WS 2014/15

Parallel Algorithms

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http://www14.in.tum.de/lehre/2014WS/pa/

Winter Term 2014/15

Part I

Organizational Matters

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Modul: IN2011

Name: "Parallel Algorithms" "Parallele Algorithmen"

ECTS: 8 Credit points

Lectures:

4 SWS
 Mon 14:00-16:00 (Room 00.08.038)
 Fri 8:30-10:00 (Room 00.08.038)

Webpage: http://www14.in.tum.de/lehre/2014WS/pa/

- Required knowledge:
 - ► IN0001, IN0003
 - "Introduction to Informatics 1/2"
 - "Einführung in die Informatik 1/2"
 ► IN0007
 - "Fundamentals of Algorithms and Data Structures"
 - "Grundlagen: Algorithmen und Datenstrukturen" (GAD)

 IN0011
 - "Basic Theoretic Informatics"
 - "Einführung in die Theoretische Informatik" (THEO)
 IN0015
 - "Discrete Structures"
 - "Diskrete Strukturen" (DS)

 N0018
 - "Discrete Probability Theory"

 "Discrete Wahrschainlighkeitstheorie" (DWT)
 - "Diskrete Wahrscheinlichkeitstheorie" (DWT)

 IN2003
 - "Efficient Algorithms and Data Structures"
 "Effiziente Algorithmen und Datenstrukturen"

The Lecturer

- ► Harald Räcke
- ► Email: raecke@in.tum.de
- Room: 03.09.044
- Office hours: (per appointment)

Tutorials

- ► Tutors:
 - Chris Pinkau
 - pinkau@in.tum.de
 - Room: 03.09.037
 - Office hours: Tue 13:00-14:00
- ▶ Room: 03.11.018
- ► Time: Tue 14:00-16:00

Assignment sheets

In order to pass the module you need to pass a 3 hour exam

Assessment

Assignment Sheets:

- An assignment sheet is usually made available on Monday on the module webpage.
- Solutions have to be handed in in the following week before the lecture on Monday.
- You can hand in your solutions by putting them in the right folder in front of room 03.09.019A.
- Solutions will be discussed in the subsequent tutorial on Tuesday.

1 Contents

- PRAM algorithms
 - Parallel Models
 - PRAM Model
 - Basic PRAM Algorithms
 - Sorting
 - Lower Bounds
- Networks of Workstations
 - Offline Permutation Routing on the Mesh
 - Oblivious Routing in the Butterfly
 - Greedy Routing
 - Sorting on the Mesh
 - ASCEND/DESCEND Programs
 - Embeddings between Networks

2 Literatur

Tom Leighton:

Introduction to Parallel Algorithms and Architecture: Arrays, Trees, Hypercubes,

Morgan Kaufmann: San Mateo, CA, 1992

Joseph JaJa:

An Introduction to Parallel Algorithms,

Addison-Wesley: Reading, MA, 1997

Jeffrey D. Ullman:

Computational Aspects of VLSI,

Computer Science Press: Rockville, USA, 1984

Selim G. Akl.:

The Design and Analysis of Parallel Algorithms,

Prentice Hall: Englewood Cliffs, NJ, 1989

Part II

Foundations

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, $S_p(n) \leq p$. 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.

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.

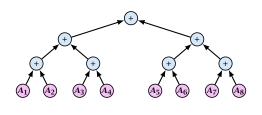
A real satisfactory model does not exist!

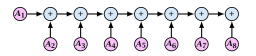
DAG model — computation graph

- 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.,

- $p_v \in \{1, ..., p\}; t_v \in \{1, ..., T\}$
- $t_u = t_v \Rightarrow p_u \neq p_v$
- $(u,v) \in E \Rightarrow t_v \ge t_u + 1$

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.

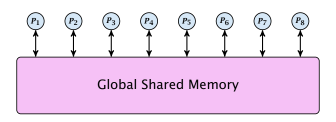
In principle, there could be a different DAG for every input size n.

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
- ▶ $id \in \{1, ..., p\}$: the id of the current processor

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

Algorithm 1 copy

- 1: **if** id = 1 **then** $round \leftarrow 1$
- 2: while $round \le p$ 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]...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]
```

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

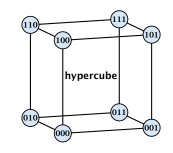
PRAM Model — remarks

- 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

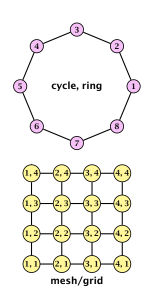
Network of Workstations — NOWs

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

Typical Topologies







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]...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]
```

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 O(T(n))
- $\mathcal{O}(C(n))$ cost and time $\mathcal{O}(T(n))$
- ▶ $\mathcal{O}(C(n)/p)$ time for any number $p \le P(n)$ 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

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

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

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

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

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

Communication Cost

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; $n = 2^k$ 2: Output: C = AB3: 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 $\log n$

7:
$$X[i, j, \ell] \leftarrow X[i, j, \ell] + X[i, j, \ell + 2^{r-1}]$$

6: for $1 \le i, j \le n$; $\ell \mod 2^r = 1$ pardo

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.

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 $1\leq j,\ell\leq n$ 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

Alternative Algorithm

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 $n^{4/3}$.

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

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

much, much better in practise

Part III

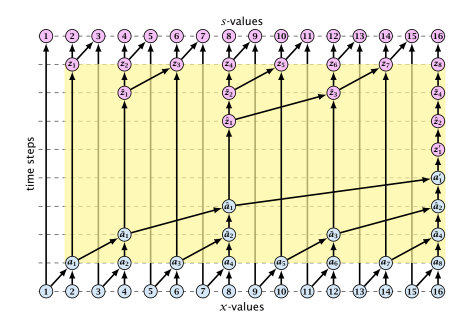
PRAM Algorithms

Prefix Sum

```
input: x[1]...x[n]
output: s[1]...s[n] with s[i] = \sum_{j=1}^{i} x[i] (w.r.t. operator *)
```

```
Algorithm 6 PrefixSum(n, x[1]...x[n])
1: // compute prefixsums; n = 2^k
2: if n = 1 then s[1] \leftarrow x[1]; return
3: for 1 \le i \le n/2 pardo
4: a[i] \leftarrow x[2i-1] * x[2i]
5: z[1],...,z[n/2] \leftarrow \text{PrefixSum}(n/2,a[1]...a[n/2])
6: for 1 \le i \le n pardo
7: i \text{ even } : s[i] \leftarrow z[i/2]
8: i = 1 : s[1] = x[1]
      i odd : s[i] \leftarrow z[(i-1)/2] * x[i]
```

Prefix Sum



Prefix Sum

The algorithm uses work $\mathcal{O}(n)$ and time $\mathcal{O}(\log n)$ for solving Prefix Sum on an EREW-PRAM with n processors.

It is clearly work-optimal.

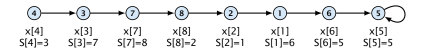
Theorem 6

On a CREW PRAM a Prefix Sum requires running time $\Omega(\log n)$ regardless of the number of processors.

Parallel Prefix

Input: a linked list given by successor pointers; a value x[i] for every list element; an operator *;

Output: for every list position ℓ the sum (w.r.t. *) of elements after ℓ in the list (including ℓ)



Parallel Prefix

Algorithm 7 ParallelPrefix 1: for $1 \le i \le n$ pardo 2: $P[i] \leftarrow S[i]$ 3: while $S[i] \ne S[S[i]]$ do 4: $x[i] \leftarrow x[i] * x[S[i]]$ 5: $S[i] \leftarrow S[S[i]]$ 6: if $P[i] \ne i$ then $x[i] \leftarrow x[i] * x[S(i)]$

The algorithm runs in time $O(\log n)$.

It has work requirement $\mathcal{O}(n \log n)$. non-optimal

This technique is also known as pointer jumping

Given two sorted sequences $A = (a_1, ..., a_n)$ and $B = (b_1, ..., b_n)$, compute the sorted squence $C = (c_1, ..., c_n)$.

Definition 7

Let $X = (x_1, ..., x_t)$ be a sequence. The rank $\operatorname{rank}(y:X)$ of y in X is

$$rank(y:X) = |\{x \in X \mid x \le y\}|$$

For a sequence $Y = (y_1, ..., y_s)$ we define $\operatorname{rank}(Y : X) := (r_1, ..., r_s)$ with $r_i = \operatorname{rank}(y_i : X)$.

Given two sorted sequences $A = (a_1 ... a_n)$ and $B = (b_1 ... b_n)$, compute the sorted squence $C = (c_1 ... c_n)$.

Observation:

We can assume wlog. that elements in A and B are different.

Then for $c_i \in C$ we have $i = \operatorname{rank}(c_i : A \cup B)$.

This means we just need to determine $rank(x : A \cup B)$ for all elements!

Observe, that $rank(x : A \cup B) = rank(x : A) + rank(x : B)$.

Clearly, for $x \in A$ we already know rank(x : A), and for $x \in B$ we know rank(x : B).

Compute $\operatorname{rank}(x:A)$ for all $x \in B$ and $\operatorname{rank}(x:B)$ for all $x \in A$. can be done in $\mathcal{O}(\log n)$ time with 2n processors by binary search

Lemma 8

On a CREW PRAM, Merging can be done in $\mathcal{O}(\log n)$ time and $\mathcal{O}(n\log n)$ work.

not optimal

$$A = (a_1, \dots, a_n); B = (b_1, \dots, b_n);$$

 $\log n$ integral; $k := n/\log n$ integral;

Algorithm 8 GenerateSubproblems

- 1: $j_0 \leftarrow 0$ 2: $j_k \leftarrow n$
- 3: for $1 \le i \le k-1$ pardo
- 4: $j_i \leftarrow \operatorname{rank}(b_{i \log n} : A)$
- 5: for $0 \le i \le k-1$ pardo
- 6: $B_i \leftarrow (b_{i \log n + 1}, \dots, b_{(i+1) \log n})$ 7: $A_i \leftarrow (a_{j_i + 1}, \dots, a_{j_{i+1}})$

If C_i is the merging of A_i and B_i then the sequence $C_0 \dots C_{k-1}$ is a sorted sequence.

We can generate the subproblems in time $O(\log n)$ and work O(n).

Note that in a sub-problem B_i has length $\log n$.

If we run the algorithm again for every subproblem, (where A_i takes the role of B) we can in time $\mathcal{O}(\log\log n)$ and work $\mathcal{O}(n)$ generate subproblems where A_j and B_j have both length at most $\log n$.

Such a subproblem can be solved by a single processor in time $\mathcal{O}(\log n)$ and work $\mathcal{O}(|A_i| + |B_i|)$.

Parallelizing the last step gives total work $\mathcal{O}(n)$ and time $\mathcal{O}(\log n)$.

the resulting algorithm is work optimal

4.4 Maximum Computation

Lemma 9

On a CRCW PRAM the maximum of n numbers can be computed in time O(1) with n^2 processors.

proof on board...

4.4 Maximum Computation

Lemma 10

On a CRCW PRAM the maximum of n numbers can be computed in time $O(\log \log n)$ with n processors and work $O(n \log \log n)$.

proof on board...

4.4 Maximum Computation

Lemma 11

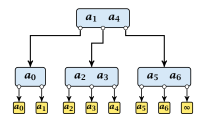
On a CRCW PRAM the maximum of n numbers can be computed in time $O(\log \log n)$ with n processors and work O(n).

proof on board...

4.5 Inserting into a (2, 3)-tree

Given a (2,3)-tree with n elements, and a sequence $x_0 < x_1 < x_2 < \cdots < x_k$ of elements. We want to insert elements x_1, \ldots, x_k into the tree $(k \ll n)$.

time: $O(\log n)$; work: $O(k \log n)$



4.5 Inserting into a (2, 3)-tree

1. determine for every x_i the leaf element before which it has to be inserted time: $\mathcal{O}(\log n)$; work: $\mathcal{O}(k\log n)$; CREW PRAM

all x_i 's that have to be inserted before the same element form a $\frac{1}{2}$

determine the largest/smallest/middle element of every chain

time: $O(\log k)$; work: O(k);

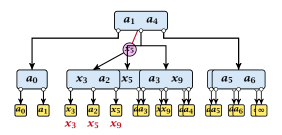
3. insert the middle element of every chain compute new chains

time: $\mathcal{O}(\log n)$; work: $\mathcal{O}(k_i \log n + k)$; k_i = #inserted elements

(computing new chains is constant time)

4. repeat Step 3 for logarithmically many rounds time: $O(\log n \log k)$; work: $O(k \log n)$;

Step 3



- each internal node is split into at most two parts
- each split operation promotes at most one element
- hence, on every level we want to insert at most one element per successor pointer
- we can use the same routine for every level

4.5 Inserting into a (2, 3)-tree

- Step 3, works in phases; one phase for every level of the tree
- Step 4, works in rounds; in each round a different set of elements is inserted

Observation

We can start with phase i of round r as long as phase i of round r-1 and (of course), phase i-1 of round r has finished.

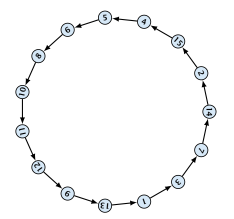
This is called Pipelining. Using this technique we can perform all rounds in Step 4 in just $O(\log k + \log n)$ many parallel steps.

The following algorithm colors an n-node cycle with $\lceil \log n \rceil$ colors.

Algorithm 9 BasicColoring

- 1: for $1 \le i \le n$ pardo
- 2: $\operatorname{col}(i) \leftarrow i$ 3: $k_i \leftarrow \operatorname{smallest} \operatorname{bitpos} \operatorname{where} \operatorname{col}(i) \operatorname{and} \operatorname{col}(S(i)) \operatorname{differ}$ 4: $\operatorname{col}'(i) \leftarrow 2k_i + \operatorname{col}(i)_{k_i}$

(bit positions are numbered starting with 0)



| | _ | _ | , |
|----|------|---|------|
| v | col | k | col' |
| 1 | 0001 | 1 | 2 |
| 3 | 0011 | 2 | 4 |
| 7 | 0111 | 0 | 1 |
| 14 | 1110 | 2 | 5 |
| 2 | 0010 | 0 | 0 |
| 15 | 1111 | 0 | 1 |
| 4 | 0100 | 0 | 0 |
| 5 | 0101 | 0 | 1 |
| 6 | 0110 | 1 | 3 |
| 8 | 1000 | 1 | 2 |
| 10 | 1010 | 0 | 0 |
| 11 | 1011 | 0 | 1 |
| 12 | 1100 | 0 | 0 |
| 9 | 1001 | 2 | 4 |
| 13 | 1101 | 2 | 5 |

Applying the algorithm to a coloring with bit-length t generates a coloring with largest color at most

$$2(t-1)+1$$

and bit-length at most

$$\lceil \log_2(2(t-1)+1) \rceil \leq \lceil \log_2(2t) \rceil = \lceil \log_2(t) \rceil + 1$$

Applying the algorithm repeatedly generates a constant number of colors after $\mathcal{O}(\log^* n)$ operations.

Note that the first inequality holds because 2(t-1)-1 is odd.

As long as the bit-length $t \ge 4$ the bit-length decreases.

Applying the algorithm with bit-length 3 gives a coloring with colors in the range 0, ..., 5 = 2t - 1.

We can improve to a 3-coloring by successively re-coloring nodes from a color-class:

```
Algorithm 10 ReColor

1: for \ell \leftarrow 5 to 3

2: for 1 \le i \le n pardo

3: if \operatorname{col}(i) = \ell then

4: \operatorname{col}(i) \leftarrow \min\{\{0,1,2\} \setminus \{\operatorname{col}(P[i]), \operatorname{col}(S[i])\}\}
```

This requires time O(1) and work O(n).

Lemma 12

We can color vertices in a ring with three colors in $O(\log^* n)$ time and with $O(n \log^* n)$ work.

not work optimal

Lemma 13

Given n integers in the range $0, \ldots, \mathcal{O}(\log n)$, there is an algorithm that sorts these numbers in $\mathcal{O}(\log n)$ time using a linear number of operations.

Proof: Exercise!

```
Algorithm 11 OptColor

1: for 1 \le i \le n pardo
```

2: $\operatorname{col}(i) \leftarrow i$

3: apply BasicColoring once

4: sort vertices by colors

5: **for** $\ell = 2\lceil \log n \rceil$ **to** 3 **do**

6: **for** all vertices i of color ℓ **pardo**

7: $\operatorname{col}(i) \leftarrow \min\{\{0, 1, 2\} \setminus \{\operatorname{col}(P[i]), \operatorname{col}(S[i])\}\}$

We can perform Lines 6 and 7 in time $\mathcal{O}(n_\ell)$ only because we sorted before. In general a state-ment like "for constraint pardo" should only contain a contraint on the id's of the processors but not something complicated (like the color) which has to be checked and, hence, induces work. Because of the sorting we can transform this complicated constraint into a constraint on just the processor id's.

Lemma 14

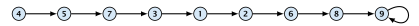
A ring can be colored with 3 colors in time $O(\log n)$ and with work O(n).

work optimal but not too fast

List Ranking

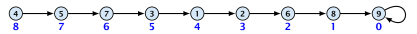
Input:

A list given by successor pointers;



Output:

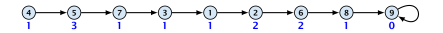
For every node number of hops to end of the list;



Observation:

Special case of parallel prefix

List Ranking

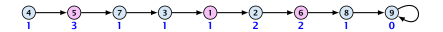


 Given a list with values; perhaps from previous iterations.

The list is given via predecessor pointers P(i) and successor pointers S(i).

$$S(4) = 5$$
, $S(2) = 6$, $P(3) = 7$, etc.

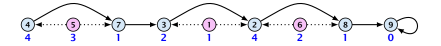
List Ranking



2. Find an independent set; time: $O(\log n)$; work: O(n).

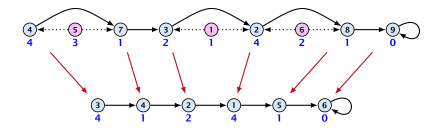
The independent set should contain a constant fraction of the vertices.

Color vertices; take local minima



3. Splice the independent set out of the list;

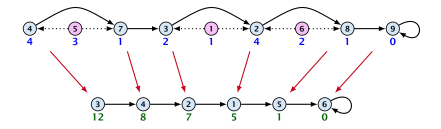
At the independent set vertices the array still contains old values for P(i) and S(i);



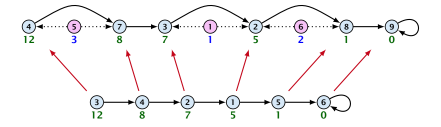
4. Compress remaining n' nodes into a new array of n' entries.

The index positions can be computed by a prefix sum in time $\mathcal{O}(\log n)$ and work $\mathcal{O}(n)$

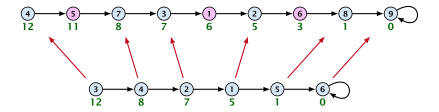
Pointers can then be adjusted in time $\mathcal{O}(1)$.



5. Solve the problem on the remaining list. If current size is less than $n/\log n$ do pointer jumping: time $\mathcal{O}(\log n)$; work $\mathcal{O}(n)$. Otherwise continue shrinking the list by finding an independent set



6. Map the values back into the larger list. Time: $\mathcal{O}(1)$; Work: $\mathcal{O}(n)$



- 7. Compute values for independent set nodes. Time: $\mathcal{O}(1)$; Work: $\mathcal{O}(1)$.
- **8.** Splice nodes back into list. Time: O(1); Work: O(1).

We need $\mathcal{O}(\log \log n)$ shrinking iterations until the size of the remaining list reaches $\mathcal{O}(n/\log n)$.

Each shrinking iteration takes time $O(\log n)$.

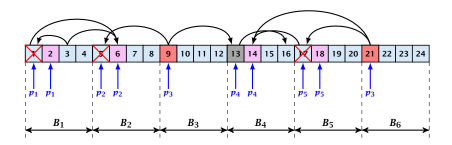
The work for all shrinking operations is just $\mathcal{O}(n)$, as the size of the list goes down by a constant factor in each round.

List Ranking can be solved in time $\mathcal{O}(\log n \log \log n)$ and work $\mathcal{O}(n)$ on an EREW-PRAM.

Optimal List Ranking

In order to reduce the work we have to improve the shrinking of the list to $\mathcal{O}(n/\log n)$ nodes.

After this we apply pointer jumping



- some nodes are active;
- active nodes without neighbouring active nodes are isolated;
- the others form sublists;

work: $\psi(n)$;

- 1 delete isolated nodes from the list;
- **2** color each sublist with $O(\log \log n)$ colors; time: O(1);

Optimal List Ranking

Each iteration requires constant time and work $O(n/\log n)$, because we just work on one node in every block.

We need to prove that we just require $O(\log n)$ iterations to reduce the size of the list to $O(n/\log n)$.

Observations/Remarks:

- ▶ If the *p*-pointer of a block cannot be advanced without leaving the block, the processor responsible for this block simply stops working; all other blocks continue.
- The p-node of a block (the node p_i is pointing to) at the beginning of a round is either a ruler with a living subject or the node will become active during the round.
- The subject nodes always lie to the left of the p-node of the respective block (if it exists).

Measure of Progress:

- a ruler will delete a subject
- an active node either
 - becomes a ruler (with a subject)
 - becomes a subject
 - is isolated and therefore gets deleted

Analysis

For the analysis we assign a weight to every node in every block as follows.

Definition 15

The weight of the i-th node in a block is

$$(1 - q)^{i}$$

with $q = \frac{1}{\log \log n}$, where the node-numbering starts from 0. Hence, a block has nodes $\{0, \dots, \log n - 1\}$.

Definition of Rulers

Properties:

- ▶ A ruler should have at most $\log \log n$ subjects.
- The weight of a ruler should be at most the weight of any of its subjects.
- Each ruler must have at least one subject.
- We must be able to remove the next subject in constant time.
- We need to make the ruler/subject decision in constant time.

Given a sublist of active nodes.

Color the sublist with $O(\log \log n)$ colors. Take the local minima w.r.t. this coloring.

If the first node is not a ruler

- if the second node is a ruler switch ruler status between first and second
- otw. just make the first node into a ruler

This partitions the sub-list into chains of length at most $\log \log n$ each starting with a ruler

Now we change the ruler definition.

Consider some chain.

We make all local minima w.r.t. the weight function into a ruler; ties are broken according to block-id (so that comparing weights gives a strict inequality).

A ruler gets as subjects the nodes left of it until the next local maximum (or the start of the chain) (including the local maximum) and the nodes right of it until the next local maximum (or the end of the chain) (excluding the local maximum).

In case the first node is a ruler the above definition could leave it without a subject. We use constant time to fix this in some arbitrary manner

Set
$$q = \frac{1}{\log \log n}$$
.

The *i*-th node in a block is assigned a weight of $(1-q)^i$, $0 \le i < \log n$

The total weight of a block is at most 1/q and the total weight of all items is at most $\frac{n}{q \log n}$.

to show:

After $\mathcal{O}(\log n)$ iterations the weight is at most $(n/\log n)(1-q)^{\log n}$

This means at most $n/\log n$ nodes remain because the smallest weight a node can have is $(1-q)^{\log n-1}$.

In every iteration the weight drops by a factor of

$$(1 - q/4)$$
.

We consider subject nodes to just have half their weight.

We can view the step of becoming a subject as a precursor to deletion.

Hence, a node looses half its weight when becoming a subject and the remaining half when deleted.

Note that subject nodes will be deleted after just an additional $\mathcal{O}(\log\log n)$ iterations.

The weight is reduced because

- An isolated node is removed.
- ► A node is labelled as ruler, and the corresponding subjects reduce their weight by a factor of 1/2.
- A node is a ruler and deletes one of its subjects.

Hence, the weight reduction comes from p-nodes (ruler/active).

Each p-node is responsible for some other nodes; it has to generate a weight reduction large enough so that the weight of all nodes it is responsible for decreases by the desired factor.

An active node is responsible for all nodes that come after it in its block.

A ruler is responsible for all nodes that come after it in its block and for all its subjects.

Note that by this definition every node remaining in the list is covered.

Case 1: Isolated Node

Suppose we delete an isolated node v that is the i-th node in its block.

The weight of all node that v is responsible for is

$$\sum_{i \le j < \log n} (1-q)^j$$

This weight reduces to

$$\sum_{i < j < \log n} (1 - q)^j \le (1 - q) \sum_{i \le j < \log n} (1 - q)^j$$

Hence, weight reduces by a factor $(1 - q) \le (1 - q/4)$.

Case 2: Creating Subjects

Suppose we generate a ruler with at least one subject.

Weight of ruler: $(1-q)^{i_1}$.

Weight of subjects: $(1-q)^{i_j}$, $2 \le j \le k$.

Initial weight:

$$Q = \sum_{j=1}^{k} \sum_{i_j \le \ell < \log n} (1 - q)^{\ell} \le \frac{1}{q} \sum_{j=1}^{k} (1 - q)^{i_j} \le \frac{2}{q} \sum_{j=2}^{k} (1 - q)^{i_j}$$

New weight:

$$Q' = Q - \frac{1}{2} \sum_{i=2}^{k} (1 - q)^{i_j} \le (1 - \frac{q}{4})Q$$

Case 3: Removing Subjects

weight of ruler: $(1-q)^{i_1}$; weight of subjects: $(1-q)^{i_j}$, $2 \le j \le k$

Assume ruler removes subject with largest weight say i_2 (why?).

Initial weight:

$$Q = \sum_{i_1 \le \ell < \log n} (1 - q)^{\ell} + \frac{1}{2} \sum_{j=2}^{k} (1 - q)^{i_j}$$

$$\le \frac{1}{q} (1 - q)^{i_1} + \frac{k}{2} (1 - q)^{i_2}$$

$$\le \frac{1}{q} (1 - q)^{i_2} + \frac{1}{2q} (1 - q)^{i_2}$$

New weight:

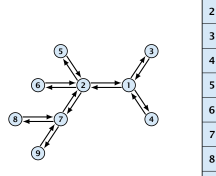
$$Q' = Q - \frac{1}{2}(1 - q)^{i_2} \le (1 - \frac{q}{3})Q$$

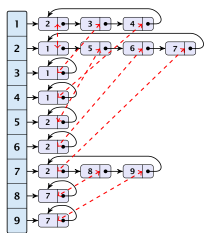
After s iterations the weight is at most

$$\frac{n}{q \log n} \left(1 - \frac{q}{4} \right)^s \stackrel{!}{\leq} \frac{n}{\log n} (1 - q)^{\log n}$$

Choosing $i = 5 \log n$ the inequality holds for sufficiently large n.

Tree Algorithms





Euler Circuits

Every node v fixes an arbitrary ordering among its adjacent nodes:

$$u_0, u_1, \ldots, u_{d-1}$$

We obtain an Euler tour by setting

$$\operatorname{succ}((u_i, v)) = (v, u_{(i+1) \bmod d})$$

Euler Circuits

Lemma 16

An Euler circuit can be computed in constant time O(1) with O(n) operations.

Rooting a tree

- split the Euler tour at node r
- this gives a list on the set of directed edges (Euler path)
- assign x[e] = 1 for every edge;
- perform parallel prefix; let $s[\cdot]$ be the result array
- if s[(u,v)] < s[(v,u)] then u is parent of v;

Postorder Numbering

- split the Euler tour at node r
- this gives a list on the set of directed edges (Euler path)
- ▶ assign x[e] = 1 for every edge (v, parent(v))
- ▶ assign x[e] = 0 for every edge (parent(v), v)
- perform parallel prefix
- ightharpoonup post(v) = s[(v, parent(v))]; post(r) = n

Level of nodes

- split the Euler tour at node r
- this gives a list on the set of directed edges (Euler path)
- ▶ assign x[e] = -1 for every edge (v, parent(v))
- ▶ assign x[e] = 1 for every edge (parent(v), v)
- perform parallel prefix
- level(v) = s[(parent(v), v)]; level(r) = 0

Number of descendants

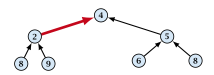
- split the Euler tour at node r
- this gives a list on the set of directed edges (Euler path)
- ▶ assign x[e] = 0 for every edge (parent(v), v)
- ▶ assign x[e] = 1 for every edge $(v, parent(v)), v \neq r$
- perform parallel prefix
- ightharpoonup size(v) = s[(v, parent(v))] s[(parent(v), v)]

Rake Operation

Given a binary tree T.

Given a leaf $u \in T$ with $p(u) \neq r$ the rake-operation does the following

- remove u and p(u)
- attach sibling of u to p(p(u))



We want to apply rake operations to a binary tree T until T just consists of the root with two children.

Possible Problems:

- we could concurrently apply the rake-operation to two siblings
- 2. we could concurrently apply the rake-operation to two leaves u and v such that p(u) and p(v) are connected

By choosing leaves carefully we ensure that none of the above cases occurs

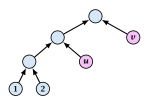
Algorithm:

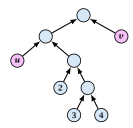
- label leaves consecutively from left to right (excluding left-most and right-most leaf), and store them in an array A
- for $\lceil \log(n+1) \rceil$ iterations
 - apply rake to all odd leaves that are left children
 - apply rake operation to remaining odd leaves (odd at start of round!!!)
 - A=even leaves

Observations

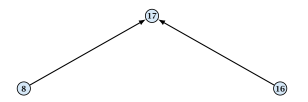
- the rake operation does not change the order of leaves
- two leaves that are siblings do not perform a rake operation in the same round because one is even and one odd at the start of the round
- two leaves that have adjacent parents either have different parity (even/odd) or they differ in the type of child (left/right)

Cases, when the left edge btw. p(u) and p(v) is a left-child edge.





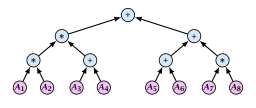
Example

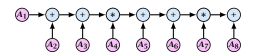


- one iteration can be performed in constant time with $\mathcal{O}(|A|)$ processors, where A is the array of leaves;
- ▶ hence, **all** iterations can be performed in $O(\log n)$ time and O(n) work;
- ▶ the intial parallel prefix also requires time $O(\log n)$ and work O(n)

Evaluating Expressions

Suppose that we want to evaluate an expression tree, containing additions and multiplications.





If the tree is not balanced this may be time-consuming.

We can use the rake-operation to do this quickly.

Applying the rake-operation changes the tree.

In order to maintain the value we introduce parameters a_{ν} and b_{ν} for every node that still allows to compute the value of a node based on the value of its children.

Invariant:

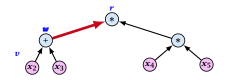
Let u be internal node with children v and w. Then

$$val(u) = (a_v \cdot val(v) + b_v) \otimes (a_w \cdot val(w) + b_w)$$

where $\otimes \in \{*, +\}$ is the operation at node u.

Initially, we can choose $a_v = 1$ and $b_v = 0$ for every node.

Rake Operation



Currently the value at u is

$$val(u) = (a_v \cdot val(v) + b_v) + (a_w \cdot val(w) + b_w)$$
$$= x_1 + (a_w \cdot val(w) + b_w)$$

In the expression for r this goes in as

$$a_{u} \cdot [x_{1} + (a_{w} \cdot \text{val}(w) + b_{w})] + b_{u}$$

$$= \underbrace{a_{u}a_{w}}_{a'_{w}} \cdot \text{val}(w) + \underbrace{a_{u}x_{1} + a_{u}b_{w} + b_{u}}_{b'_{w}}$$

If we change the a and b-values during a rake-operation according to the previous slide we can calculate the value of the root in the end.

Lemma 17

We can evaluate an arithmetic expression tree in time $O(\log n)$ and work O(n) regardless of the height or depth of the tree.

By performing the rake-operation in the reverse order we can also compute the value at each node in the tree.

Lemma 18

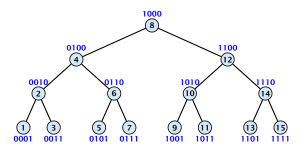
We compute tree functions for arbitrary trees in time $O(\log n)$ and a linear number of operations.

proof on board...

In the LCA (least common ancestor) problem we are given a tree and the goal is to design a data-structure that answers LCA-queries in constant time.

Least Common Ancestor

LCAs on complete binary trees (inorder numbering):

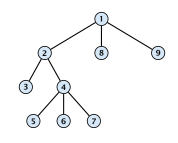


The least common ancestor of u and v is

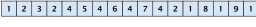
$$z_1 z_2 \dots z_i 10 \dots 0$$

where z_{i+1} is the first bit-position in which u and v differ.

Least Common Ancestor



nodes



levels

 $\ell(v)$ is index of first appearance of v in node-sequence.

r(v) is index of last appearance of v in node-squence.

 $\ell(v)$ and r(v) can be computed in constant time, given the node- and level-sequence.

Least Common Ancestor

Lemma 19

- **1.** u is ancestor of v iff $\ell(u) < \ell(v) < r(u)$
- **2.** u and v are not related iff either $r(u) < \ell(v)$ or $r(v) < \ell(u)$
- 3. suppose $r(u) < \ell(v)$ then LCA(u, v) is vertex with minimum level over interval $[r(u), \ell(v)]$.

Range Minima Problem

Given an array A[1...n], a range minimum query (ℓ,r) consists of a left index $\ell \in \{1,...,n\}$ and a right index $r \in \{1,...,n\}$.

The answer has to return the index of the minimum element in the subsequence $A[\ell \dots r]$.

The goal in the range minima problem is to preprocess the array such that range minima queries can be answered quickly (constant time).

Observation

Given an algorithm for solving the range minima problem in time T(n) and work W(n) we can obtain an algorithm that solves the LCA-problem in time $\mathcal{O}(T(n) + \log n)$ and work $\mathcal{O}(n + W(n))$.

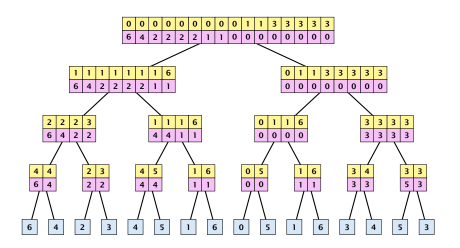
Remark

In the sequential setting the LCA-problem and the range minima problem are equivalent. This is not necessarily true in the parallel setting.

For solving the LCA-problem it is sufficient to solve the restricted range minima problem where two successive elements in the array just differ by +1 or -1.

Prefix and Suffix Minima

Tree with prefix-minima and suffix-minima:



- Suppose we have an array A of length $n = 2^k$
- ▶ We compute a complete binary tree *T* with *n* leaves.
- Each internal node corresponds to a subsequence of A. It contains an array with the prefix and suffix minima of this subsequence.

Given the tree T we can answer a range minimum query (ℓ, r) in constant time.

- we can determine the LCA x of ℓ and r in constant time since T is a complete binary tree
- ▶ Then we consider the suffix minimum of ℓ in the left child of x and the prefix minimum of r in the right child of x.
- The minimum of these two values is the result.

Lemma 20

We can solve the range minima problem in time $\mathcal{O}(\log n)$ and work $\mathcal{O}(n\log n)$.

Reducing the Work

Partition A into blocks B_i of length $\log n$

Preprocess each B_i block separately by a sequential algorithm so that range-minima queries within the block can be answered in constant time. (how?)

For each block B_i compute the minimum x_i and its prefix and suffix minima.

Use the previous algorithm on the array $(x_1, \dots, x_{n/\log n})$.

Answering a query (ℓ, r) :

- if ℓ and r are from the same block the data-structure for this block gives us the result in constant time
- if ℓ and r are from different blocks the result is a minimum of three elements:
 - ullet the suffix minmum of entry ℓ in ℓ 's block
 - the minimum among $x_{\ell+1},\ldots,x_{r-1}$
 - ullet the prefix minimum of entry r in r's block

Searching

An extension of binary search with p processors gives that one can find the rank of an element in

$$\log_{p+1}(n) = \frac{\log n}{\log(p+1)}$$

many parallel steps with p processors. (not work-optimal)

This requires a CREW PRAM model. For the EREW model searching cannot be done faster than $\mathcal{O}(\log n - \log p)$ with p processors even if there are p copies of the search key.

Merging

Given two sorted sequences $A = (a_1, ..., a_n)$ and $B = (b_1, ..., b_n)$, compute the sorted squence $C = (c_1, ..., c_n)$.

Definition 21

Let $X = (x_1, ..., x_t)$ be a sequence. The rank $\operatorname{rank}(y:X)$ of y in X is

$$rank(y:X) = |\{x \in X \mid x \le y\}|$$

For a sequence $Y = (y_1, ..., y_s)$ we define $\operatorname{rank}(Y : X) := (r_1, ..., r_s)$ with $r_i = \operatorname{rank}(y_i : X)$.

Merging

We have already seen a merging-algorithm that runs in time $\mathcal{O}(\log n)$ and work $\mathcal{O}(n)$.

Using the fast search algorithm we can improve this to a running time of $\mathcal{O}(\log\log n)$ and work $\mathcal{O}(n\log\log n)$.

Merging

Input:
$$A = a_1, ..., a_n$$
; $B = b_1, ..., b_m$; $m \le n$

- 1. if m < 4 then rank elements of B, using the parallel search algorithm with p processors. Time: $\mathcal{O}(1)$. Work: $\mathcal{O}(n)$.
- 2. Concurrently rank elements $b_{\sqrt{m}}, b_{2\sqrt{m}}, \ldots, b_m$ in A using the parallel search algorithm with $p = \sqrt{n}$. Time: $\mathcal{O}(1)$. Work: $\mathcal{O}(\sqrt{m} \cdot \sqrt{n}) = \mathcal{O}(n)$

$$j(i) := \operatorname{rank}(b_{i\sqrt{m}} : A)$$

3. Let $B_i = (b_{i\sqrt{m}+1}, \ldots, b_{(i+1)\sqrt{m}-1});$ and $A_i = (a_{j(i)+1}, \ldots, a_{j(i+1)}).$

Recursively compute $rank(B_i : A_i)$.

4. Let k be index not a multiple of \sqrt{m} . $i = \lceil \frac{k}{\sqrt{m}} \rceil$. Then $\operatorname{rank}(b_k : A) = j(i) + \operatorname{rank}(b_k : A_i)$.

The algorithm can be made work-optimal by standard techniques.

proof on board...

Mergesort

Lemma 22

A straightforward parallelization of Mergesort can be implemented in time $O(\log n \log \log n)$ and with work $O(n \log n)$.

This assumes the CREW-PRAM model.

Mergesort

Let L[v] denote the (sorted) sublist of elements stored at the leaf nodes rooted at v.

We can view Mergesort as computing L[v] for a complete binary tree where the leaf nodes correspond to nodes in the given array.

Since the merge-operations on one level of the complete binary tree can be performed in parallel we obtain time $\mathcal{O}(h\log\log n)$ and work $\mathcal{O}(hn)$, where $h=\mathcal{O}(\log n)$ is the height of the tree.

We again compute L[v] for every node in the complete binary tree.

After round s, $L_s[v]$ is an **approximation** of L[v] that will be improved in future rounds.

For $s \ge 3 \operatorname{height}(v)$, $L_s[v] = L[v]$.

In every round, a node v sends $\mathrm{sample}(L_s[v])$ (an approximation of its current list) upwards, and receives approximations of the lists of its children.

It then computes a new approximation of its list.

A node is called active in round s if $s \le 3 \operatorname{height}(v)$ (this means its list is not yet complete at the start of the round, i.e., $L_{s-1}[v] \ne L[v]$).

```
Algorithm 11 ColeSort()

1: initialize L_0[v] = A_v for leaf nodes; L_0[v] = \emptyset otw.

2: for s \leftarrow 1 to 3 \cdot \text{height}(T) do

3: for all active nodes v do

4: //u and w children of v

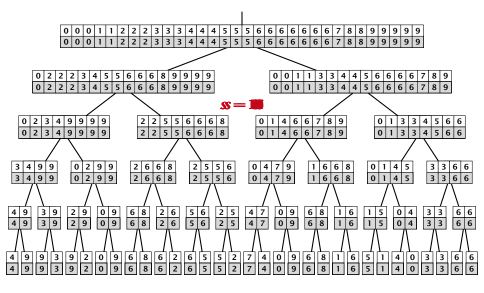
5: L'_s[u] \leftarrow \text{sample}(L_{s-1}[u])

6: L'_s[w] \leftarrow \text{sample}(L_{s-1}[w])

7: L_s[v] \leftarrow \text{merge}(L'_s[u], L'_s[w])
```

$$\operatorname{sample}(L_s[v]) = \begin{cases} \operatorname{sample}_4(L_s[v]) & s \leq 3 \operatorname{height}(v) \\ \operatorname{sample}_2(L_s[v]) & s = 3 \operatorname{height}(v) + 1 \\ \operatorname{sample}_1(L_s[v]) & s = 3 \operatorname{height}(v) + 2 \end{cases}$$

Colesort





Lemma 23

After round $s = 3 \operatorname{height}(v)$, the list $L_s[v]$ is complete.

Proof:

- clearly true for leaf nodes
- suppose it is true for all nodes up to height h;
- fix a node v on level h+1 with children u and w
- $L_{3h}[u]$ and $L_{3h}[w]$ are complete by induction hypothesis
- further sample($L_{3h+2}[u]$) = L[u] and sample($L_{3h+2}[v]$) = L[v]
- hence in round 3h + 3 node v will merge the complete list of its children; after the round L[v] will be complete

Lemma 24

The number of elements in lists $L_s[v]$ for active nodes v is at most O(n).

proof on board...

Definition 25

A sequence X is a c-cover of a sequence Y if for any two consecutive elements α, β from $(-\infty, X, \infty)$ the set $|\{y_i \mid \alpha \leq y_i \leq \beta\}| \leq c$.

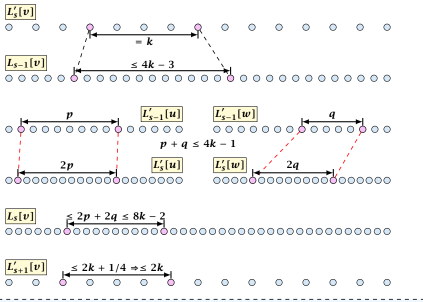
Lemma 26

 $L'_{s}[v]$ is a 4-cover of $L'_{s+1}[v]$.

If [a,b] fulfills $|[a,b] \cap (A \cup \{-\infty,\infty\})| = k$ we say [a,b] intersects $(-\infty,A,+\infty)$ in k items.

Lemma 27

If [a,b] with $a,b \in L'_s[v] \cup \{-\infty,\infty\}$ intersects $(-\infty,L'_s[v],\infty)$ in $k \ge 2$ items, then [a,b] intersects $(-\infty,L'_{s+1},\infty)$ in at most 2k items.



Note that the last step holds as long $L'_{s+1}[v] = \text{sample}_4(L_s[v])$. Otw. $L_{s-1}[v]$ has already been full, and hence, $L'_s[v], L'_{s+1}[v], L'_{s+2}[v]$ are 4-covers of the complete list L[v], and also 4-covers of each other.

Merging with a Cover

Lemma 28

Given two sorted sequences A and B. Let X be a c-cover of A and B for constant c, and let $\operatorname{rank}(X : A)$ and $\operatorname{rank}(X : B)$ be known.

We can merge A and B in time $\mathcal{O}(1)$ using $\mathcal{O}(|X|)$ operations.

Merging with a Cover

Lemma 29

Given two sorted sequences A and B. Let X be a c-cover of B for constant c, and let $\operatorname{rank}(A:X)$ and $\operatorname{rank}(X:B)$ be known.

We can compute $\operatorname{rank}(A:B)$ using $\mathcal{O}(|X|+|A|)$ operations.

Merging with a Cover

Lemma 30

Given two sorted sequences A and B. Let X be a c-cover of B for constant c, and let $\operatorname{rank}(A:X)$ and $\operatorname{rank}(X:B)$ be known.

We can compute rank(B : A) using O(|X| + |A|) operations.

Easy to do with concurrent read. Can also be done with exclusive read but non-trivial.

In order to do the merge in iteration s+1 in constant time we need to know

$$\operatorname{rank}(L_{s}[v]:L'_{s+1}[u])$$
 and $\operatorname{rank}(L_{s}[v]:L'_{s+1}[w])$

and we need to know that $L_s[v]$ is a 4-cover of $L'_{s+1}[u]$ and $L'_{s+1}[w]$.

Lemma 31

 $L_s[v]$ is a 4-cover of $L'_{s+1}[u]$ and $L'_{s+1}[w]$.

- $L_{\mathcal{S}}[v] \supseteq L_{\mathcal{S}}'[u], L_{\mathcal{S}}'[w]$
- L'_s[u] is 4-cover of $L'_{s+1}[u]$
- ▶ Hence, $L_s[v]$ is 4-cover of $L'_{s+1}[u]$ as adding more elements cannot destroy the cover-property.

Analysis

Lemma 32

Suppose we know for every internal node v with children u and w

- ▶ $\operatorname{rank}(L'_{s}[v]:L'_{s+1}[v])$
- rank $(L'_{s}[u]:L'_{s}[w])$
- $ightharpoonup \operatorname{rank}(L'_{S}[w]:L'_{S}[u])$

We can compute

- $ightharpoonup rank(L'_{s+1}[v]:L'_{s+2}[v])$
- $ightharpoonup rank(L'_{s+1}[u]:L'_{s+1}[w])$
- $ightharpoonup rank(L'_{s+1}[w]:L'_{s+1}[u])$

in constant time and $O(|L_{s+1}[v]|)$ operations, where v is the parent of u and w.

Given

- ► $rank(L'_s[u]:L'_{s+1}[u])$ (4-cover)
- $ightharpoonup \operatorname{rank}(L'_{S}[w]:L'_{S}[u])$
- $ightharpoonup \operatorname{rank}(L'_s[u]:L'_s[w])$
- ► rank($L'_s[w]: L'_{s+1}[w]$) (4-cover)

Compute

- $ightharpoonup rank(L'_{s+1}[w]:L'_{s}[u])$
- $rank(L'_{s+1}[u]:L'_s[w])$

Compute

- $ightharpoonup rank(L'_{s+1}[w]:L'_{s+1}[u])$
- ► $rank(L'_{s+1}[u]: L'_{s+1}[w])$

ranks between siblings can be computed easily

Given

- ► $\operatorname{rank}(L'_{s}[u]: L'_{s+1}[u])$ (4-cover $\rightarrow \operatorname{rank}(L'_{s+1}[u]: L'_{s}[u])$)
- ▶ $rank(L'_{s}[w]:L'_{s+1}[u])$
- $ightharpoonup rank(L'_{s}[u]:L'_{s+1}[w])$
- ► $\operatorname{rank}(L'_{s}[w]:L'_{s+1}[w])$ (4-cover $\rightarrow \operatorname{rank}(L'_{s+1}[w]:L'_{s}[w])$)

Compute (recall that $L_s[v] = merge(L'_s[u], L'_s[w])$)

- ightharpoonup rank $(L_s[v]:L'_{s+1}[u])$
- $ightharpoonup \operatorname{rank}(L_s[v]:L'_{s+1}[w])$

Compute

- rank $(L_s[v]:L_{s+1}[v])$ (by adding)
- ► $\operatorname{rank}(L'_{s+1}[v]:L'_{s+2}[v])$ (by sampling)

Definition 33

A 0-1 sequence S is bitonic if it can be written as the concatenation of subsequences S_1 and S_2 such that either

- S₁ is monotonically increasing and S₂ monotonically decreasing, or
- S₁ is monotonically decreasing and S₂ monotonically increasing.

Note, that this just defines bitonic 0-1 sequences. Bitonic sequences are defined differently.

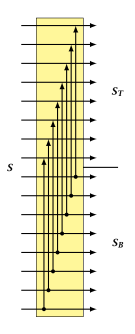
Bitonic Merger

If we feed a bitonic 0-1 sequence S into the network on the right we obtain two bitonic sequences S_T and S_B s.t.

- 1. $S_B \leq S_T$ (element-wise)
- **2.** S_B and S_T are bitonic

Proof:

- assume wlog. S more 1's than 0's.
- ► assume for contradiction two 0s at same comparator $(i, j = i + 2^d)$
 - everything 0 btw i and j means we have more than 50% zeros (ξ).
 - ▶ all 1s btw. i and j means we have less than 50% ones (\$\xeta\$).
 - ▶ 1 btw. *i* and *j* and elsewhere means *S* is not bitonic (﴿).



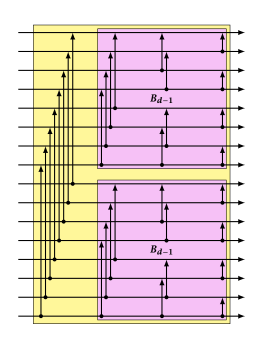
Bitonic Merger

Bitonic Merger B_d

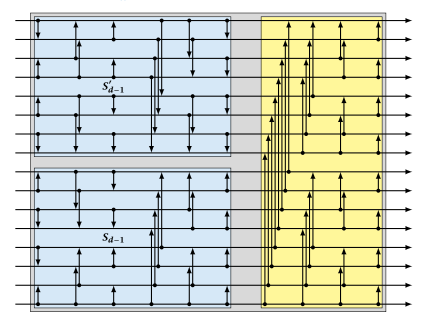
The bitonic merger B_d of dimension d is constructed by combining two bitonic mergers of dimension d-1.

If we feed a bitonic 0-1 sequence into this, the sequence will be sorted.

(actually, any bitonic sequence will be sorted, but we do not prove this)



Bitonic Sorter S_d



Bitonic Merger: $(n = 2^d)$

- ► comparators: $C(n) = 2C(n/2) + n/2 \Rightarrow C(n) = O(n \log n)$.
- ▶ depth: $D(n) = D(n/2) + 1 \Rightarrow D(d) = O(\log n)$.

Bitonic Sorter: $(n = 2^d)$

- comparators: $C(n) = 2C(n/2) + \mathcal{O}(n \log n) \Rightarrow$ $C(n) = \mathcal{O}(n \log^2 n).$
- depth: $D(n) = D(n/2) + \log n \Rightarrow D(n) = \Theta(\log^2 n)$.

Odd-Even Merge

How to merge two sorted sequences?

$$A = (a_1, a_2, ..., a_n), B = (b_1, b_2, ..., b_n), n \text{ even.}$$

Split into odd and even sequences:

$$A_{\text{odd}} = (a_1, a_3, a_5, \dots, a_{n-1}), A_{\text{even}} = (a_2, a_4, a_6, \dots a_n)$$

 $B_{\text{odd}} = (b_1, b_3, b_5, \dots, b_{n-1}), B_{\text{even}} = (b_2, b_4, b_6, \dots, b_n)$

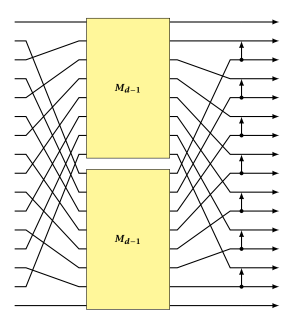
Let

$$X = \text{merge}(A_{\text{odd}}, B_{\text{odd}}) \text{ and } Y = \text{merge}(A_{\text{even}}, B_{\text{even}})$$

Then

$$S = (x_1, \min\{x_2, y_1\}, \max\{x_2, y_1\}, \min\{x_3, y_2\}, \dots, y_n)$$

Odd-Even Merge



Theorem 34

There exists a sorting network with depth $O(\log n)$ and $O(n \log n)$ comparators.

Parallel Comparison Tree Model

A parallel comparison tree (with parallelism p) is a 3^p -ary tree.

- each internal node represents a set of p comparisons btw.
 p pairs (not necessarily distinct)
- ightharpoonup a leaf v corresponds to a unique permutation that is valid for all the comparisons on the path from the root to v
- the number of parallel steps is the height of the tree

Comparison PRAM

A comparison PRAM is a PRAM where we can only compare the input elements;

- we cannot view them as strings
- we cannot do calculations on them

A lower bound for the comparison tree with parallelism $\it p$ directly carries over to the comparison PRAM with $\it p$ processors.

A Lower Bound for Searching

Theorem 35

Given a sorted table X of n elements and an element y. Searching for y in X requires $\Omega(\frac{\log n}{\log(p+1)})$ steps in the parallel comparsion tree with parallelism p < n.

A Lower Bound for Maximum

Theorem 36

A graph G with m edges and n vertices has an independent set on at least $\frac{n^2}{2m+n}$ vertices.

base case (n = 1)

▶ The only graph with one vertex has m = 0, and an independent set of size 1.

induction step $(1, \ldots, n \rightarrow n + 1)$

- Let G be a graph with n + 1 vertices, and v a node with minimum degree (d).
- Let G' be the graph after deleting v and its adjacent vertices in G.
- n' = n (d+1)
- $m' \le m \frac{d}{2}(d+1)$ as we remove d+1 vertices, each with degree at least d
- In G' there is an independent set of size $((n')^2/(2m'+n'))$.
- lacktriangle By adding v we obtain an indepent set of size

$$1 + \frac{(n')^2}{2m' + n'} \ge \frac{n^2}{2m + n}$$

A Lower Bound for Maximum

Theorem 37

Computing the maximum of n elements in the comparison tree requires $\Omega(\log \log n)$ steps whenever the degree of parallelism is $p \le n$.

Theorem 38

Computing the maximum of n elements requires $\Omega(\log\log n)$ steps on the comparison PRAM with n processors.

An adversary can specify the input such that at the end of the (i+1)-st step the maximum lies in a set C_{i+1} of size s_{i+1} such that

- ▶ no two elements of C_{i+1} have been compared
- $> s_{i+1} \ge \frac{s_i^2}{2p + c_i}$

Theorem 39

The selection problem requires $\Omega(\log n/\log\log n)$ steps on a comparison PRAM.

not proven yet

A Lower Bound for Merging

The (k,s)-merging problem, asks to merge k pairs of subsequences A^1, \ldots, A^k and B^1, \ldots, B^k where we know that all elements in $A^i \cup B^i$ are smaller than elements in $A^j \cup B^j$ for (i < j). Further $|A_i|, |B_i| \ge s$.

A Lower Bound for Merging

Lemma 40

Suppose we are given a parallel comparison tree with parallelism p to solve the (k,s) merging problem. After the first step an adversary can specify the input such that an arbitrary (k',s') merging problem has to be solved, where

$$k' = \frac{3}{4} \sqrt{pk}$$

$$s' = \frac{s}{4} \sqrt{\frac{k}{p}}$$

A Lower Bound for Merging

Partition A^is and B^is into blocks of length roughly s/ℓ ; hence ℓ blocks.

Define an $\ell \times \ell$ binary matrix M^i , where $M^i_{\mathcal{X}\mathcal{Y}}$ is 0 iff the parallel step **did not** compare an element from $A^i_{\mathcal{X}}$ with an element from $B^i_{\mathcal{Y}}$.

The matrix has $2\ell - 1$ diagonals.

Choose for every i the diagonal of M^i that has most zeros.

Pair all $A^i_{j+d_i}, B^i_j$, (where $d_i \in \{-(\ell-1), \dots, \ell-1\}$ specifies the chosen diagonal) for which the entry in M^i is zero.

We can choose value s.t. elements for the j-th pair along the diagonal are **all** smaller than for the (j+1)-th pair.

Hence, we get a (k', s') problem.

How many pairs do we have?

- there are $k\ell$ blocks in total
- there are $k \cdot \ell^2$ matrix entries in total
- there are at least $k \cdot \ell^2 p$ zeros.
- choosing a random diagonal (same for every matrix M^i) hits at least

$$\frac{k\ell^2 - p}{2\ell - 1} \ge \frac{k\ell}{2} - \frac{p}{2\ell}$$

zeroes.

► Choosing $\ell = \lceil 2\sqrt{p/k} \rceil$ gives

$$k' \ge \frac{3}{4}\sqrt{pk}$$
 and $s' = \lfloor \frac{s}{\ell} \rfloor \ge \frac{s}{4\sqrt{p/k}} = \frac{s}{4}\sqrt{\frac{k}{p}}$

where we assume $s \ge 6\sqrt{p/k}$.

Lemma 41

Let T(k, s, p) be the number of parallel steps required on a comparison tree to solve the (k, s) merging problem. Then

$$T(k, p, s) \ge \frac{1}{4} \log \frac{\log \frac{p}{k}}{\log \frac{p}{ks}}$$

provided that $p \ge 2ks$ and $p \le ks^2/36$

Induction Step:

Assume that

$$T(k', s', p) \ge \frac{1}{4} \log \frac{\log \frac{p}{k'}}{\log \frac{p}{k's'}}$$

$$\ge \frac{1}{4} \log \frac{\log \frac{4}{3} \sqrt{\frac{p}{k}}}{\log \frac{16}{3} \frac{p}{ks}}$$

$$\ge \frac{1}{4} \log \frac{\frac{1}{2} \log \frac{p}{k}}{7 \log \frac{p}{ks}}$$

$$\ge \frac{1}{4} \log \frac{\log \frac{p}{k}}{\log \frac{p}{ks}} - 1$$

This gives the induction step.

Theorem 42

Merging requires at least $\Omega(\log \log n)$ time on a CRCW PRAM with n processors.

Theorem 43

We can simulate a p-processor priority CRCW PRAM on a p-processor EREW PRAM with slowdown $O(\log p)$.

Theorem 44

We can simulate a p-processor priority CRCW PRAM on a $p \log p$ -processor common CRCW PRAM with slowdown O(1).

Theorem 45

We can simulate a p-processor priority CRCW PRAM on a p-processor common CRCW PRAM with slowdown $\mathcal{O}(\frac{\log p}{\log \log p})$.

Theorem 46

We can simulate a p-processor priority CRCW PRAM on a p-processor arbitrary CRCW PRAM with slowdown $\mathcal{O}(\log\log p)$.

Lower Bounds for the CREW PRAM

Ideal PRAM:

- every processor has unbounded local memory
- in each step a processor reads a global variable
- then it does some (unbounded) computation on its local memory
- then it writes a global variable

Lower Bounds for the CREW PRAM

Definition 47

An input index i affects a memory location M at time t on some input I if the content of M at time t differs between inputs I and I(i) (i-th bit flipped).

 $L(M, t, I) = \{i \mid i \text{ affects } M \text{ at time } t \text{ on input } I\}$

Lower Bounds for the CREW PRAM

Definition 48

An input index i affects a processor P at time t on some input I if the state of P at time t differs between inputs I and I(i) (i-th bit flipped).

 $K(P, t, I) = \{i \mid i \text{ affects } P \text{ at time } t \text{ on input } I\}$

Lower Bounds for the CREW PRAM

Lemma 49

If $i \in K(P, t, I)$ with t > 1 then either

- ▶ $i \in K(P, t 1, I)$, or
- ▶ P reads a global memory location M on input I at time t, and $i \in L(M, t-1, I)$.

Lower Bounds for the CREW PRAM

Lemma 50

If $i \in L(M, t, I)$ with t > 1 then either

- ▶ A processor writes into M at time t on input I and $i \in K(P, t, I)$, or
- No processor writes into M at time t on input I and
 - either $i \in L(M, t-1, I)$
 - or a processor P writes into M at time t on input I(i).

Let $k_0 = 0$, $\ell_0 = 1$ and define

$$k_{t+1} = k_t + \ell_t$$
 and $\ell_{t+1} = 3k_t + 4\ell_t$

Lemma 51

 $|K(P,t,I)| \le k_t$ and $|L(M,t,I)| \le \ell_t$ for any $t \ge 0$

base case (t = 0):

- No index can influence the local memory/state of a processor before the first step (hence $|K(P, 0, I)| = k_0 = 0$).
- Initially every index in the input affects exactly one memory location. Hence $|L(M,0,I)| = 1 = \ell_0$.

induction step $(t \rightarrow t + 1)$:

 $K(P, t+1, I) \subseteq K(P, t, I) \cup L(M, t, I)$, where M is the location read by P in step t+1.

Hence,

$$|K(P, t+1, I)| \le |K(P, t, I)| + |L(M, t, I)|$$

 $\le k_t + \ell_t$

induction step $(t \rightarrow t + 1)$:

For the bound on |L(M, t + 1, I)| we have two cases.

Case 1:

A processor P writes into location M at time t + 1 on input I.

Then,

$$\begin{split} |L(M,t+1,I)| &\leq |K(P,t+1,I)| \\ &\leq k_t + \ell_t \\ &\leq 3k_t + 4\ell_t = \ell_{t+1} \end{split}$$

Case 2:

No processor P writes into location M at time t+1 on input I.

An index i affects M at time t+1 iff i affects M at time t or some processor P writes into M at t+1 on I(i).

$$L(M,t+1,I) \subseteq L(M,t,I) \cup Y(M,t+1,I)$$

Y(M, t+1, I) is the set of indices u_j that cause some processor P_{w_j} to write into M at time t+1 on input I.

Y(M, t+1, I) is the set of indices u_j that cause some processor P_{w_j} to write into M at time t+1 on input I.

Fact:

For all pairs u_s , u_t with $P_{w_s} \neq P_{w_t}$ either $u_s \in K(P_{w_t}, t+1, I(u_t))$ or $u_t \in K(P_{w_s}, t+1, I(u_s))$.

Otherwise, P_{w_t} and P_{w_s} would both write into M at the same time on input $I(u_s)(u_t)$.

Let $U = \{u_1, ..., u_r\}$ denote all indices that cause some processor to write into M.

Let
$$V = \{(I(u_1), P_{w_1}), \dots\}.$$

We set up a bipartite graph between U and V, such that $(u_i, (I(u_j), P_{w_j})) \in E$ if u_i affects P_{w_j} at time t+1 on input $I(u_j)$.

Each vertex $(I(u_j), P_{w_j})$ has degree at most k_{t+1} as this is an upper bound on indices that can influence a processor P_{w_j} .

Hence, $|E| \leq r \cdot k_{t+1}$.

For an index u_j there can be at most k_{t+1} indices u_i with $P_{w_i} = P_{w_j}$.

Hence, there must be at least $\frac{1}{2}r(r-k_{t+1})$ pairs u_i,u_j with $P_{w_i} \neq P_{w_j}$.

Each pair introduces at least one edge.

Hence,

$$|E| \geq \frac{1}{2}r(r - k_{t+1})$$

This gives $r \leq 3k_{t+1} \leq 3k_t + 3\ell_t$

Recall that $L(M, t + 1, i) \subseteq L(M, t, i) \cup Y(M, t + 1, I)$

 $|L(M,t+1,i)| \leq 3k_t + 4\ell_t$

$$\begin{pmatrix} k_{t+1} \\ \ell_{t+1} \end{pmatrix} = \begin{pmatrix} 1 & 1 \\ 3 & 4 \end{pmatrix} \begin{pmatrix} k_t \\ \ell_t \end{pmatrix} \qquad \begin{pmatrix} k_0 \\ \ell_0 \end{pmatrix} = \begin{pmatrix} 0 \\ 1 \end{pmatrix}$$

Eigenvalues:

$$\lambda_1 = \frac{1}{2}(5 + \sqrt{21}) \text{ and } \lambda_2 = \frac{1}{2}(5 - \sqrt{21})$$

Eigenvectors:

$$v_1=egin{pmatrix}1\\-(1-\lambda_1)\end{pmatrix}$$
 and $v_2=egin{pmatrix}1\\-(1-\lambda_2)\end{pmatrix}$

$$v_1=\left(egin{array}{c}1\\rac{3}{2}+rac{1}{2}\sqrt{21}\end{array}
ight)$$
 and $v_2=\left(egin{array}{c}1\\rac{3}{2}-rac{1}{2}\sqrt{21}\end{array}
ight)$

$$v_1 = \begin{pmatrix} 1 \\ \frac{3}{2} + \frac{1}{2}\sqrt{21} \end{pmatrix}$$
 and $v_2 = \begin{pmatrix} 1 \\ \frac{3}{2} - \frac{1}{2}\sqrt{21} \end{pmatrix}$
$$\begin{pmatrix} k_0 \\ \ell_0 \end{pmatrix} = \begin{pmatrix} 0 \\ 1 \end{pmatrix} = \frac{1}{\sqrt{21}}(v_1 - v_2)$$

$$\begin{pmatrix} k_t \\ \ell_t \end{pmatrix} = \frac{1}{\sqrt{21}} \left(\lambda_1^t v_1 - \lambda_2^t v_2\right)$$

Solving the recurrence gives

$$k_t = \frac{\lambda_1^t}{\sqrt{21}} - \frac{\lambda_2^t}{\sqrt{21}}$$

$$\ell_t = \frac{3+\sqrt{21}}{2\sqrt{21}}\lambda_1^t + \frac{-3+\sqrt{21}}{2\sqrt{21}}\lambda_2^t$$
 with $\lambda_1 = \frac{1}{2}(5+\sqrt{21})$ and $\lambda_2 = \frac{1}{2}(5-\sqrt{21})$.

Theorem 52

The following problems require logarithmic time on a CREW PRAM.

- Sorting a sequence of x_1, \ldots, x_n with $x_i \in \{0, 1\}$
- Computing the maximum of n inputs
- Computing the sum $x_1 + \cdots + x_n$ with $x_i \in \{0, 1\}$

A Lower Bound for the EREW PRAM

Definition 53 (Zero Counting Problem)

Given a monotone binary sequence $x_1, x_2, ..., x_n$ determine the index i such that $x_i = 0$ and $x_{i+1} = 1$.

We show that this problem requires $\Omega(\log n - \log p)$ steps on a p-processor EREW PRAM.

Let I_i be the input with i zeros folled by n-i ones.

Index i affects processor P at time t if the state in step t is differs between I_{i-1} and I_i .

Index i affects location M at time t if the content of M after step t differs between inputs I_{i-1} and I_i .

Lemma 54

If $i \in K(P, t)$ then either

- $i \in K(P, t-1)$, or
- ▶ P reads some location M on input I_i (and, hence, also on I_{i-1}) at step t and $i \in L(M, t-1)$

Lemma 55

If $i \in L(M,t)$ then either

- $i \in L(M, t-1)$, or
- Some processor P writes M at step t on input I_i and $i \in K(P,t)$.
- Some processor P writes M at step t on input I_{i-1} and $i \in K(P,t)$.

Define

$$C(t) = \sum_{P} |K(P, t)| + \sum_{M} \max\{0, |L(M, t)| - 1\}$$

$$C(T) \ge n, C(0) = 0$$

Claim:

$$C(t) \le 6C(t-1) + 3|P|$$

This gives $C(T) \le \frac{6^T - 1}{5} 3|P|$ and hence $T = \Omega(\log n - \log |P|)$.

For an index i to newly appear in L(M,t) some processor must write into M on either input I_i or I_{i-1} .

Hence, any index in K(P,t) can at most generate two new indices in L(M,t).

This means that the number of new indices in any set L(M,t) (over all M) is at most

$$2\sum_{P}|K(P,t)|$$

Hence,

$$\sum_{M} |L(M,t)| \leq \sum_{M} |L(M,t-1)| + 2\sum_{P} |K(P,t)|$$

We can assume wlog. that $L(M, t - 1) \subseteq L(M, t)$. Then

$$\sum_{M} \max\{0, |L(M, t)| - 1\} \le \sum_{M} \max\{0, |L(M, t - 1)| - 1\} + 2\sum_{P} |K(P, t)|$$

For an index i to newly appear in K(P,t), P must read a memory location M with $i \in L(M,t)$ on input I_i (and also on input I_{i-1}).

Since we are in the EREW model at most one processor can do so in every step.

Let J(i,t) be memory locations read in step t on input I_i , and let $J_t = \bigcup_i J(i,t)$.

$$\sum_{P} |K(P,t)| \leq \sum_{P} |K(P,t-1)| + \sum_{M \in I_{t}} |L(M,t-1)|$$

Over all inputs I_i a processor can read at most |K(P, t-1)| + 1 different memory locations (why?).

Hence,

$$\begin{split} \sum_{P} |K(P,t)| &\leq \sum_{P} |K(P,t-1)| + \sum_{M \in J_t} |L(M,t-1)| \\ &\leq \sum_{P} |K(P,t-1)| + \sum_{M \in J_t} (|L(M,t-1)|-1) + J_t \\ &\leq 2 \sum_{P} |K(P,t-1)| + \sum_{M \in J_t} (|L(M,t-1)|-1) + |P| \\ &\leq 2 \sum_{P} |K(P,t-1)| + \sum_{M} \max\{0,|L(M,t-1)|-1\} + |P| \end{split}$$

Recall

$$\sum_{M} \max\{0, |L(M,t)| - 1\} \leq \sum_{M} \max\{0, |L(M,t-1)| - 1\} + 2\sum_{P} |K(P,t)|$$

This gives

$$\begin{split} & \sum_{P} K(P,t) + \sum_{M} \max\{0, |L(M,t)| - 1\} \\ & \leq 4 \sum_{M} \max\{0, |L(M,t-1)| - 1\} + 6 \sum_{P} |K(P,t-1)| + 3|P| \end{split}$$

Hence,

$$C(t) \le 6C(t-1) + 3|P|$$

Lower Bounds for CRCW PRAMS

Theorem 56

Let $f:\{0,1\}^n \to \{0,1\}$ be an arbitrary Boolean function. f can be computed in $\mathcal{O}(1)$ time on a common CRCW PRAM with $\leq n2^n$ processors.

Can we obtain non-constant lower bounds if we restrict the number of processors to be polynomial?

Boolean Circuits

- nodes are either AND, OR, or NOT gates or are special INPUT/OUTPUT nodes
- AND and OR gates have unbounded fan-in (indegree) and ounbounded fan-out (outdegree)
- NOT gates have unbounded fan-out
- INPUT nodes have indegree zero; OUTPUT nodes have outdegree zero
- size is the number of edges
- depth is the longest path from an input to an output

Theorem 57

Let $f: \{0,1\}^n \to \{0,1\}^m$ be a function with n inputs and $m \le n$ outputs, and circuit C computes f with depth D(n) and size S(n). Then f can be computed by a common CRCW PRAM in $\mathcal{O}(D(n))$ time using S(n) processors.

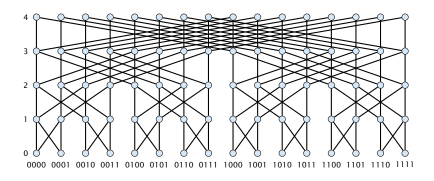
Given a family $\{C_n\}$ of circuits we may not be able to compute the corresponding family of functions on a CRCW PRAM.

Definition 58

A family $\{C_n\}$ of circuits is logspace uniform if there exists a deterministic Turing machine M s.t

- M runs in logarithmic space.
- For all n, M outputs C_n on input 1^n .

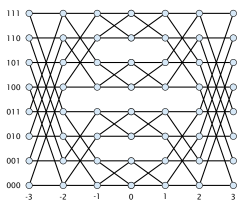
Bufferfly Network BF(*d*)



- ▶ node set $V = \{(\ell, \bar{x}) \mid \bar{x} \in [2]^d, \ell \in [d+1]\}$, where $\bar{x} = x_0 x_1 \dots x_{d-1}$ is a bit-string of length d
- edge set $E = \{\{(\ell, \bar{x}), (\ell+1, \bar{x}')\} \mid \ell \in [d], \bar{x} \in [2]^d, x_i' = x_i \text{ for } i \neq \ell\}$

Sometimes the first and last level are identified.

Beneš Network

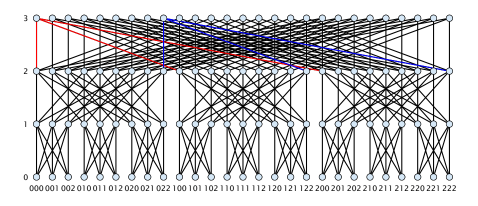


- node set $V = \{(\ell, \bar{x}) \mid \bar{x} \in [2]^d, \ell \in \{-d, ..., d\}\}$
- edge set

$$E = \{ \{ (\ell, \bar{x}), (\ell+1, \bar{x}') \} \mid \ell \in [d], \bar{x} \in [2]^d, x_i' = x_i \text{ for } i \neq \ell \}$$

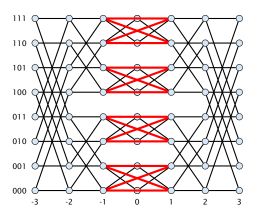
$$\cup \{ \{ (-\ell, \bar{x}), (\ell-1, \bar{x}') \} \mid \ell \in [d], \bar{x} \in [2]^d, x_i' = x_i \text{ for } i \neq \ell \}$$

n-ary Bufferfly Network BF(n, d)



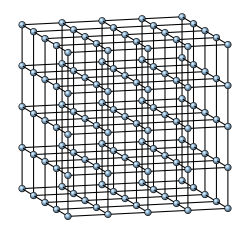
- ▶ node set $V = \{(\ell, \bar{x}) \mid \bar{x} \in [n]^d, \ell \in [d+1]\}$, where $\bar{x} = x_0 x_1 \dots x_{d-1}$ is a bit-string of length d
- edge set $E = \{\{(\ell, \bar{x}), (\ell + 1, \bar{x}')\} \mid \ell \in [d], \bar{x} \in [n]^d, x_i' = x_i \text{ for } i \neq \ell\}$

Permutation Network PN(n, d)



- ► There is an *n*-ary version of the Benes network (2 *n*-ary butterflies glued at level 0).
- ▶ identifying levels 0 and 1 (or 0 and -1) gives PN(n, d).

The d-dimensional mesh M(n, d)



- ▶ node set $V = [n]^d$
- edge set $E = \{\{(x_0, ..., x_i, ..., x_{d-1}), (x_0, ..., x_i + 1, ..., x_{d-1})\} \mid x_s \in [n] \text{ for } s \in [d] \setminus \{i\}, x_i \in [n-1]\}$

Remarks

M(2, d) is also called d-dimensional hypercube.

M(n, 1) is also called linear array of length n.

Permutation Routing

Lemma 59

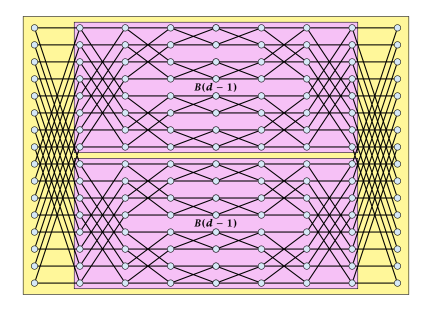
On the linear array M(n, 1) any permutation can be routed online in 2n steps with buffersize 3.

Permutation Routing

Lemma 60

On the Beneš network any permutation can be routed offline in 2d steps between the sources level (+d) and target level (-d).

Recursive Beneš Network



Permutation Routing

base case d = 0 trivial

induction step $d \rightarrow d + 1$

- ► The packets that start at (\bar{a}, d) and $(\bar{a}(d), d)$ have to be sent into different sub-networks.
- ▶ The packets that end at $(\bar{a}, -d)$ and $(\bar{a}(d), -d)$ have to come out of different sub-networks.

We can generate a graph on the set of packets.

- Every packet has an incident source edge (connecting it to the conflicting start packet)
- Every packet has an incident target edge (connecting it to the conflicting packet at its target)
- ► This clearly gives a bipartite graph; Coloring this graph tells us which packet to send into which sub-network.

Permutation Routing on the n-ary Beneš Network

Instead of two we have n sub-networks B(n, d-1).

All packets starting at positions $\{(x_0,\ldots,x_{d-2},x_{d-1},d)\mid x_{d-1}\in[n]\}$ have to be send to different sub-networks.

All packets ending at positions $\{(x_0,\ldots,x_{d-2},x_{d-1},d)\mid x_{d-1}\in[n]\}$ have to come from different sub-networks.

The conflict graph is an n-uniform 2-regular hypergraph.

We can color such a graph with n colors such that no two nodes in a hyperedge share a color.

This gives the routing.

On a d-dimensional mesh with sidelength n we can route any permutation (offline) in 4dn steps.

We can simulate the algorithm for the n-ary Beneš Network.

Each step can be simulated by routing on disjoint linear arrays. This takes at most 2n steps.

We simulate the behaviour of the Beneš network on the $n\text{-}\mathrm{dimensional}$ mesh.

In round $r \in \{-d, \ldots, -1, 0, 1, \ldots, d-1\}$ we simulate the step of sending from level r of the Beneš network to level r+1.

Each node $\bar{x} \in [n]^d$ of the mesh simulates the node (r, \bar{x}) .

Hence, if in the Beneš network we send from (r, \bar{x}) to $(r+1, \bar{x}')$ we have to send from \bar{x} to \bar{x}' in the mesh.

All communication is performed along linear arrays. In round r<0 the linear arrays along dimension -r-1 (recall that dimensions are numbered from 0 to d-1) are used

$$\bar{X}_{d-1} \dots \bar{X}_{-r} \alpha \bar{X}_{-r-2} \dots \bar{X}_0$$

In rounds $r \ge 0$ linear arrays along dimension r are used.

Hence, we can perform a round in O(n) steps.

We can route any permutation on the Beneš network in $\mathcal{O}(d)$ steps with constant buffer size.

The same is true for the butterfly network.

The nodes are of the form (ℓ, \bar{x}) , $\bar{x} \in [n]^d$, $\ell \in -d, ..., d$.

We can view nodes with same first coordinate forming columns and nodes with the same second coordinate as forming rows. This gives rows of length 2d - 1 and columns of length n^d .

We route in 3 phases:

- 1. Permute packets along the rows such that afterwards no column contains packets that have the same target row. $\mathcal{O}(d)$ steps.
- **2.** We can use pipeling to permute **every** column, so that afterwards every packet is in its target row. O(2d + 2d) steps.
- **3.** Every packet is in its target row. Permute packets to their right destinations. O(d) steps.

We can do offline permutation routing of (partial) permutations in 2d steps on the hypercube.

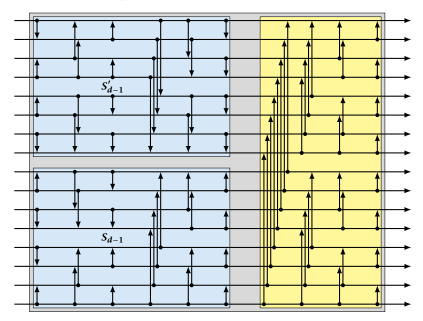
Lemma 64

We can sort on the hypercube M(2,d) in $O(d^2)$ steps.

Lemma 65

We can do online permutation routing of permutations in $\mathcal{O}(d^2)$ steps on the hypercube.

Bitonic Sorter S_d



ASCEND/DESCEND Programs

Algorithm 11 ASCEND(procedure *oper*)

- 1: **for** dim = 0 **to** d 1
- 2: for all $\bar{a} \in [2]^d$ pardo
- 3: $\operatorname{oper}(\bar{a}, \bar{a}(\operatorname{dim}), \operatorname{dim})$

Algorithm 11 DESCEND(procedure *oper*)

- 1: **for** dim = d 1 **to** 0
- 2: for all $\bar{a} \in [2]^d$ pardo
- 3: oper(\bar{a} , \bar{a} (dim), dim)

oper should only depend on the dimension and on values stored in the respective processor pair $(\bar{a}, \bar{a}(dim), V[\bar{a}], V[\bar{a}(dim)])$.

oper should take constant time.

Algorithm 11 oper $(a, a', dim, T_a, T_{a'})$ 1: if $a_{dim}, \dots, a_0 = 0^{dim+1}$ then 2: $T_a = \min\{T_a, T_{a'}\}$

Performing an ASCEND run with this operation computes the minimum in processor 0.

We can sort on M(2, d) by using d DESCEND runs.

We can do offline permutation routing by using a DESCEND run followed by an ASCEND run.

We can perform an ASCEND/DESCEND run on a linear array $M(2^d,1)$ in $\mathcal{O}(2^d)$ steps.

The CCC network is obtained from a hypercube by replacing every node by a cycle of degree d.

- nodes $\{(\ell, \bar{x}) \mid \bar{x} \in [2]^d, \ell \in [d]\}$
- edges $\{\{(\ell, \bar{x}), (\ell, \bar{x}(\ell)) \mid x \in [2]^d, \ell \in [d]\}$

constand degree

Let $d = 2^k$. An ASCEND run of a hypercube M(2, d + k) can be simulated on CCC(d) in O(d) steps.

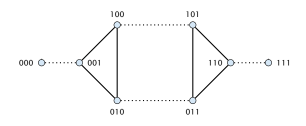
The shuffle exchange network SE(d) is defined as follows

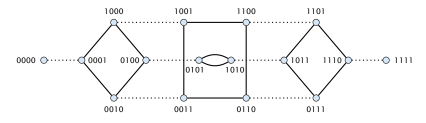
- nodes: $V = [2]^d$
- edges: $E = \left\{ \{ x \bar{\alpha}, \bar{\alpha} x \} \mid x \in [2], \bar{\alpha} \in [2]^{d-1} \right\} \cup \left\{ \{ \bar{\alpha} 0, \bar{\alpha} 1 \} \mid \bar{\alpha} \in [2]^{d-1} \right\}$

constand degree

Edges of the first type are called shuffle edges. Edges of the second type are called exchange edges

Shuffle Exchange Networks





We can perform an ASCEND run of M(2,d) on SE(d) in $\mathcal{O}(d)$ steps.

For the following observations we need to make the definition of parallel computer networks more precise.

Each node of a given network corresponds to a processor/RAM.

In addition each processor has a read register and a write register.

In one (synchronous) step each neighbour of a processor P_i can write into P_i 's write register or can read from P_i 's read register.

Usually we assume that proper care has to be taken to avoid concurrent reads and concurrent writes from/to the same register.

Definition 68

A configuration C_i of processor P_i is the complete description of the state of P_i including local memory, program counter, read-register, write-register, etc.

Suppose a machine M is in configuration (C_0, \ldots, C_{p-1}) , performs t synchronous steps, and is then in configuration $C = (C'_0, \ldots, C'_{p-1})$.

 C'_i is called the *t*-th successor configuration of *C* for processor *i*.

Definition 69

Let $C=(C_0,\ldots,C_{p-1})$ a configuration of M. A machine M' with $q\geq p$ processors weakly simulates t steps of M with slowdown k if

- ▶ in the beginning there are p non-empty processors sets $A_0, \ldots, A_{p-1} \subseteq M'$ so that all processors in A_i know C_i ;
- ▶ after at most $k \cdot t$ steps of M' there is a processor $Q^{(i)}$ that knows the t-th successors configuration of C for processor P_i .

Definition 70

M' simulates M with slowdown k if

- ightharpoonup M' weakly simulates machine M with slowdown k
- ▶ and **every** processor in A_i knows the t-th successor configuration of C for processor P_i .

We have seen how to simulate an ASCEND/DESCEND run of the hypercube M(2, d + k) on CCC(d) with $d = 2^k$ in O(d) steps.

Hence, we can simulate d+k steps (one ASCEND run) of the hypercube in $\mathcal{O}(d)$ steps. This means slowdown $\mathcal{O}(1)$.

Suppose a network S with n processors can route any permutation in time $\mathcal{O}(t(n))$. Then S can simulate any constant degree network M with at most n vertices with slowdown $\mathcal{O}(t(n))$.

Map the vertices of M to vertices of S in an arbitrary way.

Color the edges of M with $\Delta+1$ colors, where $\Delta=\mathcal{O}(1)$ denotes the maximum degree.

Each color gives rise to a permutation.

We can route this permutation in S in t(n) steps.

Hence, we can perform the required communication for one step of M by routing $\Delta + 1$ permutations in S. This takes time t(n).

A processor of ${\cal M}$ is simulated by the same processor of ${\cal S}$ throughout the simulation.

Suppose a network S with n processors can sort n numbers in time $\mathcal{O}(t(n))$. Then S can simulate any network M with at most n vertices with slowdown $\mathcal{O}(t(n))$.

There is a constant degree network on $\mathcal{O}(n^{1+\epsilon})$ nodes that can simulate any constant degree network with slowdown $\mathcal{O}(1)$.

Suppose we allow concurrent reads, this means in every step all neighbours of a processor P_i can read P_i 's read register.

Lemma 74

A constant degree network M that can simulate any n-node network has slowdown $\Omega(\log n)$ (independent of the size of M).

We show the lemma for the following type of simulation.

- ► There are representative sets A_i^t for every step t that specify which processors of M simulate processor P_i in step t (know the configuration of P_i after the t-th step).
- The representative sets for different processors are disjoint.
- ▶ for all $i \in \{1, ..., n\}$ and steps $t, A_i^t \neq \emptyset$.

This is a step-by-step simulation.

Suppose processor P_i reads from processor P_{j_i} in step t.

Every processor $Q \in M$ with $Q \in A_i^{t+1}$ must have a path to a processor $Q' \in A_i^t$ and to $Q'' \in A_{j_i}^t$.

Let k_t be the largest distance (maximized over all i, j_i).

Then the simulation of step t takes time at least k_t .

The slowdown is at least

$$k = \frac{1}{\ell} \sum_{t=1}^{\ell} k_t$$

We show

- ▶ The simulation of a step takes at least time $y \log n$, or
- the size of the representative sets shrinks by a lot

$$\sum_{i} |A_i^{t+1}| \le \frac{1}{n^{\epsilon}} \sum_{i} |A_i^t|$$

Suppose there is no pair (i, j) such that i reading from j requires time $\gamma \log n$.

- ► For every i the set $\Gamma_{2k}(A_i)$ contains a node from A_j .
- ▶ Hence, there must exist a j_i such that $\Gamma_{2k}(A_i)$ contains at most

$$|C_{j_i}| := \frac{|A_i| \cdot c^{2k}}{n-1} \le \frac{|A_i| \cdot c^{3k}}{n}$$
.

processors from $|A_{j_i}|$

If we choose that i reads from j_i we get

$$|A'_{i}| \le |C_{j_{i}}| \cdot c^{k}$$

$$\le c^{k} \cdot \frac{|A_{i}| \cdot c^{3k}}{n}$$

$$= \frac{1}{n} |A_{i}| \cdot c^{4k}$$

Choosing $k = \Theta(\log n)$ gives that this is at most $|A_i|/n^{\epsilon}$.

Let ℓ be the total number of steps and s be the number of short steps when $k_t < \gamma \log n$.

In a step of time k_t a representative set can at most increase by c^{k_t+1} .

Let h_ℓ denote the number of representatives after step ℓ .

$$n \le h_{\ell} \le h_0 \left(\frac{1}{n^{\epsilon}}\right)^s \prod_{t \in \text{long}} c^{k_t + 1} \le \frac{n}{n^{\epsilon s}} \cdot c^{\ell + \sum_t k_t}$$

If $\sum_t k_t \ge \ell(\frac{\epsilon}{2} \log_c n - 1)$, we are done. Otw.

$$n \le n^{1-\epsilon s + \ell \frac{\epsilon}{2}}$$

This gives $s \le \ell/2$.

Hence, at most 50% of the steps are short.

Lemma 75

A permutation on an $n \times n$ -mesh can be routed online in $\mathcal{O}(n)$ steps.

Definition 76 (Oblivious Routing)

Specify a path-system \mathcal{W} with a path $P_{u,v}$ between u and v for every pair $\{u,v\} \in V \times V$.

A packet with source u and destination v moves along path $P_{u,v}$.

Definition 77 (Oblivious Routing)

Specify a path-system $\mathcal W$ with a path $P_{u,v}$ between u and v for every pair $\{u,v\}\in V\times V$.

Definition 78 (node congestion)

For a given path-system the node congestion is the maximum number of path that go through any node $v \in V$.

Definition 79 (edge congestion)

For a given path-system the edge congestion is the maximum number of path that go through any edge $e \in E$.

Definition 80 (dilation)

For a given path system the dilation is the maximum length of a path.

Lemma 81

Any oblivious routing protocol requires at least $\max\{C_f, D_f\}$ steps, where C_f and D_f , are the congestion and dilation, respectively, of the path-system used. (node congestion or edge congestion depending on the communication model)

Lemma 82

Any reasonable oblivious routing protocol requires at most $\mathcal{O}(D_f \cdot C_f)$ steps (unbounded buffers).

Theorem 83 (Borodin, Hopcroft)

For any path system W there exists a permutation $\pi:V\to V$ and an edge $e\in E$ such that at least $\Omega(\sqrt{n}/\Delta)$ of the paths go through e.

Let $\mathcal{W}_v = \{P_{v,u} \mid u \in V\}.$

We say that an edge e is z-popular for v if at least z paths from \mathcal{W}_v contain e.

For any node v there are many edges that are are quite popular for v.

 $|V| \times |E|$ -matrix A(z):

$$A_{v,e}(z) = \begin{cases} 1 & e \text{ is } z\text{-popular for } v \\ 0 & \text{otherwise} \end{cases}$$

Define

$$A_{v}(z) = \sum_{e} A_{v,e}(z)$$

•

$$A_e(z) = \sum_{v} A_{v,e}(z)$$

Lemma 84

Let $z \leq \frac{n-1}{\Delta}$.

For every node $v \in V$ there exist at least $\frac{n}{2\Delta z}$ edges that are z popular for v. This means

$$A_v(z) \ge \frac{n}{2\Delta z}$$

Lemma 85

There exists an edge e' that is z-popular for at least z nodes with $z = \Omega(\sqrt{n}\Delta)$.

$$\sum_{e} A_{e}(z) = \sum_{v} A_{v}(z) \ge \frac{n^{2}}{2\Delta z}$$

There must exist an edge e'

$$A_{e'}(z) \ge \left\lceil \frac{n^2}{|E| \cdot 2\Delta z} \right\rceil \ge \left\lceil \frac{n}{2\Delta^2 z} \right\rceil$$

where the last step follows from $|E| \leq \Delta n$.

We choose z such that $z = \frac{n}{2\Delta^2 z}$ (i.e., $z = \sqrt{n}/(\sqrt{2}\Delta)$).

This means e' is $\lceil z \rceil$ -popular for $\lceil z \rceil$ nodes.

We can construct a permutation such that z paths go through e'.

Deterministic oblivious routing may perform very poorly.

What happens if we have a random routing problem in a butterfly?

Suppose every source on level 0 has p packets, that are routed to random destinations.

How many packets go over node v on level i?

From v we can reach $2^d/2^i$ different targets.

Hence,

$$\Pr[\text{packet goes over } v] \le \frac{2^{d-i}}{2^d} = \frac{1}{2^i}$$

Expected number of packets:

E[packets over
$$v$$
] = $p \cdot 2^i \cdot \frac{1}{2^i} = p$

since only $p2^i$ packets can reach v.

But this is trivial.

What is the probability that at least r packets go through v.

$$\begin{aligned} \Pr[\text{at least } r \text{ path through } v] &\leq \binom{p \cdot 2^i}{r} \cdot \left(\frac{1}{2^i}\right)^r \\ &\leq \left(\frac{p2^i \cdot e}{r}\right)^r \cdot \left(\frac{1}{2^i}\right) \\ &= \left(\frac{pe}{r}\right)^r \end{aligned}$$

 $\Pr[\text{there exists a node } v \text{ sucht that at least } r \text{ path through } v]$

$$\leq d2^d \cdot \left(\frac{pe}{r}\right)^r$$

 $\Pr[\text{there exists a node } v \text{ sucht that at least } r \text{ path through } v]$

$$\leq d2^d \cdot \left(\frac{pe}{r}\right)^r$$

Choose r as $2ep + (\ell + 1)d + \log d = \mathcal{O}(p + \log N)$, where N is number of sources in BF(d).

 $\Pr[\text{exists node } v \text{ with more than } r \text{ paths over } v] \leq \frac{1}{N\ell}$

Scheduling Packets

Assume that in every round a node may forward at most one packet but may receive up to two.

We select a random rank $R_p \in [k]$. Whenever, we forward a packet we choose the packet with smaller rank. Ties are broken according to packet id.

Random Rank Protocol

Definition 86 (Delay Sequence of length *s***)**

- \triangleright delay path \mathcal{W}
- ▶ lengths $\ell_0, \ell_1, \dots, \ell_s$, with $\ell_0 \ge 1, \ell_1, \dots, \ell_s \ge 0$ lengths of delay-free sub-paths
- ightharpoonup collision nodes $v_0, v_1, \dots, v_s, v_{s+1}$
- ightharpoonup collision packets P_0, \ldots, P_s

Properties

- ▶ $rank(P_0) \ge rank(P_1) \ge \cdots \ge rank(P_s)$
- $\sum_{i=0}^{s} \ell_i = d$
- if the routing takes d + s steps than the delay sequence has length s

Definition 87 (Formal Delay Sequence)

- ightharpoonup a path $\mathcal W$ of length d from a source to a target
- s integers $\ell_0 \ge 1$, $\ell_1, \dots, \ell_s \ge 0$ and $\sum_{i=0}^s \ell_i = d$
- nodes $v_0, \dots v_s, v_{s+1}$ on $\mathcal W$ with v_i being on level $d-\ell_0-\dots-\ell_{i-1}$
- ▶ s + 1 packets $P_0, ..., P_s$, where P_i is a packet with path through v_i and v_{i-1}
- ▶ numbers $R_s \le R_{s-1} \le \cdots \le R_0$

We say a formal delay sequence is active if $rank(P_i) = k_i$ holds for all i.

Let N_s be the number of formal delay sequences of length at most s. Then

$$\Pr[\text{routing needs at least } d + s \text{ steps}] \le \frac{N_s}{k^{s+1}}$$

Lemma 88

$$N_{s} \le \left(\frac{2eC(s+k)}{s+1}\right)^{s+1}$$

- there are N^2 ways to choose \mathcal{W}
- ▶ there are $\binom{s+d-1}{s}$ ways to choose ℓ_i 's with $\sum_{i=0}^{s} \ell_i = d$
- the collision nodes are fixed
- ▶ there are at most C^{s+1} ways to choose the collision packets where C is the node congestion
- ► there are at most $\binom{s+k}{s+1}$ ways to choose $0 \le k_s \le \cdots \le k_0 < k$

Hence the probability that the routing takes more than d + s steps is at most

$$N^3 \cdot \left(\frac{2e \cdot C \cdot (s+k)}{(s+1)k}\right)^{s+1}$$

We choose $s = 8eC - 1 + (\ell + 3)d$ and k = s + 1. This gives that the probability is at most $\frac{1}{N\ell}$.

- With probability $1 \frac{1}{N^{\ell_1}}$ the random routing problem has congestion at most $\mathcal{O}(p + \ell_1 d)$.
- With probability $1 \frac{1}{N^{\ell_2}}$ the packet scheduling finishes in at most $\mathcal{O}(C + \ell_2 d)$ steps.

Hence, with high probability routing random problems with p packets per source in a butterfly requires only $\mathcal{O}(p+d)$ steps.

What do we do for arbitrary routing problems?

Where did the scheduling analysis use the butterfly?

We only used

- all routing paths are of the same length d
- there are a polynomial number of delay paths

Choose paths as follows:

- route from source to random destination on target level
- route to real target column (albeit on source level)
- route to target

All phases run in time $\mathcal{O}(p+d)$ with high probability.

Multicommodity Flow Problem

- undirected (weighted) graph G = (V, E, c)
- ▶ commodities (s_i, t_i) , $i \in \{1, ..., k\}$
- ▶ a multicommodity flow is a flow $f: E \times \{1, ..., k\} \rightarrow \mathbb{R}^+$
 - ▶ for all edges $e \in E$: $\sum_i f_i(e) \le c(e)$
 - for all nodes $v \in V \setminus \{s_i, t_i\}$: $\sum_{u:(u,v)\in F} f_i((u,v)) = \sum_{w:(v,w)\in F} f_i((v,w))$

Goal A (Maximum Multicommodity Flow) maximize $\sum_i \sum_{e=(s_i,x)\in E} f_i(e)$

Goal B (Maximum Concurrent Multicommodity Flow) maximize $\min_i \sum_{e=(s_i,x)\in E} f_i(e)/d_i$ (throughput fraction), where d_i is demand for commodity i

A Balanced Multicommodity Flow Problem is a concurrent multicommodity flow problem in which incoming and outgoing flow is equal to

$$c(v) = \sum_{e=(v,x)\in E} c(e)$$

For a multicommodity flow S we assume that we have a decomposition of the flow(s) into flow-paths.

We use C(S) to denote the congestion of the flow problem (inverse of throughput fraction), and D(S) the length of the longest routing path.

For a network G = (V, E, c) we define the characteristic flow problem via

• demands $d_{u,v} = \frac{c(u)c(v)}{c(V)}$

Suppose the characteristic flow problem has a solution S with $C(S) \le F$ and $D(S) \le F$.

Definition 89

A (randomized) oblivious routing scheme is given by a path system $\mathcal P$ and a weight function w such that

$$\sum_{p\in\mathcal{P}_{s,t}}w(p)=1$$

Construct an oblivious routing scheme from S as follows:

• let $f_{x,y}$ be the flow between x and y in S

$$f_{x,y} \ge d_{x,y}/C(S) \ge d_{x,y}/F = \frac{1}{F} \frac{c(x)c(y)}{c(V)}$$

• for $p \in \mathcal{P}_{x,y}$ set $w(p) = f_p/f_{x,y}$

gives an oblivious routing scheme.

We apply this routing scheme twice:

- first choose a path from $\mathcal{P}_{s,v}$, where v is chosen uniformly according to c(v)/c(V)
- then choose path according to $\mathcal{P}_{v,t}$

If the input flow problem/packet routing problem is balanced doing this randomization results in flow solution S (twice).

Hence, we have an oblivious scheme with congestion and dilation at most 2F for (balanced inputs).

Example: hypercube.

Oblivious Routing for the Mesh

We can route any permutation on an $n \times n$ mesh in $\mathcal{O}(n)$ steps, by x-y routing. Actually $\mathcal{O}(d)$ steps where d is the largest distance between a source-target pair.

What happens if we do not have a permutation?

x-y routing may generate large congestion if some pairs have a lot of packets.

Valiants trick may create a large dilation.

Let for a multicommodity flow problem P $C_{\mathrm{opt}}(P)$ be the optimum congestion, and $D_{\mathrm{opt}}(P)$ be the optimum dilation (by perhaps different flow solutions).

Lemma 90

There is an oblivious routing scheme for the mesh that obtains a flow solution S with $C(S) = \mathcal{O}(C_{\text{opt}}(P)\log n)$ and $D(S) = \mathcal{O}(D_{\text{opt}}(P))$.

Lemma 91

For any oblivious routing scheme on the mesh there is a demand *P* such that routing *P* will give congestion $\Omega(\log n \cdot C_{\text{opt}})$.