchronziation is very unrealistic; in real parallel machines a global synchronization is very costly
- model is good for understanding basic parallel mechanisms/techniques but not for algorithm development
- model is good for lower bounds

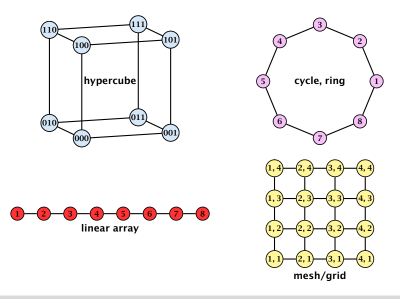


Network of Workstations — NOWs

- interconnection network represented by a graph G = (V, E)
- each $v \in V$ represents a processor
- ► an edge {u, v} ∈ E represents a two-way communication link between processors u and v
- network is asynchronous
- all coordination/communiation has to be done by explicit message passing



Typical Topologies





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Network of Workstations — NOWs

Computing the sum on a *d*-dimensional hypercube. Note that $x[0] \dots x[2^d - 1]$ are stored at the individual nodes.

Processors are numbered consecutively starting from 0

```
Algorithm 4 sum
1: // computes sum of x[0] \dots x[2^d - 1]
2: r \leftarrow 1
3: while 2^r \le 2^d \text{ do } // p = 2^d
   if id mod 2^r = 0 then
4:
              temp \leftarrow receive(id + 2<sup>r-1</sup>)
5:
             x[id] = x[id] + temp
6:
7: if id mod 2^r = 2^{r-1} then
8:
             send(x[id], id -2^{r-1})
9: r \leftarrow r + 1
10: if id = 0 then return x[id]
```

Network of Workstations — NOWs

Remarks

- One has to ensure that at any point in time there is at most one active communication along a link
- There also exist synchronized versions of the model, where in every round each link can be used once for communication
- In particular the asynchronous model is quite realistic
- Difficult to develop and analyze algorithms as a lot of low level communication has to be dealt with
- Results only hold for one specific topology and cannot be generalized easily



Performance of PRAM algorithms

Suppose that we can solve an instance of a problem with size n with P(n) processors and time T(n).

We call $C(n) = T(n) \cdot P(n)$ the time-processor product or the cost of the algorithm.

The following statements are equivalent

- P(n) processors and time $\mathcal{O}(T(n))$
- $\mathcal{O}(\mathcal{C}(n))$ cost and time $\mathcal{O}(T(n))$
- $\mathcal{O}(\mathcal{C}(n)/p)$ time for any number $p \leq P(n)$ processors
- $\mathcal{O}(C(n)/p + T(n))$ for any number p of processors



Performance of PRAM algorithms

Suppose we have a PRAM algorithm that takes time T(n) and work W(n), where work is the total number of operations.

We can nearly always obtain a PRAM algorithm that uses time at most

$$W(n)/p \rfloor + T(n)$$

parallel steps on p processors.

Idea:

- ▶ $W_i(n)$ denotes operations in parallel step *i*, $1 \le i \le T(n)$
- simulate each step in $[W_i(n)/p]$ parallel steps
- then we have

$$\sum_{i} \lceil W_{i}(n)/p \rceil \leq \sum_{i} \left(\lfloor W_{i}(n)/p \rfloor + 1 \right) \leq \lfloor W(n)/p \rfloor + T(n)$$

Performance of PRAM algorithms

Why nearly always?

We need to assign processors to operations.

- every processor p_i needs to know whether it should be active
- in case it is active it needs to know which operations to perform

design algorithms for an arbitrary number of processors; keep total time and work low



Optimal PRAM algorithms

Suppose the optimal sequential running time for a problem is $T^*(n)$.

We call a PRAM algorithm for the same problem work optimal if its work W(n) fulfills

$$W(n) = \Theta(T^*(n))$$

If such an algorithm has running time T(n) it has speedup

$$S_p(n) = \Omega\left(\frac{T^*(n)}{T^*(n)/p + T(n)}\right) = \Omega\left(\frac{pT^*(n)}{T^*(n) + pT(n)}\right) = \Omega(p)$$

for $p = \mathcal{O}(T^*(n)/T(n)).$



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This means by improving the time T(n), (while using same work) we improve the range of p, for which we obtain optimal speedup.

We call an algorithm worktime (WT) optimal if T(n) cannot be asymptotically improved by any work optimal algorithm.



Example

Algorithm for computing the sum has work W(n) = O(n). optimal

 $T(n) = O(\log n)$. Hence, we achieve an optimal speedup for $p = O(n/\log n)$.

One can show that any CREW PRAM requires $\Omega(\log n)$ time to compute the sum.



When we differentiate between local and global memory we can analyze communication cost.

We define the communication cost of a PRAM algorithm as the worst-case traffic between the local memory of a processor and the global shared memory.

Important criterion as communication is usually a major bottleneck.



Communication Cost

```
Algorithm 5 MatrixMult(A, B, n)1: Input: n \times n matrix A and B; n = 2^k2: Output: C = AB3: for 1 \le i, j, \ell \le n pardo4: X[i, j, \ell] \leftarrow A[i, \ell] \cdot B[\ell, j]5: for r \leftarrow 1 to log n6: for 1 \le i, j \le n; \ell \mod 2^r = 1 pardo7: X[i, j, \ell] \leftarrow X[i, j, \ell] + X[i, j, \ell + 2^{r-1}]8: C[i, j] \leftarrow X[i, j, \ell]
```

On n^3 processors this algorithm runs in time $O(\log n)$. It uses n^3 multiplications and $O(n^3)$ additions.



What happens if we have n processors?

Phase 1

 p_i computes $X[i, j, \ell] = A[i, \ell] \cdot B[\ell, j]$ for all $1 \le j, \ell \le n$ n^2 time; n^2 communication for every processor

Phase 2 (round r)

 p_i updates $X[i, j, \ell]$ for all $1 \le j \le n; 1 \le \ell \mod 2^r = 1$ $n \cdot n/2^r$ time; no communication

Phase 3

 p_i writes *i*-th row into C[i, j]'s.

n time; n communication



Alternative Algorithm

Split matrix into blocks of size $n^{2/3} \times n^{2/3}$.

$A_{1,1}$ $A_{1,2}$ $A_{1,3}$ $A_{1,4}$		<i>B</i> _{1,1}	B _{1,2}	B _{1,3}	<i>B</i> _{1,4}		<i>C</i> _{1,1}	<i>C</i> _{1,2}	<i>C</i> _{1,3}	<i>C</i> _{1,4}
$A_{2,1} A_{2,2} A_{2,3} A_{2,4}$		B _{2,1}	B _{2,2}	B _{2,3}	<i>B</i> _{2,4}	_	<i>C</i> _{2,1}	C _{2,2}	<i>C</i> _{2,3}	<i>C</i> _{2,4}
$\begin{array}{c c} & \mathbf{A} \\ A_{3,1} & A_{3,2} & A_{3,3} & A_{3,4} \end{array}$	-	B _{3,1}	B _{3,2}	B _{3,3}	B _{3,4}	_	<i>C</i> _{3,1}	C _{3,2}	C _{3,3}	<i>C</i> _{3,4}
A _{4,1} A _{4,2} A _{4,3} A _{4,4}		B4,1	B4,2	B4,3	B4,4		<i>C</i> _{4,1}	C _{4,2}	<i>C</i> _{4,3}	C _{4,4}

Note that $C_{i,j} = \sum_{\ell} A_{i,\ell} B_{\ell,j}$.

Now we have the same problem as before but $n' = n^{1/3}$ and a single multiplication costs time $\mathcal{O}((n^{2/3})^3) = \mathcal{O}(n^2)$. An addition costs $n^{4/3}$.

work for multiplications: $\mathcal{O}(n^2 \cdot (n')^3) = \mathcal{O}(n^3)$ work for additions: $\mathcal{O}(n^{4/3} \cdot (n')^3) = \mathcal{O}(n^3)$ time: $\mathcal{O}(n^2) + \log n' \cdot \mathcal{O}(n^{4/3}) = \mathcal{O}(n^2)$

Alternative Algorithm

The communication cost is only $\mathcal{O}(n^{4/3}\log n')$ as a processor in the original problem touches at most $\log n$ entries of the matrix.

Each entry has size $O(n^{4/3})$.

The algorithm exhibits less parallelism but still has optimum work/runtime for just n processors.

much, much better in practise

