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.

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- space utilization
- energy consumption
- programmability
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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 $S_{\mathcal{P}}(n)$ of a parallel algorithm A that requires time $T_{\mathcal{P}}(n)$ for solving P with \mathcal{P} processors is defined as

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

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



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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 $p \geq T_1(n)/T_{\infty}(n)$.



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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.



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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.



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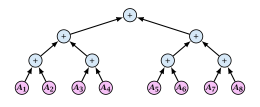
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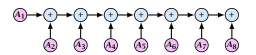


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Example: Addition





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

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

where a non-internal node x (an input node) has $t_{\scriptscriptstyle X}=0.$ T is the length of the schedule.



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The parallel complexity of a DAG is defined as

$$T_p(n) = \min_{\text{schedule } S} \{T(S)\}$$
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 $T_1(n)$: #internal nodes in DAG

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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.

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

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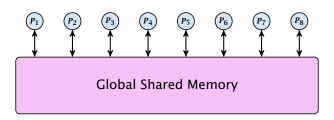


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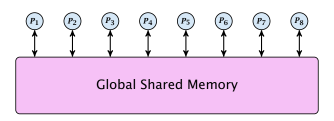




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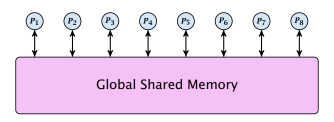




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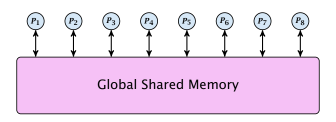




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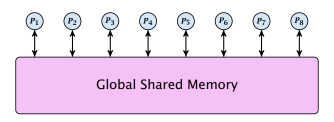




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Every processor executes the same program.

However, the program has access to two special variables:

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

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Algorithm 1 copy

1: if id = 1 then round - 1

2: while round \le p and id = round do

3: x[id + 1] - x[id]

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Algorithm 2 sum
1: // computes sum of x[1]...x[p]
2: // red part is executed only by processor 1
3: \gamma \leftarrow 1
4: while 2^r \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}]
   r \leftarrow r + 1
9: return x[1]
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- 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





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Algorithm 3 sum

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1: // computes sum of x[1]...x[p]
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- 2: $r \leftarrow 1$
- 3: while $2^r \le p$ do
- 4: **for** $id \mod 2^{\gamma} = 1$ **pardo**
- 5: $x[id] = x[id] + x[id + 2^{r-1}]$
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- 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



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- 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 synchronization 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



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- ightharpoonup each $v \in V$ represents a processor
- ▶ an edge $\{u, v\} \in E$ represents a two-way communication link between processors u and v
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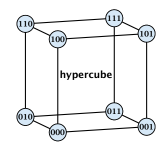
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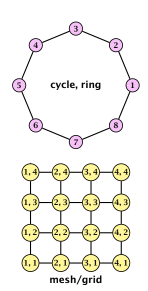
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Typical Topologies









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

Processors are numbered consecutively starting from 0

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```
Algorithm 4 sum
1: // computes sum of x[0]...x[2^d-1]
2: r \leftarrow 1
3: while 2^r \le 2^d do //p = 2^d
   if id mod 2^r = 0 then
4:
             temp \leftarrow receive(id + 2^{r-1})
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]
```

- 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



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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.

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



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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

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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 $\{W_i(n)\}_{\mathcal{D}}$ parallel steps

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 $\sum_{i} |W_{i}(n)/p| \le \sum_{i} (|W_{i}(n)/p| + 1) \le |W(n)/p| + T(n)$

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- then we have

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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.

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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.



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Algorithm 5 MatrixMult(A, B, n)

- 1: **Input:** $n \times n$ matrix A and B; $n = 2^k$
- 2: Output: C = AB
- 3: for $1 \le i, j, \ell \le n$ pardo
- 4: $X[i,j,\ell] \leftarrow A[i,\ell] \cdot B[\ell,j]$
- 5: **for** $r \leftarrow 1$ **to** $\log n$
- 6: **for** $1 \le i, j \le n$; $\ell \mod 2^r = 1$ **pardo**
- 7: $X[i, j, \ell] \leftarrow X[i, j, \ell] + X[i, j, \ell + 2^{r-1}]$
- 8: for $1 \le i, j \le n$ pardo
- 9: $C[i,j] \leftarrow X[i,j,1]$

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



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Phase 1

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

Phase 2 (round r) p_i updates $X[i,j,\ell]$ for all $1 \le j \le n; 1 \le \ell \mod 2^r = 1$ $\mathcal{O}(n \cdot n/2^r)$ time; no communication

Phase 3 p_i writes i-th row into C[i, j]'s n time; n communication



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

$A_{1,1}$ $A_{1,2}$ $A_{1,3}$ $A_{1,4}$	$B_{1,1}$	$B_{1,2}$	$B_{1,3}$	$B_{1,4}$		$C_{1,1}$	C _{1,2}	C _{1,3}	$C_{1,4}$
$A_{2,1}$ $A_{2,2}$ $A_{2,3}$ $A_{2,4}$	$B_{2,1}$	B _{2,2}	$B_{2,3}$	$B_{2,4}$	_	$C_{2,1}$	C _{2,2}	$C_{2,3}$	$C_{2,4}$
$A_{3,1}$ $A_{3,2}$ $A_{3,3}$ $A_{3,4}$	$B_{3,1}$	B _{3,2}	$B_{3,3}$	$B_{3,4}$	_	C _{3,1}	$C_{3,2}$	$C_{3,3}$	$C_{3,4}$
A _{4,1} A _{4,2} A _{4,3} A _{4,4}	B _{4,1}	B _{4,2}	B _{4,3}	B _{4,4}		$C_{4,1}$	$C_{4,2}$	$C_{4,3}$	$C_{4,4}$

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 $n^{4/3}$.

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

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 $n^{4/3}$.

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$\begin{bmatrix} A_{1,1} & A_{1,2} & A_{1,3} & A_{1,4} \end{bmatrix}$	$B_{1,1}$	$B_{1,2}$	$B_{1,3}$	$B_{1,4}$		$C_{1,1}$	C _{1,2}	C _{1,3}	$C_{1,4}$
$A_{2,1}$ $A_{2,2}$ $A_{2,3}$ $A_{2,4}$	$B_{2,1}$	B _{2,2}	$B_{2,3}$	$B_{2,4}$	_	$C_{2,1}$	C _{2,2}	C _{2,3}	$C_{2,4}$
$A_{3,1}$ $A_{3,2}$ $A_{3,3}$ $A_{3,4}$	$B_{3,1}$	B _{3,2}	$B_{3,3}$	$B_{3,4}$	_	$C_{3,1}$	C _{3,2}	C _{3,3}	C _{3,4}
A _{4,1} A _{4,2} A _{4,3} A _{4,4}	B _{4,1}	B _{4,2}	B _{4,3}	B _{4,4}		$C_{4,1}$	C _{4,2}	C _{4,3}	C _{4,4}

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$A_{3,1}$ $A_{3,2}$ $A_{3,3}$ $A_{3,4}$	$B_{3,1}$	B _{3,2}	$B_{3,3}$	$B_{3,4}$	_	$C_{3,1}$	C _{3,2}	C _{3,3}	C _{3,4}
A _{4,1} A _{4,2} A _{4,3} A _{4,4}	B _{4,1}	B _{4,2}	B _{4,3}	B _{4,4}		$C_{4,1}$	C _{4,2}	C _{4,3}	C _{4,4}

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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 $\mathcal{O}(n^{4/3})$

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



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