Efficient Parallel Algorithms

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- 2 Parallel computation models
- 3 Basic parallel algorithms
- 4 Further parallel algorithms
- 5 Parallel matrix algorithms
- 6 Parallel graph algorithms



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Model: abstraction of reality allowing qualitative and quantitative reasoning

Examples:

- atom
- biological cell
- galaxy
- Kepler's universe
- Newton's universe
- Einstein's universe
- . . .

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Computation model: abstract computing device to reason about computations and algorithms

Examples:

- scales+weights (for "counterfeit coin" problems)
- Turing machine
- von Neumann machine ("ordinary computer")
- JVM
- quantum computer
- . . .

Computation: input \rightarrow (computation steps) \rightarrow output

Algorithm: a finite description of a (usually infinite) set of computations on different inputs

Assumes a specific computation model and input/output encoding

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Similarly for other resources (e.g. memory, communication)

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Computation models and algorithms

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 $f(n) \ge 0 \quad n \to \infty$

Asymptotic growth classes relative to f: O(f), o(f), $\Omega(f)$, $\omega(f)$, $\Theta(f)$

Computation models and algorithms

 $f(n), g(n) \ge 0$ $n \to \infty$ g = O(f): "g grows at the same rate or slower than f"...

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$$\begin{split} f(n), g(n) &\geq 0 \quad n \to \infty \\ g &= O(f): \text{ "g grows at the same rate or slower than } f'' \dots \\ g &= O(f) \iff \exists C : \exists n_0 : \forall n \geq n_0 : g(n) \leq C \cdot f(n) \end{split}$$

In words: we can scale f up by a specific (possibly large) constant, so that f will eventually overtake and stay above g

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In words: even if we scale f down by any (however small) constant, f will still eventually overtake and stay above g

Overtaking point depends on the constant!

Exercise: $\exists n_0 : \forall c : \forall n \ge n_0 : g(n) \le c \cdot f(n)$ — what does this say?

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 $g = \Omega(f)$: "g grows at the same rate or faster than f" $g = \omega(f)$: "g grows (strictly) faster than f" $g = \Omega(f)$ iff f = O(g) $g = \omega(f)$ iff f = o(g)

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Computation models and algorithms

 $f(n), g(n) \ge 0$ $n \to \infty$ The maximum rule: $f + g = \Theta(\max(f, g))$

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Computation models and algorithms

 $f(n), g(n) \ge 0$ $n \to \infty$ The maximum rule: $f + g = \Theta(\max(f, g))$ Proof:

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Computation models and algorithms

 $f(n), g(n) \ge 0 \quad n \to \infty$ The maximum rule: $f + g = \Theta(\max(f, g))$ Proof: for all *n*, we have $\max(f(n) + g(n)) \le f(n) + g(n) \le 2\max(f(n) + g(n))$

Example usage: sorting an array of size n

All good comparison-based sorting algorithms run in time $O(n \log n)$

If only pairwise comparisons between elements are allowed, no algorithm can run faster than $\Omega(n \log n)$

Hence, comparison-based sorting has complexity $\Theta(n \log n)$

If we are not restricted to just making comparisons, we can often sort in time $o(n \log n)$, or even O(n)

Example usage: multiplying $n \times n$ matrices

All good algorithms run in time $O(n^3)$, where n is matrix size

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Hence, $(+, \times)$ matrix multiplication has complexity $\Theta(n^3)$

If subtraction is allowed, everything changes! The best known matrix multiplication algorithm (with subtraction) runs in time $O(n^{2.373})$

It is conjectured that $O(n^{2+\epsilon})$ for any $\epsilon > 0$ is possible – open problem! Matrix multiplication cannot run faster than $\Omega(n^2 \log n)$ even with subtraction (under some natural assumptions)

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Computation models and algorithms

Algorithm complexity depends on the model

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E.g. sorting *n* items:

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- E.g. factoring large numbers:
 - hard in a von Neumann-type (standard) model
 - not so hard on a quantum computer
- E.g. deciding if a program halts on a given input:
 - impossible in a standard (or even quantum) model
 - can be added to the standard model as an oracle, to create a more powerful model

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The circuit model

Basic special-purpose parallel model: a circuit

$$a^2 + 2ab + b^2$$
$$a^2 - b^2$$



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The circuit model

Basic special-purpose parallel model: a circuit

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Directed acyclic graph (dag), fixed number of inputs/outputs Models oblivious computation: control sequence independent of the input

The circuit model

Basic special-purpose parallel model: a circuit

 $a^2 + 2ab + b^2$ $a^2 - b^2$



Directed acyclic graph (dag), fixed number of inputs/outputs Models oblivious computation: control sequence independent of the input Computation on varying number of inputs: an (infinite) circuit family May or may not admit a finite description (= algorithm)

The circuit model

In a circuit family, node indegree/outdegree may be bounded (by a constant), or unbounded: e.g. two-argument vs *n*-argument sum Elementary operations:

- arithmetic/Boolean/comparison
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- *size* = number of nodes
- *depth* = max path length from input to output

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Other uses of circuits:

- arbitrary (non-oblivious) computation can be thought of as a circuit that is not given in advance, but revealed gradually
- timed circuits with feedback: systolic arrays

The comparison network model

A comparison network is a circuit of comparator nodes

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Input/output: sequences of equal length, taken from a totally ordered set

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The comparison network model

A merging network is a comparison network that takes two sorted input sequences of length n', n'', and produces a sorted output sequence of length n = n' + n''

A sorting network is a comparison network that takes an arbitrary input sequence, and produces a sorted output sequence

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General merging: O(n) comparisons, non-oblivious

General sorting: $O(n \log n)$ comparisons by mergesort, non-oblivious

What is the complexity of oblivious sorting?

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Naive sorting networks

BUBBLE-SORT(n) size $n(n-1)/2 = O(n^2)$ depth 2n - 3 = O(n)



Image: A matrix

Naive sorting networks

BUBBLE-SORT(n) size $n(n-1)/2 = O(n^2)$ depth 2n - 3 = O(n)

BUBBLE-SORT(8)

size 28

depth 13



Naive sorting networks

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Naive sorting networks

INSERTION-SORT(n) size $n(n-1)/2 = O(n^2)$ depth 2n - 3 = O(n)

INSERTION-SORT(8)

size 28

depth 13

Identical to BUBBLE-SORT!



The zero-one principle

Zero-one principle: A comparison network is sorting, if and only if it sorts all input sequences of 0s and 1s



The zero-one principle

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Proof.

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Proof. "Only if": trivial. "If": by contradiction.

Assume a given network does not sort input $x = \langle x_1, \ldots, x_n \rangle$

 $\langle x_1, \ldots, x_n \rangle \mapsto \langle y_1, \ldots, y_n \rangle \qquad \exists k, l : k < l : y_k > y_l$

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$$\begin{aligned} \langle x_1, \dots, x_n \rangle &\mapsto \langle y_1, \dots, y_n \rangle & \exists k, l : k < l : y_k > y_l \\ \text{Let } X_i &= \begin{cases} 0 & \text{if } x_i < y_k \\ 1 & \text{if } x_i \ge y_k \end{cases} \text{ and run the network on input } X = \langle X_1, \dots, X_n \rangle \end{aligned}$$

For all i, j we have $x_i \le x_j \Rightarrow X_i \le X_j$, therefore each X_i follows the same path through the network as x_i

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$$\langle X_1,\ldots,X_n\rangle\mapsto\langle Y_1,\ldots,Y_n\rangle$$
 $Y_k=1>0=Y_l$

We have k < l but $Y_k > Y_l$, so the network does not sort 0s and 1s

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The zero-one principle

The zero-one principle applies to sorting, merging and other comparison problems (e.g. selection)



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It allows one to test:

- a sorting network by checking only 2^n input sequences, instead of a much larger number $n! = (1 + o(1))(2\pi n)^{1/2} \cdot (n/e)^n$
- a merging network by checking only $(n'+1) \cdot (n''+1)$ pairs of input sequences, instead of a (typically) very much larger number $\binom{n}{n'} = \binom{n}{n''}$, e.g. for n = 2n' = 2n'': $\binom{n}{n'} = (1 + o(1))(\pi n/2)^{-1/2} \cdot 2^n$

General merging: O(n) comparisons, non-oblivious How fast can we merge obliviously?

Image: A matrix

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General merging: O(n) comparisons, non-oblivious

How fast can we merge obliviously?

$$\langle x_1 \leq \cdots \leq x_{n'} \rangle, \langle y_1 \leq \cdots \leq y_{n''} \rangle \mapsto \langle z_1 \leq \cdots \leq z_n \rangle$$

Odd-even merging

When n' = n'' = 1 compare (x_1, y_1) , otherwise by recursion:

- merge $\langle x_1, x_3, \dots \rangle, \langle y_1, y_3, \dots \rangle \mapsto \langle u_1 \leq u_2 \leq \dots \leq u_{\lceil n'/2 \rceil + \lceil n''/2 \rceil} \rangle$
- merge $\langle x_2, x_4, \dots \rangle, \langle y_2, y_4, \dots \rangle \mapsto \langle v_1 \leq v_2 \leq \dots \leq v_{\lfloor n'/2 \rfloor + \lfloor n''/2 \rfloor} \rangle$
- compare pairwise: (*u*₂, *v*₁), (*u*₃, *v*₂), ...

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- compare pairwise: (*u*₂, *v*₁), (*u*₃, *v*₂), . . .

 $size(OEM(n', n'')) \le 2 \cdot size(OEM(n'/2, n''/2)) + O(n) = O(n \log n)$ $depth(OEM(n', n'')) \le depth(OEM(n'/2, n''/2)) + 1 = O(\log n)$

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OEM(n', n'')size $O(n \log n)$ depth $O(\log n)$



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OEM(4,4) size 9 depth 3





Efficient merging and sorting networks

Correctness proof of odd-even merging:



Correctness proof of odd-even merging: induction, zero-one principle Induction base: trivial (2 inputs, 1 comparator) Inductive step. Inductive hypothesis: odd, even merge both work correctly Let the input consist of 0s and 1s. We have for all k, l: $\langle 0^{\lceil k/2 \rceil} 11 \dots \rangle, \langle 0^{\lceil l/2 \rceil} 11 \dots \rangle \mapsto \langle 0^{\lceil k/2 \rceil + \lceil l/2 \rceil} 11 \dots \rangle$ in the odd merge $\langle 0^{\lfloor k/2 \rfloor} 11 \dots \rangle, \langle 0^{\lfloor l/2 \rfloor} 11 \dots \rangle \mapsto \langle 0^{\lfloor k/2 \rfloor + \lfloor l/2 \rfloor} 11 \dots \rangle$ in the even merge

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The final stage of comparators corrects the wrong pair $\langle 0^k 11 \dots \rangle, \langle 0^l 11 \dots \rangle \mapsto \langle 0^{k+l} 11 \dots \rangle$

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Sorting an arbitrary input $\langle x_1, \ldots, x_n \rangle$

Odd-even merge sorting

[Batcher: 1968]

When n = 1 we are done, otherwise by recursion:

- sort $\langle x_1, \ldots, x_{\lceil n/2 \rceil} \rangle$
- sort $\langle x_{\lceil n/2 \rceil+1}, \ldots, x_n \rangle$
- merge results by $OEM(\lceil n/2 \rceil, \lfloor n/2 \rfloor)$



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 $size(OEM-SORT)(n) \le 2 \cdot size(OEM-SORT(n/2)) + size(OEM(n/2, n/2)) = 2 \cdot size(OEM-SORT(n/2)) + O(n \log n) = O(n(\log n)^2)$ $depth(OEM-SORT(n)) \le 0$

 $depth(OEM-SORT(n/2)) + depth(OEM(n/2, n/2)) = depth(OEM-SORT(n/2)) + O(\log n) = O((\log n)^2)$

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OEM-SORT(n)size $O(n(\log n)^2)$ depth $O((\log n)^2)$



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OEM-SORT(n)size $O(n(\log n)^2)$ depth $O((\log n)^2)$

OEM-SORT(8) size 19

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depth 6



Efficient merging and sorting networks

A bitonic sequence:
$$\langle x_1 \ge \cdots \ge x_m \le \cdots \le x_n \rangle$$
 $1 \le m \le n$



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Exercise: prove correctness (by zero-one principle)

Note: cannot exchange \geq and \leq in definition of bitonic!

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BM(n)size $O(n \log n)$ depth $O(\log n)$



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Computation by circuits Efficient merging and sorting networks

BM(n)size $O(n \log n)$ depth $O(\log n)$



BM(8) size 12 depth 3





Bitonic merge sorting

[Batcher: 1968]

When n = 1 we are done, otherwise by recursion:

- sort $\langle x_1, \dots, x_{\lceil n/2 \rceil} \rangle \mapsto \langle y_1 \ge \dots \ge y_{\lceil n/2 \rceil} \rangle$ in reverse
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- comparators are actually nodes in a circuit, which can always be drawn using "standard comparators"
- a network drawn with "inverted comparators" can be converted into one with only "standard comparators" by a top-down rearrangement

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Computation by circuits Efficient merging and sorting networks

BM-SORT(n)size $O(n(\log n)^2)$ depth $O((\log n)^2)$



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Computation by circuits Efficient merging and sorting networks

BM-SORT(n)
size $O(n(\log n)^2)$
depth $O((\log n)^2)$

BM-SORT(8) size 24 depth 6



BM-SORT

 $(\lfloor n/2 \rfloor)$

BM-SORT

 $(\lceil n/2 \rceil)$

BM(n)

Both *OEM-SORT* and *BM-SORT* have size $\Theta(n(\log n)^2)$ Is it possible to sort obliviously in size $o(n(\log n)^2)$? $O(n \log n)$?

Image: A matrix

Both *OEM-SORT* and *BM-SORT* have size $\Theta(n(\log n)^2)$ Is it possible to sort obliviously in size $o(n(\log n)^2)$? $O(n \log n)$? AKS sorting [Ajtai, Komlós, Szemerédi: 1983] [Paterson: 1990]; [Seiferas: 2009]

Sorting network: size $O(n \log n)$, depth $O(\log n)$

Uses sophisticated graph theory (expanders)

Asymptotically optimal, but has huge constant factors

Computation by circuits

2 Parallel computation models

- 3 Basic parallel algorithms
- 4 Further parallel algorithms
- 5 Parallel matrix algorithms
- 6 Parallel graph algorithms

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Parallel Random Access Machine (PRAM)

Simple, idealised general-purpose parallel model



Parallel Random Access Machine (PRAM)

Simple, idealised general-purpose parallel model



Contains

- unlimited number of processors (1 time unit/op)
- global shared memory (1 time unit/access)

Operates in full synchrony

PRAM computation: sequence of parallel steps Communication and synchronisation taken for granted Not scalable in practice!

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PRAM computation: sequence of parallel steps

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Not scalable in practice!

PRAM variants:

- concurrent/exclusive read
- concurrent/exclusive write

CRCW, CREW, EREW, (ERCW) PRAM

PRAM computation: sequence of parallel steps

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- concurrent/exclusive read
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CRCW, CREW, EREW, (ERCW) PRAM

E.g. a linear system solver: $O((\log n)^2)$ steps using n^4 processors :-0

PRAM algorithm design: minimising number of steps, sometimes also number of processors

Bulk-Synchronous Parallel (BSP) computer

Simple, realistic general-purpose parallel model

Goals: scalability, portability, predictability



Bulk-Synchronous Parallel (BSP) computer

Simple, realistic general-purpose parallel model

Goals: scalability, portability, predictability



[Valiant: 1990]

Contains

- *p* processors, each with local memory (1 time unit/operation)
- communication environment, including a network and an external memory (g time units/data unit communicated)
- barrier synchronisation mechanism (*I* time units/synchronisation)

Some elements of a BSP computer can be emulated by others, e.g.

- external memory by local memory + network communication
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Communication network parameters:

- g is communication gap (inverse bandwidth), worst-case time for a data unit to enter/exit the network
- *I* is latency, worst-case time for a data unit to get across the network

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Every parallel system can be (approximately) described by p, g, lNetwork efficiency grows slower than processor efficiency and costs more energy: $g, l \gg 1$. E.g. for Cray T3E: p = 64, $g \approx 78$, $l \approx 1825$

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FIG. 1.13. Time of an h-relation on a 64-processor Cray T3E.

TABLE 1.2. Benchmarked BSP parameters p, g, l and the time of a 0-relation for a Cray T3E. All times are in flop units (r = 35 Mflop/s)

p	9	l	$T_{comm}(0)$
1	36	47	38
2	28	486	325
4	31	679	437
8	31	1193	580
16	31	2018	757
32	72	1145	871
64	78	1825	1440

is a used, rather than a forus. Increasing the number of processor nucleon the subpartition look more like a torus, with richer connectivity.) The time of a Gredzian (i.e. the time of a superstep without communication) displays a sanotcher behaviour than that of (a, nd) it is presented here for comparison. This time is a layer bound on l_i since it represents only part of the fixed cast of a superstep.

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Alexander Tiskin (Warwick)

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BSP computation: sequence of parallel supersteps



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BSP computation: sequence of parallel supersteps



Asynchronous computation/communication within supersteps (includes data exchange with external memory)

Synchronisation before/after each superstep

BSP computation: sequence of parallel supersteps



Asynchronous computation/communication within supersteps (includes data exchange with external memory)

Synchronisation before/after each superstep

Cf. CSP: parallel collection of sequential processes

Compositional cost model

For individual processor proc in superstep sstep:

- comp(sstep, proc): the amount of local computation and local memory operations by processor proc in superstep sstep
- comm(sstep, proc): the amount of data sent and received by processor proc in superstep sstep

Compositional cost model

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For the whole BSP computer in one superstep *sstep*:

- comp(sstep) = max_{0≤proc<p} comp(sstep, proc)
- comm(sstep) = max_{0≤proc<p} comm(sstep, proc)
- $cost(sstep) = comp(sstep) + comm(sstep) \cdot g + l$

For the whole BSP computation with *sync* supersteps:

•
$$comp = \sum_{0 \le sstep < sync} comp(sstep)$$

• $comm = \sum_{0 \le sstep < sync} comm(sstep)$
• $cost = \sum_{0 \le sstep < sync} cost(sstep) = comp + comm \cdot g + sync \cdot I$

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The input/output data are stored in the external memory; the cost of input/output is included in *comm*

E.g. for a particular linear system solver with an $n \times n$ matrix:

$$comp = O(n^3/p)$$
 $comm = O(n^2/p^{1/2})$ $sync = O(p^{1/2})$

- problem size $n \gg p$ (slackness)
- input/output in external memory, counts as one-sided communication

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BSP algorithm design: minimising comp, comm, sync

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BSP algorithm design: minimising *comp*, *comm*, *sync* Main principles:

- computation load balancing: ideally, $comp = O(\frac{seq \ work}{p})$
- data locality: ideally $comm = O(\frac{input/output}{p})$
- coarse granularity: ideally, sync function of p not n (or better, O(1))

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Data locality exploited, network locality ignored!

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BSP software: industrial projects

- Google's Pregel [2010]
- Apache Hama, Spark, Giraph (apache.org) [2010–16]

BSP software: research projects

- Oxford BSP (www.bsp-worldwide.org/implmnts/oxtool) [1998]
- Paderborn PUB (www2.cs.uni-paderborn.de/~pub) [1998]
- BSML (traclifo.univ-orleans.fr/BSML)
- BSPonMPI (bsponmpi.sourceforge.net)
- Multicore BSP (www.multicorebsp.com)
- Epiphany BSP (www.codu.in/ebsp)
- Petuum (petuum.org)

[1998]

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Broadcasting:

- initially, one designated processor holds a value a
- at the end, every processor must hold a copy of a

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Combining (complementary to broadcasting):

- initially, every processor r holds a value a_r
- at the end, one designated processor must hold $\sum_r a_r$
- addition can be replaced by any given associative operator •:
 a (b c) = (a b) c, computable in time O(1)

Examples: numerical +, \cdot , min, max, Boolean \land , \lor , ...

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By symmetry, we only need to consider broadcasting

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Fundamental communication patterns

Direct broadcast:

 designated processor makes *p* - 1 copies of *a* and sends them directly to destinations



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$$comp = O(p)$$

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Fundamental communication patterns

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Cost components will be shaded when they are optimal, i.e. cannot be improved by another algorithm (under certain explicit assumptions)

Fundamental communication patterns

Binary tree broadcast:

- initially, only designated processor is awake
- processors wake up each other in log p rounds
- in every round, every awake processor makes a copy of *a* and send it to a sleeping processor, waking it up



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$$comm = O(\log p)$$

$$sync = O(\log p)$$

 $comp = O(\log p)$

Array broadcasting:

- initially, one designated processor holds array a of size $n \ge p$
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By symmetry, we only need to consider array broadcasting

Fundamental communication patterns

Two-phase array broadcast:

- partition array into p blocks of size n/p
- scatter blocks
- total-exchange blocks



Fundamental communication patterns

Two-phase array broadcast:

- partition array into p blocks of size n/p
- scatter blocks
- total-exchange blocks

$$\begin{array}{c} A & B \\ C & D \\ \hline \\ A & B \\ \hline \\ C & D \\ \hline \\ A & B \\ C & D \\ \hline \\ C & D \\ \hline \\ C & D \\ \hline \\ \\ Sync = O(1) \end{array}$$

$$comp = O(n)$$

$$comm = O(n)$$



Array broadcasting/combining enables concurrent access to external memory in blocks of size $\geq p$

Concurrent reading: a designated processor

- reads block from external memory
- broadcasts block

Concurrent writing, resolved by •: a designated processor

- combines blocks from each processor by •
- writes combined block to external memory

Two-phase array broadcast/combine used as subroutine

Network routing

BSP network model: complete graph, uniformly accessible (access efficiency described by parameters g, l)

Has to be implemented on concrete networks

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Network routing

BSP network model: complete graph, uniformly accessible (access efficiency described by parameters g, l)

Has to be implemented on concrete networks

Parameters of a network topology (i.e. the underlying graph):

- degree number of links per node
- diameter maximum distance between nodes

Low degree — easier to implement

Low diameter — more efficient

Network routing

2D array network $p = q^2$ processors degree 4 diameter $p^{1/2} = q$



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Network routing

3D array network $p = q^3$ processors degree 6 diameter $3/2 \cdot p^{1/3} = 3/2 \cdot q$



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Network routing

Butterfly network

 $p = q \log q$ processors

degree 4

diameter $\approx \log p \approx \log q$



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Network routing

Hypercube network

 $p = 2^q$ processors

degree $\log p = q$

diameter $\log p = q$



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Network routing

Network	Degree	Diameter
1D array	2	$1/2 \cdot p$
2D array	4	$p^{1/2}$
3D array	6	$3/2 \cdot p^{1/3}$
Butterfly	4	log p
Hypercube	log p	log p
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BSP parameters g, l depend on degree, diameter, routing strategy

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Assume store-and-forward routing (alternative — wormhole)

Assume distributed routing: no global control

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Butterfly	4	log p
Hypercube	log p	log p
• • •		• • •

BSP parameters g, l depend on degree, diameter, routing strategy
Assume store-and-forward routing (alternative — wormhole)
Assume distributed routing: no global control
Oblivious routing: path determined only by source and destination
E.g. greedy routing: a packet always takes the shortest path

Network routing

h-relation (*h*-superstep): every processor sends and receives $\leq h$ packets

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Parallel computation models Network routing

h-relation (*h*-superstep): every processor sends and receives $\leq h$ packets Sufficient to consider permutations (1-relations): once we can route any permutation in *k* steps, we can route any *h*-relation in *hk* steps



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Any routing method may be forced to make $\Omega(diameter)$ steps



h-relation (*h*-superstep): every processor sends and receives $\leq h$ packets Sufficient to consider permutations (1-relations): once we can route any permutation in *k* steps, we can route any *h*-relation in *hk* steps Any routing method may be forced to make $\Omega(diameter)$ steps Any oblivious routing method may be forced to make $\Omega(p^{1/2}/degree)$ steps Many practical patterns force such "hot spots" on traditional networks



Parallel computation models Network routing

Routing based on sorting networks

- Each processor corresponds to a wire
- Each link corresponds to (possibly several) comparators
- Routing corresponds to sorting by destination address
- Each stage of routing corresponds to a stage of sorting

Parallel computation models Network routing

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Such routing is non-oblivious (for individual packets)!

Network	Degree	Diameter
OEM-SORT/BM-SORT	$O((\log p)^2)$	$O((\log p)^2)$
AKS	$O(\log p)$	$O(\log p)$

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Network	Degree	Diameter
OEM-SORT/BM-SORT	$O((\log p)^2)$	$O((\log p)^2)$
AKS	$O(\log p)$	$O(\log p)$

No "hot spots": can always route a permutation in O(diameter) steps Requires a specialised network, too messy and impractical

Parallel computation models Network routing

Two-phase randomised routing:

[Valiant: 1980]

- send every packet to random intermediate destination
- forward every packet to final destination

Both phases oblivious (e.g. greedy), but non-oblivious overall due to randomness

Parallel computation models Network routing

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Hot spots very unlikely: on a 2D array, butterfly, hypercube, can route a permutation in O(diameter) steps with high probability

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On a hypercube, the same holds even for a log p-relation

Hence constant g, I in the BSP model



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Network routing

BSP implementation: processes placed at random, communication delayed until end of superstep

All packets with same source and destination sent together, hence message overhead absorbed in I



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Network routing

BSP implementation: processes placed at random, communication delayed until end of superstep

All packets with same source and destination sent together, hence message overhead absorbed in ${\it I}$

Network	g	1
1D array	<i>O</i> (<i>p</i>)	<i>O</i> (<i>p</i>)
2D array	$O(p^{1/2})$	$O(p^{1/2})$
3D array	$O(p^{1/3})$	$O(p^{1/3})$
Butterfly	$O(\log p)$	$O(\log p)$
Hypercube	O(1)	$O(\log p)$

Actual values of g, I obtained by running benchmarks



Parallel computation models

3 Basic parallel algorithms

- 4 Further parallel algorithms
- 5 Parallel matrix algorithms
- 6 Parallel graph algorithms



Balanced tree and prefix sums

The balanced binary tree dag:

tree(n) 1 input, n outputs size n-1depth log n b_0 b_1 b_2 b_3 b_4 b_5 b_6 b_7 b_8 b_9 b_{10} b_{11} b_{12} b_{13} b_{14} b_{15}

Balanced tree and prefix sums

The balanced binary tree dag:



Can be directed

- top-down (one input at root, *n* outputs at leaves)
- bottom-up (n inputs at leaves, one output at root)

Sequential work O(n)

Balanced tree and prefix sums

Parallel balanced tree computation, p = 4



From now on, we always assume that a problem's input/output is stored in the external memory; reading/writing will also refer to the external memory

Balanced tree and prefix sums

Parallel balanced tree computation, p = 4



From now on, we always assume that a problem's input/output is stored in the external memory; reading/writing will also refer to the external memory Partition tree(n) into

- one top block, isomorphic to *tree*(*p*)
- a bottom layer of p blocks, each isomorphic to tree(n/p)

Balanced tree and prefix sums

Parallel balanced tree computation (contd.)

For top-down computation, a designated processor

- is assigned the top block
- reads block's input, computes block, writes block's p outputs

Then every processor

- is assigned a different bottom block
- reads block's input, computes block, writes block's n/p outputs

For bottom-up computation, reverse the steps

Balanced tree and prefix sums

Parallel balanced tree computation (contd.)

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For bottom-up computation, reverse the steps

$$comm = O(n/p)$$

$$sync = O(1)$$

Required slackness $n \ge p^2$

comp = O(n/p)

Balanced tree and prefix sums

The described parallel balanced tree algorithm is fully optimal:

- optimal $comp = O(n/p) = O(\frac{\text{sequential work}}{p})$
- optimal $comm = O(n/p) = O(\frac{input/output size}{p})$
- optimal sync = O(1)

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Balanced tree and prefix sums

The described parallel balanced tree algorithm is fully optimal:

• optimal $comp = O(n/p) = O(\frac{\text{sequential work}}{p})$

• optimal
$$comm = O(n/p) = O(rac{input/output size}{p})$$

• optimal
$$sync = O(1)$$

For other problems, we may not be so lucky to get a fully-optimal BSP algorithm. However, we are typically interested in algorithms that are optimal in *comp* (under reasonable assumptions).

Optimality in comm and sync is considered subject to optimality in comp

For example, we are not allowed to "cheat" by running the whole computation in a single processor, sacrificing *comp* and *comm* to guarantee optimal sync = O(1)

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Balanced tree and prefix sums

The prefix sums problem

Given array $a = [a_0, \ldots, a_{n-1}]$

Compute
$$b_{-1} = 0$$
 $b_i = a_i + b_{i-1}$ $0 \le i < n$

Addition can be replaced by any given associative operator • Operator identity ϵ (can be introduced formally if missing) Compute $b_{-1} = \epsilon$ $b_i = a_i \bullet b_{i-1}$ $0 \le i < n$ $b_0 = a_0$ $b_1 = a_0 \bullet a_1$ $b_2 = a_0 \bullet a_1 \bullet a_2$... $b_{n-1} = a_0 \bullet a_1 \bullet \dots \bullet a_{n-1}$

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Balanced tree and prefix sums

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Sequential work O(n) by trivial circuit of size n-1, depth n-1

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Balanced tree and prefix sums



where $a_{k:l} = a_k \bullet a_{k+1} \bullet \ldots \bullet a_l$

The underlying dag is called the prefix dag

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Balanced tree and prefix sums

The prefix circuit (contd.)

prefix(n) n inputs

n outputs

size 2*n* – 2

depth $2 \log n$



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Balanced tree and prefix sums

Parallel prefix sums computation

Dag prefix(n) consists of

- a top subtree similar to bottom-up *tree*(*n*)
- transfer of values from top subtree to bottom subtree
- a bottom subtree similar to top-down *tree*(*n*)

Balanced tree and prefix sums

Parallel prefix sums computation Dag prefix(n) consists of

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Both trees can be computed by the previous algorithm Transfer stage: communication cost O(n/p)

Balanced tree and prefix sums

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- a bottom subtree similar to top-down tree(n)

Both trees can be computed by the previous algorithm

Transfer stage: communication cost O(n/p)

$$np = O(n/p)$$
 $comm = O(n/p)$

$$sync = O$$

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Required slackness $n \ge p^2$

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Balanced tree and prefix sums

Application: generic first-order linear recurrence Given arrays $a = [a_0, ..., a_{n-1}]$, $b = [b_0, ..., b_{n-1}]$ Compute $c_{-1} = 0$ $c_i = a_i + b_i \cdot c_{i-1}$ $0 \le i < n$ $c_0 = a_0$ $c_1 = a_1 + b_1 \cdot c_0$ $c_2 = a_2 + b_2 \cdot c_1$... $c_{n-1} = a_{n-1} + b_{n-1} \cdot c_{n-2}$

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Balanced tree and prefix sums

Application: generic first-order linear recurrence (contd.)

$$c_{-1} = 0 \quad c_i = a_i + b_i \cdot c_{i-1} \quad 0 \le i < n$$
Let $A_i = \begin{bmatrix} 1 & 0 \\ a_i & b_i \end{bmatrix} \quad C_i = \begin{bmatrix} 1 \\ c_i \end{bmatrix} \quad A_i C_{i-1} = \begin{bmatrix} 1 & 0 \\ a_i & b_i \end{bmatrix} \begin{bmatrix} 1 \\ c_{i-1} \end{bmatrix} = \begin{bmatrix} 1 \\ c_i \end{bmatrix} = C_i$

$$C_0 = A_0 \cdot C_{-1}$$

$$C_1 = A_1 A_0 \cdot C_{-1}$$

$$C_2 = A_2 A_1 A_0 \cdot C_{-1}$$

$$\cdots$$

$$C_{n-1} = A_{n-1} \cdots A_1 A_0 \cdot C_{-1}$$

Application: generic first-order linear recurrence (contd.)

Computing the generic first-order linear recurrence:

- suffix sums (= prefix sums in reverse) of [A_{n-1},..., A₀], with operator defined by 2 × 2-matrix multiplication
- each suffix sum multiplied by C_{-1}
- output obtained as bottom component of resulting 2-vectors

Resulting circuit: size O(n), depth $O(\log n)$

Application: generic first-order linear recurrence (contd.)

Operators +, \cdot can be replaced by any given $\oplus,$ $\odot,$ where

- operators \oplus , \odot computable in time O(1)
- operator \oplus associative: $a \oplus (b \oplus c) = (a \oplus b) \oplus c$
- operator \odot associative: $a \odot (b \odot c) = (a \odot b) \odot c$
- operator \odot (left-)distributive over \oplus : $a \odot (b \oplus c) = (a \odot b) \oplus (a \odot c)$

Examples of possible \oplus , \odot :

- numerical +, ·
- numerical min, +; numerical max, +
- Boolean \land , \lor ; Boolean \lor , \land

Balanced tree and prefix sums

Application: polynomial evaluation

Given $a = [a_0, ..., a_{n-1}]$, x

Compute $y = a_0 + a_1 \cdot x + \ldots + a_{n-2} \cdot x^{n-2} + a_{n-1} \cdot x^{n-1}$

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Balanced tree and prefix sums

Application: polynomial evaluation

Given $a = [a_0, ..., a_{n-1}]$, x

Compute $y = a_0 + a_1 \cdot x + \ldots + a_{n-2} \cdot x^{n-2} + a_{n-1} \cdot x^{n-1}$

Evaluating the polynomial:

- $1, x, x^2, \dots, x^{n-1}$ by prefix sums with operator \cdot
- sum y by bottom-up balanced binary tree with operator +

Resulting circuit: size O(n), depth $O(\log n)$

Balanced tree and prefix sums

Application: polynomial evaluation by Horner's rule Given $a = [a_0, \ldots, a_{n-1}], x$ Compute $y = a_0 + a_1 \cdot x + \ldots + a_{n-2} \cdot x^{n-2} + a_{n-1} \cdot x^{n-1}$ $y = a_0 + x \cdot (a_1 + x \cdot (a_2 + x \cdot (\dots + x \cdot a_{n-1}) \dots))$ $y_0 = a_{n-1}$ $y_1 = a_{n-2} + x \cdot y_0$ $V_2 = a_{n-3} + X \cdot V_1$. . . $V_{n-1} = a_0 + x \cdot V_{n-2}$ Generic first-order linear recurrence over $[a_{n-1}, \ldots, a_0]$, $[x, x, \ldots, x]$ Resulting circuit: size O(n), depth $O(\log n)$

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Balanced tree and prefix sums

Application: binary addition via Boolean logic x + y = z , x, y, z represented as binary arrays $x = [x_{n-1}, \dots, x_0]$ $y = [y_{n-1}, \dots, y_0]$ $z = [z_n, z_{n-1}, \dots, z_0]$ The binary adder problem: given x, y, compute z

Boolean operators as primitives: bitwise \land ("and"), \lor ("or"), \oplus ("xor") Let $c = [c_{n-1}, \ldots, c_0]$, where c_i is the *i*-th carry bit We have: $x_i + y_i + c_{i-1} = z_i + 2c_i$ $0 \le i < n$

Balanced tree and prefix sums

Application: binary addition via Boolean logic (contd.) Define bit arrays $u = [u_{n-1}, \ldots, u_0]$, $v = [v_{n-1}, \ldots, v_0]$ $u_i = x_i \land y_i$ $v_i = x_i \oplus y_i$ $0 \le i < n$ $z_0 = v_0$ $c_0 = u_0$ $z_1 = v_1 \oplus c_0$ $c_1 = u_1 \lor (v_1 \land c_0)$ \ldots $z_{n-1} = v_{n-1} \oplus c_{n-2}$ $c_{n-1} = u_{n-1} \lor (v_{n-1} \land c_{n-2})$ $z_n = c_{n-1}$

Alexander Tiskin (Warwick)

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Balanced tree and prefix sums

Application: binary addition via Boolean logic (contd.) Define bit arrays $u = [u_{n-1}, ..., u_0], v = [v_{n-1}, ..., v_0]$ $u_i = x_i \wedge y_i$ $v_i = x_i \oplus v_i$ $0 \leq i < n$ $z_0 = v_0$ $c_0 = u_0$ $c_1 = u_1 \vee (v_1 \wedge c_0)$ $z_1 = v_1 \oplus c_0$ $z_{n-1} = v_{n-1} \oplus c_{n-2}$ $c_{n-1} = u_{n-1} \lor (v_{n-1} \land c_{n-2})$ $Z_n = C_{n-1}$ Resulting circuit has size and depth O(n)

Equivalent to a ripple-carry adder. Can we do better?

Balanced tree and prefix sums

Application: binary addition via Boolean logic (contd.)

$$c_{-1} = 0$$
 $c_i = u_i \lor (v_i \land c_{i-1})$

Compute

- c as generic first-order linear recurrence with inputs u, v and operators ∨, ∧: size O(n), depth O(log n)
- z in extra size O(n), depth O(1)

Resulting circuit has size O(n), depth $O(\log n)$

Equivalent to a carry-lookahead adder

Integer sorting

The integer sorting problem

Given $a = [a_0, ..., a_{n-1}]$, arrange elements of a in increasing order $a_i \in \{0, 1, ..., n-1\}$ $0 \le i < n$

Elements of *a* assumed to be distinguishable keys even if values equal

A bucket: subset of keys with equal values

Stable integer sorting: order of keys preserved within each bucket Sequential work O(n) e.g. by bucket sort or counting sort

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Integer sorting

Parallel integer sorting

Initially assume
$$a_i \in \{0, 1, \dots, \frac{n}{p} - 1\}$$
, i.e. $\frac{n}{p}$ buckets

Every processor

- reads subarray of a of size n/p
- counts subarray elements in each bucket

A designated processor

- adds subarray counts for each bucket (array combining)
- determines bucket boundaries, broadcasts them (array broadcasting)

Every processor

• writes each element at appropriate offset from bucket boundary

Integer sorting

Parallel integer sorting (contd.)

Now consider $a_i \in \{0, 1, \dots, p-1\}$, i.e. p buckets

Consider keys as pairs: $a_i = (a_i \mod \frac{n}{p}, a_i \dim \frac{n}{p})$

Perform 2-fold radix sort on pairs:

- left ("least significant") position
- right ("most significant") position

In each position, perform stable sorting over range $\{0, 1, \dots, \frac{n}{p} - 1\}$

$$comp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(1)$
Required slackness $n \ge p^2$

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A complex number ω is called a primitive root of unity of degree *n*, if $\omega, \omega^2, \ldots, \omega^{n-1} \neq 1$, and $\omega^n = 1$

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A complex number ω is called a primitive root of unity of degree *n*, if $\omega, \omega^2, \ldots, \omega^{n-1} \neq 1$, and $\omega^n = 1$

The Discrete Fourier Transform problem: given complex vector a, compute b, where $F_{n,\omega} \cdot a = b$, and $F_{n,\omega} = \left[\omega^{ij}\right]_{i,j=0}^{n-1}$ is the Fourier matrix

$$\begin{bmatrix} 1 & 1 & 1 & \cdots & 1 \\ 1 & \omega & \omega^2 & \cdots & \omega^{n-1} \\ 1 & \omega^2 & \omega^4 & \cdots & \omega^{n-2} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & \omega^{n-1} & \omega^{n-2} & \cdots & \omega \end{bmatrix} \cdot \begin{bmatrix} a_0 \\ a_1 \\ a_2 \\ \vdots \\ a_{n-1} \end{bmatrix} = \begin{bmatrix} b_0 \\ b_1 \\ b_2 \\ \vdots \\ b_{n-1} \end{bmatrix}$$
$$\sum_j \omega^{ij} a_j = b_i \qquad i, j = 0, \dots, n-1$$

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Sequential work $O(n^2)$ by matrix-vector multiplication

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$$\sum_j \omega^{ij} a_j = b_i \qquad i, j = 0, \dots, n-1$$

Sequential work $O(n^2)$ by matrix-vector multiplication

Applications: digital signal processing (amplitude vs frequency); polynomial multiplication; long integer multiplication

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Efficient Parallel Algorithms

The Fast Fourier Transform (FFT) algorithm ("four-step" version) Assume $n = m^2$

Let $A_{u,v} = a_{mu+v}$ $B_{s,t} = b_{ms+t}$ s, t, u, v = 0, ..., m-1

Matrices A, B are vectors a, b written out as $m \times m$ matrices

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The Fast Fourier Transform (FFT) algorithm ("four-step" version) Assume $n = m^2$ Let $A_{u,v} = a_{mu+v}$ $B_{s,t} = b_{ms+t}$ s, t, u, v = 0, ..., m-1Matrices A, B are vectors a, b written out as $m \times m$ matrices $B_{s,t} = \sum_{u,v} \omega^{(ms+t)(mu+v)} A_{u,v} = \sum_{u,v} \omega^{msv+tv+mtu} A_{u,v} =$ $\sum_{\nu} ((\omega^m)^{s\nu} \cdot \omega^{t\nu} \cdot \sum_{\mu} (\omega^m)^{t\mu} A_{\mu,\nu}), \text{ thus } B = F_{m,\omega^m} \cdot (G_{m,\omega} \circ (F_{m,\omega^m} \cdot A))^T$ $F_{m,\omega^m} \cdot A$ is *m* independent DFTs of size *m* on each column of A $G_{m,\omega} = \left[\omega^{tv}\right]_{t,v=0}^{m-1}$ is the twiddle-factor matrix Operator \circ is the Hadamard product (elementwise matrix multiplication)

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Basic parallel algorithms Fast Fourier Transform and the butterfly dag

The Fast Fourier Transform (FFT) algorithm (contd.)

$$B = F_{m,\omega^m} \cdot (G_{m,\omega} \circ (F_{m,\omega^m} \cdot A))^7$$

Thus, DFT of size *n* in four steps:

- *m* independent DFTs of size *m*
- transposition and twiddle-factor scaling
- *m* independent DFTs of size *m*

The Fast Fourier Transform (FFT) algorithm (contd.) We reduced DFT of size $n = m^2$ to DFTs of size mSimilarly, we can reduce DFT of size n = kl to DFTs of sizes k and lAssume $n = 2^{2^r}$, then $m = 2^{2^{r-1}}$

By recursion, we have the FFT circuit

The Fast Fourier Transform (FFT) algorithm (contd.) We reduced DFT of size $n = m^2$ to DFTs of size mSimilarly, we can reduce DFT of size n = kl to DFTs of sizes k and lAssume $n = 2^{2^r}$, then $m = 2^{2^{r-1}}$

By recursion, we have the FFT circuit

 $size_{FFT}(n) = O(n) + 2 \cdot n^{1/2} \cdot size_{FFT}(n^{1/2}) = O(1 \cdot n \cdot 1 + 2 \cdot n^{1/2} \cdot n^{1/2} + 4 \cdot n^{3/4} \cdot n^{1/4} + \dots + \log n \cdot n \cdot 1) = O(n + 2n + 4n + \dots + \log n \cdot n) = O(n \log n)$ $depth_{FFT}(n) = 1 + 2 \cdot depth_{FFT}(n^{1/2}) = O(1 + 2 + 4 + \dots + \log n) = O(\log n)$

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Basic parallel algorithms Fast Fourier Transform and the butterfly dag

The FFT circuit



The underlying dag is called butterfly dag

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Basic parallel algorithms Fast Fourier Transform and the butterfly dag

The FFT circuit and the butterfly dag (contd.)

bfly(n) *n* inputs *n* outputs size $\frac{n \log n}{2}$ depth log *n*



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The FFT circuit and the butterfly dag (contd.) Dag bfly(n) consists of

- a top layer of $n^{1/2}$ blocks, each isomorphic to $bfly(n^{1/2})$
- a bottom layer of $n^{1/2}$ blocks, each isomorphic to $bfly(n^{1/2})$

The data exchange pattern between the top and bottom layers corresponds to $n^{1/2} \times n^{1/2}$ matrix transposition

Parallel butterfly computation

To compute bfly(n), every processor

- reads inputs for $\frac{n^{1/2}}{p}$ blocks from top layer; computes blocks; writes outputs
- reads inputs for $\frac{n^{1/2}}{p}$ blocks from bottom layer; computes blocks; writes outputs

In each layer, the processor reads the total of n/p inputs, performs $O(n \log n/p)$ computation, then writes the total of n/p outputs

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Parallel butterfly computation

To compute bfly(n), every processor

- reads inputs for $\frac{n^{1/2}}{p}$ blocks from top layer; computes blocks; writes outputs
- reads inputs for $\frac{n^{1/2}}{p}$ blocks from bottom layer; computes blocks; writes outputs

In each layer, the processor reads the total of n/p inputs, performs $O(n \log n/p)$ computation, then writes the total of n/p outputs

$$O(\frac{n \log n}{p})$$
 $comm = O(n/p)$ $sync = O(1)$

Required slackness: $n \ge p^2$

comp =

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The ordered 2D grid dag

 $grid_2(n)$

nodes arranged in an $n \times n$ grid

edges directed top-to-bottom, left-to-right

- $\leq 2n$ inputs (to left/top borders)
- $\leq 2n$ outputs (from right/bottom borders) size n^2 depth 2n - 1



```
The ordered 2D grid dag
```

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nodes arranged in an $n \times n$ grid

edges directed top-to-bottom, left-to-right

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\leq 2n inputs (to left/top borders)
```

```
\leq 2n outputs (from right/bottom borders)
size n^2 depth 2n-1
```



Applications: triangular linear system; discretised PDE via Gauss–Seidel iteration (single step); 1D cellular automata; dynamic programming Sequential work $O(n^2)$

Parallel ordered 2D grid computation

 $grid_2(n)$

Partition into a $p \times p$ grid of blocks, each isomorphic to $grid_2(n/p)$



Parallel ordered 2D grid computation

 $grid_2(n)$

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Parallel ordered 2D grid computation (contd.)

The computation proceeds in 2p - 1 stages, each computing a layer of blocks. In a stage:

- every block assigned to a different processor (some processors idle)
- the processor reads the 2n/p block inputs, computes the block, and writes back the 2n/p block outputs

Parallel ordered 2D grid computation (contd.)

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comp: $(2p-1) \cdot O((n/p)^2) = O(p \cdot n^2/p^2) = O(n^2/p)$ comm: $(2p-1) \cdot O(n/p) = O(n)$

Parallel ordered 2D grid computation (contd.)

The computation proceeds in 2p-1 stages, each computing a layer of blocks. In a stage:

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$$comm = O(n)$$
 sync

$$sync = O(p)$$

Required slackness $n \ge p$

 $comp = O(n^2/p)$

Application: string comparison

Let *a*, *b* be strings of characters

A subsequence of string a is obtained by deleting some (possibly none, or all) characters from a

The longest common subsequence (LCS) problem: find the longest string that is a subsequence of both a and b

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In computational molecular biology, the LCS problem and its variants are referred to as sequence alignment

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LCS computation by dynamic programming[Wagner, Fischer: 1974]Let lcs(a, b) denote the LCS length

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$$lcs(a\alpha, b\beta) = \begin{cases} max(lcs(a\alpha, b), lcs(a, b\beta)) & \text{if } \alpha \neq \beta \\ lcs("", b) = 0 & \text{if } \alpha = \beta \end{cases}$$

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	*	D	Е	F	Ι	N	Е
*	0	0	0	0	0	0	0
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Data dependence in the table corresponds to the 2D grid dag

Parallel LCS computation

The 2D grid algorithm solves the LCS problem (and many others) by dynamic programming

$$comp = O(n^2/p)$$
 $comm = O(n)$ $sync = O(p)$

Image: Image:
Parallel LCS computation

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comm is not scalable (i.e. does not decrease with increasing p)Can scalable comm be achieved for the LCS problem?

:-(

Parallel LCS computation

Solve the more general semi-local LCS problem:

- each string vs all substrings of the other string
- all prefixes of each string against all suffixes of the other string

Divide-and-conquer on substrings of a, b: log p recursion levels

Each level assembles substring LCS from smaller ones by parallel seaweed multiplication

Base level: p semi-local LCS subproblems, each of size $n/p^{1/2}$ Sequential time still $O(n^2)$

Parallel LCS computation (cont.)

Communication vs synchronisation tradeoff

Parallelising normal $O(n \log n)$ seaweed multiplication: [Krusche, T: 2010]

$$comp = O(n^2/p)$$

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Special seaweed multiplication

Sacrifices some *comp*, *comm* for *sync*

[Krusche, T: 2007]

 $sync = O(\log p)$

Parallel LCS computation (cont.)

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Special seaweed multiplication

Sacrifices some comp, comm for sync

$$omp = O(n^2/p)$$
 $comm = O(\frac{n\log p}{p^{1/2}})$ $sync = O(\log p)$

Open problem: can we achieve $comm = O(\frac{n}{p^{1/2}})$, $sync = O(\log p)$?

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The ordered 3D grid dag

 $grid_3(n)$

nodes arranged in an $n \times n \times n$ grid

edges directed top-to-bottom, left-to-right, front-to-back

```
\leq 3n^2 inputs (to front/left/top)
```

 $\leq 3n^2$ outputs (from back/right/bottom)

size n^3 depth 3n - 2



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Applications: Gaussian elimination; discretised PDE via Gauss–Seidel iteration; 2D cellular automata; dynamic programming

Sequential work $O(n^3)$

Parallel ordered 3D grid computation

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Partition into $p^{1/2} \times p^{1/2} \times p^{1/2}$ grid of blocks, each isomorphic to $grid_3(n/p^{1/2})$



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Parallel ordered 3D grid computation (contd.)

The computation proceeds in $3p^{1/2} - 2$ stages, each computing a layer of blocks. In a stage:

- every processor is either assigned a block or is idle
- a non-idle processor reads the $3n^2/p$ block inputs, computes the block, and writes back the $3n^2/p$ block outputs

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$$comm = O(n^2/p^{1/2})$$

$$sync = O(p^{1/2})$$

Required slackness $n \ge p^{1/2}$

 $comp = O(n^3/p)$

Basic parallel algorithms Discussion

Costs *comp*, *comm*, *sync*: functions of n, pTypically, realistic slackness requirements: $n \gg p$ The goals:

- comp = comp_{opt} = comp_{seq}/p
- comm scales down with increasing p
- sync is a function of p, independent of n

Basic parallel algorithms Discussion

Costs *comp*, *comm*, *sync*: functions of n, pTypically, realistic slackness requirements: $n \gg p$ The goals:

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- sync is a function of p, independent of n

The challenges:

- efficient (optimal) algorithms
- good (sharp) lower bounds



- 2 Parallel computation models
- 3 Basic parallel algorithms
- 4 Further parallel algorithms
- 5 Parallel matrix algorithms
- 6 Parallel graph algorithms

Linked list: array of *n* nodes

Each node contains data and a pointer to (= index of) successor node

Nodes may be placed in array in an arbitrary order



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Logical structure linear: head, succ(head), succ(succ(head)), ...

- a pointer can be followed in time O(1)
- no global ranks/indexing/comparison



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- a pointer can be followed in time O(1)
- no global ranks/indexing/comparison



Pointer jumping at node *u*

Let • be an associative operator, computable in time O(1)

Pointer v and data a, b are kept, so that pointer jumping can be reversed: $succ(u) \leftarrow v \qquad data(u) \leftarrow a \qquad data(v) \leftarrow b$

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Further parallel algorithms

List contraction and colouring

Abstract view: node merging, allows e.g. for bidirectional links



Data a, b are kept, so that node merging can be reversed

Further parallel algorithms

List contraction and colouring

Abstract view: node merging, allows e.g. for bidirectional links



Data *a*, *b* are kept, so that node merging can be reversed

The list contraction problem: reduce the list to a single node by successive merging (note the result is independent on the merging order)

The list expansion problem: restore the original list, reversing contraction

Application: list ranking

Node's rank: distance from *head*

rank(head) = 0, rank(succ(head)) = 1, ...

The list ranking problem: each node to hold its rank

$$\bullet - 0 \rightarrow 1 \rightarrow 2 \rightarrow 3 \rightarrow 4 \rightarrow 5 \rightarrow 6 \rightarrow 7 - 1$$

Note the solution should be independent of the merging order

Application: list ranking (contd.)

Each intermediate node during contraction/expansion represents a contiguous sublist in the original list

Contraction phase: each node u holds length l(u) of corresponding sublist Initially, $l(u) \leftarrow 1$ for each node u

Merging v, w into u: $l(u) \leftarrow l(v) + l(w)$, keeping l(v), l(w)

Fully contracted list: single node t holding I(t) = n

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Application: list ranking (contd.) Expansion phase: each node holds

- length I(u) of corresponding sublist (as before)
- rank r(u) of the sublist's starting node

Fully contracted list: single node t holding

$$l(t) = n \quad r(t) \leftarrow 0$$

Un-merging u to v, w: restore I(u), I(v), then

$$r(v) \leftarrow r(u) \quad r(w) \leftarrow r(v) + l(v)$$

After full expansion: each node *u* holds

$$l(u) = 1$$
 $r(u) = rank(u)$

Further parallel algorithms

List contraction and colouring

Application: list prefix sums

Initially, each node u holds value $a_{rank(u)}$

 a_6 a₃ a_4 a_5 a_1 a_2 a_7 a_0

Application: list prefix sums

Initially, each node u holds value $a_{rank(u)}$

$$\bullet - a_0 \longrightarrow a_1 \longrightarrow a_2 \longrightarrow a_3 \longrightarrow a_4 \longrightarrow a_5 \longrightarrow a_6 \longrightarrow a_7 - _$$

Let ullet be an associative operator with identity ϵ

The list prefix sums problem: each node u to hold prefix sum $a_{0:rank(u)} = a_0 \bullet a_1 \bullet \cdots \bullet a_{rank(u)}$

Note the solution should be independent of the merging order

Application: list prefix sums (contd.)

Each intermediate node during contraction/expansion represents a contiguous sublist in the original list

Contraction phase: each node u holds the \bullet -sum l(u) corresponding sublist Initially, $l(u) \leftarrow a_{rank(u)}$ for each node uMerging v, w into u: $l(u) \leftarrow l(v) \bullet l(w)$, keeping l(v), l(w)Fully contracted list: single node t with $l(t) = a_{0:n-1}$

Application: list prefix sums (contd.) Expansion phase: each node holds

- •-sum *l*(*u*) of corresponding sublist (as before)
- •-sum r(u) of all nodes before the sublist

Fully contracted list: single node t holding

$$I(t) = a_{0:n-1} \quad r(t) \leftarrow e^{-t}$$

Un-merging u to v, w: restore I(u), I(v), then

$$r(v) \leftarrow r(u) \quad r(w) \leftarrow r(v) \bullet l(v)$$

After full expansion: each node *u* holds

$$l(u) = a_{rank(u)}$$
 $r(u) = a_{0:rank(u)}$
In general, only need to consider contraction phase (expansion by symmetry)

Sequential contraction: always merge head with succ(head), time O(n)

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In general, only need to consider contraction phase (expansion by symmetry)

Sequential contraction: always merge head with succ(head), time O(n)

Parallel contraction must be based on local merging decisions: a node can be merged with either its successor or predecessor, but not both

Therefore, we need either node splitting, or efficient symmetry breaking

List contraction and colouring

Wyllie's mating

[Wyllie: 1979]

Image: A matrix



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List contraction and colouring



Alexander Tiskin (Warwick)

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List contraction and colouring



Merge mating node pairs, obtaining two lists of size $\approx n/2$



Parallel list contraction by Wyllie's mating

In the first round, every processor

- inputs n/p nodes (not necessarily contiguous in input list), overall n nodes forming input list across p processors
- performs node splitting and labelling
- merges mating pairs; each merge involves communication between two processors; the merged node placed arbitrarily on either processor
- outputs the resulting ≤ 2n/p nodes (not necessarily contiguous in output list), overall n nodes forming output lists across p processors

Subsequent rounds similar

Parallel list contraction by Wyllie's mating (contd.)

Parallel list contraction:

- perform log *n* rounds of Wyllie's mating, reducing original list to *n* fully contracted lists of size 1
- select one fully contracted list

Parallel list contraction by Wyllie's mating (contd.)

Parallel list contraction:

- perform log *n* rounds of Wyllie's mating, reducing original list to *n* fully contracted lists of size 1
- select one fully contracted list

Total work $O(n \log n)$, not optimal vs. sequential work O(n)

$$comp = O(\frac{n \log n}{p})$$
 $comm = O(\frac{n \log n}{p})$ $sync = O(\log n)$ $n \ge p$

List contraction and colouring

Random mating

[Miller, Reif: 1985]

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Label every node "forward" $\overleftarrow{\mathbb{G}}$ or "backward" $\overleftarrow{\mathbb{G}}$ independently with probability $\frac{1}{2}$

 $\bullet - \overleftarrow{\mathbb{Q}} \to \overleftarrow{\mathbb{Q}} \to$



List contraction and colouring

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probability $\frac{1}{2}$ $\bullet - \emptyset ? \to \emptyset$

Merge mating node pairs



List contraction and colouring

Random mating

Label every node "forward" 3 or "backward" 3 independently with probability $\frac{1}{2}$

Merge mating node pairs



On average $\frac{n}{2}$ nodes mate, therefore new list has expected size $\frac{3n}{4}$ Moreover, size $\leq \frac{15n}{16}$ with high probability (whp), i.e. with probability exponentially close to 1 (as a function of n)

$$Prob\left(\mathsf{new size} \leq rac{15n}{16}
ight) \geq 1 - e^{-n/64}$$

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[Miller, Reif: 1985]

Parallel list contraction by random mating

In the first round, every processor

- inputs ⁿ/_p nodes (not necessarily contiguous in input list), overall n nodes forming input list across p processors
- performs node randomisation and labelling
- merges mating pairs; each merge involves communication between two processors; the merged node placed arbitrarily on either processor
- outputs the resulting $\leq \frac{n}{p}$ nodes (not necessarily contiguous in output list), overall $\leq \frac{15n}{16}$ nodes (whp), forming output list across p processors

Subsequent rounds similar, on a list of decreasing size (whp)

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Parallel list contraction by random mating (contd.)

Parallel list contraction:

- perform $\log_{16/15} p$ rounds of random mating, reducing original list to size $\frac{n}{p}$ whp
- a designated processor inputs the remaining list, contracts it sequentially

Parallel list contraction by random mating (contd.)

Parallel list contraction:

- perform $\log_{16/15} p$ rounds of random mating, reducing original list to size $\frac{n}{p}$ whp
- a designated processor inputs the remaining list, contracts it sequentially

Total work O(n), optimal but randomised

comp = O(n/p) whpcomm = O(n/p) whp $sync = O(\log p)$ Required slackness $n > p^2$

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Further parallel algorithms List contraction and colouring

Block mating

Will mate nodes deterministically

Contract local chains (if any)

Build distribution graph:

- complete weighted digraph on *p* supernodes
- $w(i,j) = |\{u \rightarrow v : u \in proc_i, v \in proc_j\}|$

Each processor holds a supernode's outgoing edges





Block mating (contd.)

Designated processor collects the distribution graph

Label every supernode F ("forward") or B ("backward"), so that $\sum_{i \in F, j \in B} w(i, j) \ge \frac{1}{4} \cdot \sum_{i, j} w(i, j)$ by a sequential greedy algorithm

Distribute supernode labels to processors

Merge mating node pairs

By construction of supernode labelling, $\geq \frac{n}{2}$ nodes mate, therefore new list has size $\leq \frac{3n}{4}$



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B

Parallel list contraction by block mating

In the first round, every processor

- inputs ⁿ/_p nodes (not necessarily contiguous in input list), overall n nodes forming input list across p processors
- participates in construction of distribution graph and communicating it to the designated processor

The designated processor collects distribution graph, computes and distributes labels

Parallel list contraction by block mating (contd.)

Continuing the first round, every processor

- receives its label from the designated processor
- merges mating pairs; each merge involves communication between two processors; the merged node placed arbitrarily on either processor
- outputs the resulting $\leq \frac{n}{p}$ nodes (not necessarily contiguous in output list), overall $\leq \frac{3n}{4}$ nodes, forming output list across p processors

Subsequent rounds similar, on a list of decreasing size

Parallel list contraction by block mating (contd.)

Parallel list contraction:

- perform $\log_{4/3} p$ rounds of block mating, reducing the original list to size n/p
- a designated processor collects the remaining list and contracts it sequentially

Parallel list contraction by block mating (contd.)

Parallel list contraction:

- perform $\log_{4/3} p$ rounds of block mating, reducing the original list to size n/p
- a designated processor collects the remaining list and contracts it sequentially

Total work O(n), optimal and deterministic

comp = O(n/p)comm = O(n/p) $sync = O(\log p)$ Required slackness $n > p^4$



Using list contraction, k-colouring for any k can be done in

$$comp = O(n/p)$$

$$comm = O(n/p)$$

$$sync = O(\log p)$$

Is list contraction really necessary for list k-colouring?

Can list k-colouring be done more efficiently?



Using list contraction, k-colouring for any k can be done in

$$op = O(n/p)$$
 comm = $O(n/p)$

$$sync = O(\log p)$$

Is list contraction really necessary for list k-colouring?

Can list k-colouring be done more efficiently?

For k = p: we can easily (how?) do *p*-colouring in

$$comp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(n/p)$

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Using list contraction, k-colouring for any k can be done in

$$mp = O(n/p)$$
 $comm = O(n/p)$

$$sync = O(\log p)$$

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Is list contraction really necessary for list k-colouring?

Can list *k*-colouring be done more efficiently?

For k = p: we can easily (how?) do *p*-colouring in

$$comp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(1)$

Can this be extended to any $k \leq p$, e.g. k = O(1)?

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List contraction and colouring

Deterministic coin tossing

Given a k-colouring, k > 6

[Cole, Vishkin: 1986]

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Deterministic coin tossing

Given a k-colouring, k > 6

Consider every node v. We have $col(v) \neq col(succ(v))$.

If col(v) differs from col(succ(v)) in *i*-th bit, re-colour v in

- 2*i*, if *i*-th bit in col(v) is 0, and in col(succ(v)) is 1
- 2i + 1, if *i*-th bit in col(v) is 1, and in col(succ(v)) is 0

Model assumption: can find lowest nonzero bit in an integer in time O(1)After re-colouring, still have $col(v) \neq col(succ(v))$ Number of colours reduced from k to $2\lceil \log k \rceil \ll k$ comp, comm: O(n/p)

[Cole, Vishkin: 1986]

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Parallel list colouring by deterministic coin tossing

Reducing the number of colours from p to 6: need $O(\log^* p)$ rounds of deterministic coin tossing

The iterated log function

 $\log^* k = \min r : \log \ldots \log k \le 1$ (r times)

Parallel list colouring by deterministic coin tossing

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The iterated log function

 $\log^* k = \min r : \log \ldots \log k \le 1$ (r times)

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Number of particles in observable universe: $10^{81} \approx 2^{270}$

 $\log^* 2^{270} = \log^* 2^{65536} = \log^* 2^{2^{2^2}} = 5$

Parallel list colouring by deterministic coin tossing (contd.)

Initially, each processor reads a subset of n/p nodes

- partially contract the list to size O(n/log* p) by log_{4/3} log* p rounds of block mating
- compute a *p*-colouring of the resulting list
- reduce the number of colours from p to 6 by O(log* p) rounds of deteministic coin tossing

comp, comm: $O\left(\frac{n}{p} + \frac{n}{p \log^* p} \cdot \log^* p\right) = O(n/p)$ sync: $O(\log^* p)$

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Parallel list colouring by deterministic coin tossing (contd.)

We have a 6-coloured, partially contacted list of size $O(n/\log^* p)$

- select node v as a pivot, if col(pred(v)) > col(v) < col(succ(v)); no two pivots are adjacent or further than 12 nodes apart
- re-colour all pivots in one colour
- $\bullet\,$ from each pivot, 2-colour the next ≤ 12 non-pivots sequentially; we now have a 3-coloured list
- reverse the partial contraction, maintaining the 3-colouring

We have now 3-coloured the original list

comp = O(n/p)

$$comm = O(n/p)$$

$$\boxed{sync = O(\log^* p)} \qquad n \ge \mu$$

The sorting problem

Given $a = [a_0, \ldots, a_{n-1}]$, arrange elements of a in increasing order

May assume all a_i are distinct (otherwise, attach unique tags)

Assume the comparison model: primitives \langle , \rangle , no arithmetic or bit operations on a_i

Sequential work $O(n \log n)$ e.g. by mergesort

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The sorting problem

Given $a = [a_0, \ldots, a_{n-1}]$, arrange elements of a in increasing order

May assume all a_i are distinct (otherwise, attach unique tags)

Assume the comparison model: primitives \langle , \rangle , no arithmetic or bit operations on a_i

Sequential work $O(n \log n)$ e.g. by mergesort

Parallel sorting based on an AKS sorting network

$$comp = O\left(\frac{n\log n}{p}\right)$$

$$comm = O\left(\frac{n\log n}{p}\right)$$

$$sync = O(\log n)$$

Parallel sorting by regular sampling Every processor [Shi, Schaeffer: 1992]

- reads subarray of a of size n/p and sorts it sequentially
- selects from it p samples from base index 0 at steps n/p^2

Samples define *p* equal-sized, contiguous blocks in local subarray

Parallel sorting by regular sampling Every processor [Shi, Schaeffer: 1992]

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- reads subarray of a of size n/p and sorts it sequentially
- selects from it p samples from base index 0 at steps n/p^2

Samples define p equal-sized, contiguous blocks in local subarray

A designated processor

- collects all p^2 samples and sorts them sequentially
- selects from them *p* splitters from base index 0 at steps *p*
- broadcasts the splitters

Splitters define *p* unequal-sized, rank-contiguous buckets in global array *a*

Further parallel algorithms Sorting

Parallel sorting by regular sampling (contd.)



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Parallel sorting by regular sampling (contd.)

Every processor

- receives the splitters and is assigned a bucket
- scans its subarray and sends each element to the appropriate bucket
- receives the elements of its bucket and sorts them sequentially
- writes the sorted bucket back to external memory

We will need to prove that bucket sizes, although not uniform, are still well-balanced ($\leq 2n/p$)

$$comp = O(\frac{n \log n}{p})$$
 $comm = O(n/p)$ $sync = O(1)$

Required slackness $n \ge p^3$
Further parallel algorithms Sorting

Parallel sorting by regular sampling (contd.)

Claim: each bucket has size $\leq 2n/p$



Further parallel algorithms Sorting

Parallel sorting by regular sampling (contd.)

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Claim: each bucket has size \leq 2n/p
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Proof (sketch). Relative to a fixed bucket B, a block b is

- low, if lower boundary of b is \leq lower boundary of B
- high otherwise

A bucket may only intersect

- ≤ 1 low block per processor, hence $\leq p$ low blocks overall
- $\leq p$ high blocks overall

Therefore, bucket size $\leq (p+p) \cdot n/p^2 = 2n/p$

The selection problem

Given $a = [a_0, \ldots, a_{n-1}]$, target rank k

Find k-th smallest element of a; e.g. median selection: k = n/2

As with sorting, we assume the comparison model

Sequential work $O(n \log n)$ by naive sorting

Sequential work O(n) by median sampling

[Blum+: 1973]

Selection by median sampling

[Blum+: 1973]

Proceed in rounds. In the first round:

- partition array a into subarrays of size 5
- in each subarray, select median e.g. by 5-element sorting
- select median-of-medians by recursion: $n \leftarrow n/5$, $k \leftarrow n/10$
- find rank / of median-of-medians in array a by linear search

If l = k, return a_l ; otherwise, eliminate elements on the wrong side of median-of-medians; adjust size and target rank for next round:

• if
$$l < k$$
, discard all $a_i \le a_l$; adjust $n \leftarrow n - l - 1$, $k \leftarrow k - l - 1$

• if l > k, discard all $a_i \ge a_l$; adjust $n \leftarrow l$, k unchanged

Subsequents rounds similar, with adjusted n, k

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Selection by median sampling (contd.)

Claim: Each round removes $\geq \frac{3n}{10}$ of elements of *a*

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Selection by median sampling (contd.) Claim: Each round removes $\geq \frac{3n}{10}$ of elements of a Proof (sketch). We have $\frac{n}{5}$ subarrays In at least $\frac{1}{2} \cdot \frac{n}{5}$ subarrays, subarray median $\leq a_{l}$ In every such subarray, three elements \leq subarray median $\leq a_l$ Hence, at least $\frac{1}{2} \cdot \frac{3n}{5} = \frac{3n}{10}$ elements $\leq a_{l}$ Symmetrically, at least $\frac{3n}{10}$ elements $\geq a_l$ Therefore, in a round, at least $\frac{3n}{10}$ elements are eliminated With each round, array shrinks exponentially $T(n) \leq T(\frac{n}{5}) + T(n - \frac{3n}{10}) + O(n) = T(\frac{2n}{10}) + T(\frac{7n}{10}) + O(n)$, therefore T(n) = O(n)

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Parallel selection by median sampling

In the first round, every processor

• reads a subarray of size n/p, selects the median

A designated processor

- collects all p subarray medians
- selects and broadcasts the median-of-medians

Every processor

• determines rank of median-of-medians in local subarray

Parallel selection by median sampling (contd.)

A designated processor

- adds up local ranks to determine global rank of median-of-medians
- compares it against target rank to determine direction of elimination
- broadcasts info on this direction

Every processor

- performs elimination on local subarray, discarding elements on wrong side of median-of-medians
- writes remaining elements
- $\leq 3n/4$ elements remain overall in array a

Subsequents rounds similar, with adjusted n, k

Parallel selection by median sampling (contd.)

Overall algorithm:

- perform $\log_{4/3} p$ rounds of median sampling and elimination, reducing original array to size n/p
- a designated processor collects the remaining array and performs selection sequentially

Parallel selection by median sampling (contd.)

Overall algorithm:

- perform $\log_{4/3} p$ rounds of median sampling and elimination, reducing original array to size n/p
- a designated processor collects the remaining array and performs selection sequentially

$$comp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(\log p)$

Parallel selection by regular sampling (generalised median sampling) In the first round, every processor

- reads a subarray of size n/p
- selects from it s = O(1) samples from base rank 0 at rank steps $\frac{n}{sp}$

Splitters define s equal-sized, rank-contiguous blocks in local subarray

A designated processor

- collects all *sp* samples
- selects from them s splitters from base rank 0 at rank steps p
- broadcasts the splitters

Splitters define *s* unequal-sized, rank-contiguous buckets in global array *a* Every processor

• determines rank of every splitter in local subarray

Alexander Tiskin (Warwick)

Parallel selection by regular sampling (contd.)

A designated processor

- adds up local ranks to determine global rank of every splitter
- compares these against target rank to determine target bucket
- broadcasts info on target bucket

Every processor

- performs elimination on subarray, discarding elements outside target bucket
- writes remaining elements
- $\leq 2n/s$ elements remain overall in array a

Subsequents rounds similar, with adjusted n, k

Parallel selection by accelerated regular sampling

In the original median sampling, sampling frequency s = 2 fixed across all rounds (samples at base rank 0 and local median rank $\frac{n}{2p}$); array shrinks exponentially

We now increase *s* from round to round, accelerating array reduction; array now shrinks superexponentially

Round 0: selecting samples and determining splitter ranks in time $O(\frac{n \log s}{p})$; set s = 2, time O(n/p)

Round 1: array size O(n/s), we can afford sampling frequency 2^s Round 2: ...

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Parallel selection by accelerated regular sampling

Overall algorithm:

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- perform O(log* p) rounds of regular sampling (with increasing frequency) and elimination, reducing original array to size n/p
- a designated processor collects the remaining array and performs selection sequentially

$$omp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(\log^* p)$

Parallel selection comp = O(n/p)comm = O(n/p) $sync = O(\log p)$ [Ishimizu+: 2002] $sync = O(\log \log n)$ [Fujiwara+: 2000] sync = O(1) whp randomised [Gerbessiotis, Siniolakis: 2003] $sync = O(\log \log p)$ [T: 2010] $sync = O(\log^* p)$ [T: NEW]

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Set $S \subseteq \mathbb{R}^d$ is convex, if for all x, y in S, every point between x and y is also in S

 $A \subseteq \mathbb{R}^d$

The convex hull conv A is the smallest convex set containing A

conv A is a polytope, defined by its vertices $A_i \in A$

Set A is in convex position, if every its point is a vertex of conv A

Definition of convexity requires arithmetic on coordinates, hence we assume the arithmetic model



d = 2

Fundamental arithmetic primitive: signed area of a triangle

Let
$$a_0 = (x_0, y_0)$$
, $a_1 = (x_1, y_1)$, $a_2 = (x_2, y_2)$

$$\Delta(a_0, a_1, a_2) = \frac{1}{2} \begin{vmatrix} x_0 & y_0 & 1 \\ x_1 & y_1 & 1 \\ x_2 & y_2 & 1 \end{vmatrix} = \frac{1}{2} ((x_1 - x_0)(y_2 - y_0) - (x_2 - x_0)(y_1 - y_0))$$

$$\Delta(a_0, a_1, a_2) \begin{cases} < 0 \text{ if } a_0, a_1, a_2 \text{ clockwise} \\ = 0 \text{ if } a_0, a_1, a_2 \text{ collinear} \\ > 0 \text{ if } a_0, a_1, a_2 \text{ counterclockwise} \end{cases}$$

An easy O(1) check: a_0 is to the left/right of directed line from a_1 to a_2 ? All of A is to the left of every edge of conv A, traversed counterclockwise

The (discrete) convex hull problem

Given $a = [a_0, \ldots, a_{n-1}]$, $a_i \in \mathbb{R}^d$

Output (a finite representation of) conv a

More precisely, output each k-dimensional face of conv a, $1 \le k < d$

E.g. in 3D: 1D vertices, 2D edges, 3D facets

Output must be structured, i.e. should give

- for *d* = 2, all vertex-edge incidence pairs; every vertex should "know" its both neighbours
- for general d, all incidence pairs between a k-D and a (k + 1)-D face

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The (discrete) convex hull problem (contd.)

Claim: Convex hull problem in \mathbb{R}^2 is at least as hard as sorting

Image: A matrix

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The (discrete) convex hull problem (contd.)

Claim: Convex hull problem in \mathbb{R}^2 is at least as hard as sorting

Proof. Let $x_0, \ldots, x_{n-1} \in \mathbb{R}$

To sort $[x_0, ..., x_{n-1}]$:

• compute conv $\left\{ (x_i, x_i^2) \in \mathbb{R}^2 : 0 \le i < n \right\}$

follow the edges to obtain sorted output

The (discrete) convex hull problem (contd.)

 $d=2: \leq n$ vertices, $\leq n$ edges, output size $\leq 2n$

d = 3: O(n) vertices, edges and facets, output size O(n)

d > 3: much bigger output...

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The (discrete) convex hull problem (contd.)

 $d=2: \leq n$ vertices, $\leq n$ edges, output size $\leq 2n$

d = 3: O(n) vertices, edges and facets, output size O(n)

d > 3: much bigger output...

For general d, conv a contains $O(n^{\lfloor d/2 \rfloor})$ faces of various dimensions

- d = 4,5: output size $O(n^2)$
- d = 6,7: output size $O(n^3)$

From now on, will concentrate on d = 2 and will sketch d = 3

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The (discrete) convex hull problem (contd.)

 $d=2: \leq n$ vertices, $\leq n$ edges, output size $\leq 2n$

d = 3: O(n) vertices, edges and facets, output size O(n)

d > 3: much bigger output...

For general d, conv a contains $O(n^{\lfloor d/2 \rfloor})$ faces of various dimensions

d = 4,5: output size $O(n^2)$

d = 6,7: output size $O(n^3)$

From now on, will concentrate on d = 2 and will sketch d = 3Sequential work $O(n \log n)$: Graham's scan (2D); mergehull (2D, 3D) '

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 $A \subseteq \mathbb{R}^d$ Let $0 \le \epsilon \le 1$

Set $E \subseteq A$ is an ϵ -net for A, if any halfspace with no points in E covers $\leq \epsilon |A|$ points in A

An ϵ -net may always be assumed to be in convex position

 $A \subseteq \mathbb{R}^d$ Let $0 \le \epsilon \le 1$

Set $E \subseteq A$ is an ϵ -net for A, if any halfspace with no points in E covers $\leq \epsilon |A|$ points in A

An ϵ -net may always be assumed to be in convex position

Set $E \subseteq A$ is an ϵ -approximation for A, if for all α , $0 \le \alpha \le 1$, any halfspace with $\alpha |E|$ points in E covers $(\alpha \pm \epsilon)|A|$ points in A

An ϵ -approximation may not be in convex position

Both are easy to construct in 2D, much harder in 3D and higher

Claim. An ϵ -approximation for A is an ϵ -net for A

The converse does not hold!

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Claim. An ϵ -approximation for A is an ϵ -net for A

The converse does not hold!

Claim. Union of ϵ -approximations for A', A'' is ϵ -approximation for $A'' \cup A''$

Claim. An ϵ -approximation for A is an ϵ -net for A

- The converse does not hold!
- Claim. Union of ϵ -approximations for A', A'' is ϵ -approximation for $A'' \cup A''$
- Claim. An ϵ -net for a δ -approximation for A is an $(\epsilon + \delta)$ -net for A

Claim. An ϵ -approximation for A is an ϵ -net for A

The converse does not hold!

Claim. Union of ϵ -approximations for A', A'' is ϵ -approximation for $A'' \cup A''$

Claim. An ϵ -net for a δ -approximation for A is an ($\epsilon + \delta$)-net for A

Proofs: Easy by definitions, independently of d.

$$d=2$$
 $A\subseteq \mathbb{R}^2$ $|A|=n$ $\epsilon=1/r$ $r\geq 1$

Claim. A 1/r-net for A of size $\leq 2r$ exists and can be computed in sequential work $O(n \log n)$.

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d=2 $A\subseteq \mathbb{R}^2$ |A|=n $\epsilon=1/r$ $r\geq 1$

Claim. A 1/r-net for A of size $\leq 2r$ exists and can be computed in sequential work $O(n \log n)$.

Proof. Consider convex hull of A and an arbitrary interior point vPartition A into triangles: base at a hull edge, apex at v

A triangle is heavy if it contains > n/r points of A, otherwise light

d=2 $A\subseteq \mathbb{R}^2$ |A|=n $\epsilon=1/r$ $r\geq 1$

Claim. A 1/r-net for A of size $\leq 2r$ exists and can be computed in sequential work $O(n \log n)$.

Proof. Consider convex hull of A and an arbitrary interior point vPartition A into triangles: base at a hull edge, apex at vA triangle is heavy if it contains > n/r points of A, otherwise light Heavy triangles: for each triangle, put both hull vertices into E

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d=2 $A\subseteq \mathbb{R}^2$ |A|=n $\epsilon=1/r$ $r\geq 1$

Claim. A 1/r-net for A of size $\leq 2r$ exists and can be computed in sequential work $O(n \log n)$.

Proof. Consider convex hull of A and an arbitrary interior point v Partition A into triangles: base at a hull edge, apex at v A triangle is heavy if it contains > n/r points of A, otherwise light Heavy triangles: for each triangle, put both hull vertices into E Light triangles: for each triangle chain, greedy next-fit bin packing

- combine adjacent triangles into bins with $\leq n/r$ points
- for each bin, put both boundary hull vertices into E

In total $\leq 2r$ heavy triangles and bins, hence $|E| \leq 2r$

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d=2 $A\subseteq \mathbb{R}^2$ |A|=n $\epsilon=1/r$

Claim. If A is in convex position, then a 1/r-approximation for A of size $\leq r$ exists and can be computed in sequential work $O(n \log n)$.



d = 2 $A \subseteq \mathbb{R}^2$ |A| = n $\epsilon = 1/r$

Claim. If A is in convex position, then a 1/r-approximation for A of size $\leq r$ exists and can be computed in sequential work $O(n \log n)$.

Proof. Sort points of A in circular order they appear on the convex hull Put every n/r-th point into E. We have $|E| \le r$. Parallel 2D hull computation by generalised regular sampling

 $a = [a_0, \ldots, a_{n-1}]$ $a_i \in \mathbb{R}^2$

Every processor

- $\bullet\,$ reads a subset of n/p points, computes its hull, discards the rest
- selects *p* samples at regular intervals on the hull

Set of all samples: 1/p-approximation for set *a* (after discarding local interior points)
Parallel 2D hull computation by generalised regular sampling

 $a = [a_0, \ldots, a_{n-1}]$ $a_i \in \mathbb{R}^2$

Every processor

- $\bullet\,$ reads a subset of n/p points, computes its hull, discards the rest
- selects *p* samples at regular intervals on the hull

Set of all samples: 1/p-approximation for set *a* (after discarding local interior points)

A designated processor

- collects all p^2 samples (and does not compute its hull)
- selects from the samples a 1/p-net of $\leq 2p$ points as splitters

Set of splitters: 1/p-net for samples, therefore a 2/p-net for set a

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Parallel 2D hull computation by generalised regular sampling (contd.)

The 2p splitters can be assumed to be in convex position (like any ϵ -net), and therefore define a splitter polygon with at most 2p edges

Each vertex of splitter polygon defines a **bucket**: the subset of set *a* visible when sitting at this vertex (assuming the polygon is opaque)

Each bucket can be covered by two half-planes not containg any splitters. Therefore, bucket size is at most $2 \cdot (2/p) \cdot n = 4n/p$.

Parallel 2D hull computation by generalised regular sampling (contd.) The designated processor broadcasts the splitters

Every processor

- receives the splitters and is assigned 2 buckets
- scans its hull and sends each point to the appropriate bucket
- receives the points of its buckets and computes their hulls sequentially
- writes the bucket hulls back to external memory

$$comp = O(\frac{n \log n}{p}) \qquad comm = O(n/p) \qquad sync = O(1)$$

Requires slackness $n \ge p^3$

Further parallel algorithms Convex hull

d = 3 $A \subseteq \mathbb{R}^3$ |A| = n $\epsilon = 1/r$

Claim. A 1/r-net for A of size O(r) exists and can be computed in sequential work $O(rn \log n)$.

Proof: [Brönnimann, Goodrich: 1995]

Further parallel algorithms Convex hull

d = 3 $A \subseteq \mathbb{R}^3$ |A| = n $\epsilon = 1/r$

Claim. A 1/r-net for A of size O(r) exists and can be computed in sequential work $O(rn \log n)$.

Proof: [Brönnimann, Goodrich: 1995]

Claim. A 1/r-approximation for A of size $O(r^3(\log r)^{O(1)})$ exists and can be computed in sequential work $O(n \log r)$.

Proof: [Matoušek: 1992]

Better approximations are possible, but are slower to compute

Parallel 3D hull computation by generalised regular sampling

 $a = [a_0, \ldots, a_{n-1}]$ $a_i \in \mathbb{R}^3$

Every processor

- reads a subset of n/p points
- selects a 1/p-approximation of $O(p^3(\log p)^{O(1)})$ points as samples

Set of all samples: 1/p-approximation for set a

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Parallel 3D hull computation by generalised regular sampling

 $a = [a_0, \ldots, a_{n-1}]$ $a_i \in \mathbb{R}^3$

Every processor

- reads a subset of n/p points
- selects a 1/p-approximation of $O(p^3(\log p)^{O(1)})$ points as samples

Set of all samples: 1/p-approximation for set a

A designated processor

- collects all $O(p^4(\log p)^{O(1)})$ samples
- selects from the samples a 1/p-net of O(p) points as splitters

Set of splitters: 1/p-net for samples, therefore a 2/p-net for set a

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Parallel 3D hull computation by generalised regular sampling (contd.)

The O(p) splitters can be assumed to be in convex position (like any ϵ -net), and therefore define a splitter polytope with O(p) edges

Each edge of splitter polytope defines a **bucket**: the subset of *a* visible when sitting on this edge (assuming the polytope is opaque)

Each bucket can be covered by two half-spaces not containg any splitters. Therefore, bucket size is at most $2 \cdot (2/p) \cdot n = 4n/p$.

Parallel 3D hull computation by generalised regular sampling (contd.) The designated processor broadcasts the splitters

Every processor

- receives the splitters and is assigned a bucket
- scans its hull and sends each point to the appropriate bucket
- receives the points of its bucket and computes their convex hull sequentially
- writes the bucket hull back to external memory

$$comp = O(\frac{n \log n}{p})$$
 $comm = O(n/p)$ $sync = O(1)$

Requires slackness $n \gg p$

The suffix sorting problem

Given string $a = a_0 \dots a_{n-1}$ \$

 $a_i \in \{0, 1, \dots, n-1\}$ $0 \le i < n$ $\$ = -\infty$ is a sentinel

Sort all suffixes of a in lexicographic order (implicitly, by returning ranks) Character sorting: time O(n) e.g. by counting sort

Naive suffix sorting: time $O(n^2)$ by *n*-fold radix sort, performing character sorting successively in every position from least to most significant

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Suffix sorting by DC mod 3 sampling Difference cover (DC) modulo 3, aka skew algorithm [Kärkkäinen, Sanders: 2003] Denote *a_i* by [*i*]

Consider 3-substrings as super-characters: [012], [123], [234], ... Sort all super-characters by 3-fold radix sort; replace each by its rank

Call indices 3i, 3i + 1 (but not 3i + 2) for any *i* sample indices

Sample indices define

- sample suffixes: [012...], [123...], (not [234...]), [345...], ...
- sample super-characters: [012], [123], (not [234]), [345], ...

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Suffix sorting by DC mod 3 sampling (contd.)

```
b = [012][345][678]...\$[123][456][789]...\$[345][678][9 \ 10 \ 11]...\$
```

b is composed of sample suffixes of *a*, each broken up into sample super-characters; overall, *b* is of length 2n/3 super-characters

Sort sample suffixes

• suffix sorting on b by recursion

Sort non-sample suffixes

• 2-fold radix sort on a at non-sample indices

[234...] = [2][345...][567...] = [5][678...]

. . .

Suffix sorting by DC mod 3 sampling (contd.)

We have two sorted sets of suffixes:

- sample [012...], [123...], (not [234...]), [345...], [456...], (not [567...]), ...
- non-sample [234...], [567...], . . .

Perform comparison-based merging of suffix sets

Each comparison: either $[3i \dots]$ or $[3i+1 \dots]$ vs $[3i+2 \dots]$, performed as

- [3*i*][3*i*+1 3*i*+2 ...] vs [3*j*+2][3*j*+3 3*j*+4 ...]
- [3*i*+1 3*i*+2][3*i*+3 3*i*+4 ...] vs [3*j*+2 3*j*+3][3*j*+4 3*j*+5 ...]

Comparing pairs of the form (sample super-character, sample suffix) Comparison time O(1)

Overall running time
$$T(n) = O(n) + T(2n/3) = O(n)$$

Parallel suffix sorting by DC mod 3 sampling

 $a = a_0 \dots a_{n-1}$ \$

At the top recursion level, every processor

- reads substring of a of length n/p
- sorts super-characters locally by 3-fold radix sort

The processors collectively

- sort super-characters globally by regular sampling
- form string b
- sort sample suffixes of *a* by recursion on *b*
- sort non-sample suffixes of a by 2-fold radix sort at non-sample indices

Parallel suffix sorting by DC mod 3 sampling (contd.)

Every processor

• merges sample vs non-sample suffixes locally

The processors collectively

• merge sample vs non-sample suffixes globally by regular sampling

Subsequent recursion levels similar, with n adjusted

Parallel suffix sorting by DC mod 3 sampling (contd.)

Overall algorithm:

- perform $\log_{3/2} p$ recursion levels of suffix sorting by DC mod 3 sampling, obtaining a string of length n/p
- a designated processor collects the resulting string and performs suffix sorting sequentially

Parallel suffix sorting by DC mod 3 sampling (contd.)

Overall algorithm:

- perform $\log_{3/2} p$ recursion levels of suffix sorting by DC mod 3 sampling, obtaining a string of length n/p
- a designated processor collects the resulting string and performs suffix sorting sequentially

$$comp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(\log p)$

Further parallel algorithms Suffix sorting

Suffix sorting by DC mod d sampling

Difference cover (DC) modulo d: set S of integers mod d, such that for all $i \mod d$, there are $j, k \in S$ with $k - j = i \mod d$

Examples:

4 3 4 3 4 3 4

Suffix sorting by DC mod d sampling (contd.) Claim: For any d, there is a DC mod d of size $O(d^{1/2})$ [Colbourn, Ling: 2000]

DC mod 3 algorithm can be generalised to DC mod d for any $d \ge 3$ [Kärkkäinen, Sanders: 2003]

Given *d*, consider *d*-substrings as super-characters

Sort all super-characters by *d*-fold radix sort; replace each by its rank

Fix a DC mod *d* as sample indices

Sample indices define sample suffixes, sample super-characters

Suffix sorting by DC mod *d* sampling (contd.)

Form string *b*, composed of sample suffixes of *a*, each broken up into sample super-characters; overall length of *b* is $O(n/d^{1/2})$ super-characters

Sort sample suffixes

• suffix sorting on b by recursion

Sort non-sample suffixes in < d separate subsets according to index mod d

• 2-fold radix sort on *a* for each non-sample index mod *d*

Suffix sorting by DC mod *d* sampling (contd.)

We have $\leq d$ ordered sets of suffixes:

- sample suffixes
- < d subsets of non-sample suffixes according to index mod d

Perform $\leq d$ -way comparison-based merging of suffix sets Comparing pairs of the form (sample super-character, sample suffix) Comparison time O(1)

Overall running time $T(n) = O(nd) + T(O(n/d^{1/2})) = O(nd)$

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Parallel suffix sorting by accelerated DC mod *d* sampling

In parallel DC mod 3 sampling, modulus d = 3 fixed across all levels; string shrinks exponentially

We now increase modulus from each recursion level to the next, accelerating string reduction; string shrinks superexponentially, allowing further increase in modulus while keeping work $O(size \cdot modulus) = O(n)$

Level 0: array size *n*; can only afford DC mod d = O(1)

Level 1: array size $O(\frac{n}{d^{1/2}})$; can now afford DC mod $d^{3/2}$

Level 2: array size $O(\frac{n}{d^{1/2} \cdot d^{3/4}}) = O(\frac{n}{d^{5/4}})$; can now afford DC mod $d^{9/4}$ Level 3: array size $O(\frac{n}{d^{5/4} \cdot d^{9/8}}) = O(\frac{n}{d^{19/8}})$; ...

Level $O(\log \log p)$: array size O(n/p)

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Parallel suffix sorting by accelerated DC mod *d* sampling Overall algorithm:

- perform O(log log p) recursion levels of suffix sorting by DC mod d sampling (with increasing d), obtaining a string of length n/p
- a designated processor collects the resulting string and performs suffix sorting sequentially

Parallel suffix sorting by accelerated DC mod *d* sampling Overall algorithm:

- perform O(log log p) recursion levels of suffix sorting by DC mod d sampling (with increasing d), obtaining a string of length n/p
- a designated processor collects the resulting string and performs suffix sorting sequentially

$$comp = O(n/p)$$
 $comm = O(n/p)$ $sync = O(\log \log p)$

Computation by circuits

- 2 Parallel computation models
- 3 Basic parallel algorithms
- 4 Further parallel algorithms
- 5 Parallel matrix algorithms
 - 6 Parallel graph algorithms

Matrix-vector multiplication

The matrix-vector multiplication problem

 $A \cdot b = c$ A: *n*-matrix; *b*, *c*: *n*-vectors

Given A, B, compute C

 $c_i = \sum_j A_{ij} \cdot b_j \quad 0 \leq i, j < n$



Matrix-vector multiplication

The matrix-vector multiplication problem

$$A \cdot b = c$$
 A: *n*-matrix; b, c: *n*-vectors

Given A, B, compute C

$$c_i = \sum_j A_{ij} \cdot b_j \quad 0 \le i, j < n$$



Consider elements of b as inputs and of c as outputs

Elements of A are considered to be problem parameters, do not count as inputs (motivation: iterative linear algebra methods)

Overall, n^2 elementary products $A_{ij} \cdot b_j = c_j^i$ Sequential work $O(n^2)$

Matrix-vector multiplication

The matrix-vector multiplication circuit

 $c \leftarrow 0$ For all *i*, *j*: $c_i \stackrel{+}{\leftarrow} c_j^i \leftarrow A_{ij} \cdot b_j$ (adding each c_j^i to c_i asynchronously) *n* input nodes of outdegree *n*, one per element of *b* n^2 computation nodes of in- and outdegree 1, one per elementary product

n output nodes of indegree *n*, one per element of *c* A size $O(n^2)$, depth O(1)



Matrix-vector multiplication

Parallel matrix-vector multiplication

Partition computation nodes into a regular grid of $p = p^{1/2} \cdot p^{1/2}$ square $\frac{n}{p^{1/2}}$ -blocks

Matrix A gets partitioned into p square $\frac{n}{p^{1/2}}$ -blocks A_{IJ} ($0 \le I, J < p^{1/2}$)

Vectors *b*, *c* each gets partitioned into $p^{1/2}$ linear $\frac{n}{p^{1/2}}$ -blocks *b*_J, *c*_I

Overall, *p* block products $A_{IJ} \cdot b_J = c_I^J$

$$c_I = \sum_{0 \leq J < p^{1/2}} c_I^J$$
 for all I

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Matrix-vector multiplication

Parallel matrix-vector multiplication (contd.)

 $c \leftarrow 0$ For all *I*, *J*: $c_I \stackrel{+}{\leftarrow} c_I^J \leftarrow A_{IJ} \cdot b_J$



Matrix-vector multiplication

Parallel matrix-vector multiplication (contd.)

Initialise $c \leftarrow 0$ in external memory



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Matrix-vector multiplication

Parallel matrix-vector multiplication (contd.)

```
Initialise c \leftarrow 0 in external memory
```

Every processor

- is assigned I, J and block A_{IJ}
- reads block b_J and computes $c_I^J \leftarrow A_{IJ} \cdot b_J$
- updates $c_I \stackrel{+}{\leftarrow} c_I^J$ in external memory
- concurrent writing resolved by operator + (recall concurrent block writing by array combining)

$$comp = O\left(\frac{n^2}{p}\right)$$

$$comm = O\left(\frac{n}{p^{1/2}}\right)$$

$$sync = O(1)$$

Slackness required $n \geq p$ (as $\frac{n}{p^{1/2}} \geq p^{1/2}$ needed for concurrent write)

Matrix multiplication

The matrix multiplication problem

 $A \cdot B = C$ A, B, C: *n*-matrices Given A, B, compute C $C_{ik} = \sum_{j} A_{ij} \cdot B_{jk}$ $0 \le i, j, k < n$



Image: Image:

4 3 4 3 4 3 4

Matrix multiplication

The matrix multiplication problem

 $A \cdot B = C \quad A, B, C: n-\text{matrices}$ Given A, B, compute C $C_{ik} = \sum_{j} A_{ij} \cdot B_{jk}$ $0 \le i, j, k \le n$ $B = \begin{bmatrix} C \\ C \end{bmatrix}$

Overall, n^3 elementary products $A_{ij} \cdot B_{jk} = C_{ik}^j$ Sequential work $O(n^3)$

Matrix multiplication

The matrix multiplication circuit

 $C_{ik} \leftarrow 0$ For all *i*, *j*, *k*: $C_{ik} \stackrel{+}{\leftarrow} C^{j}_{ik} \leftarrow A_{ij} \cdot B_{jk}$ (adding each C^{j}_{ik} to C_{ik} asynchronously) 2*n* input nodes of outdegree *n*, one per element of *A*, *B*

 n^2 computation nodes of in- and outdegree 1, one per elementary product

n output nodes of indegree n, one per element of C

size $O(n^3)$, depth O(1)



Parallel matrix multiplication

Partition computation nodes into a regular grid of $p = p^{1/3} \cdot p^{1/3} \cdot p^{1/3}$ cubic $\frac{n}{p^{1/3}}$ -blocks

Matrices A, B, C each gets partitioned into $p^{2/3}$ square $\frac{n}{p^{1/2}}$ -blocks A_{IJ} , B_{JK} , C_{IK} ($0 \le I, J, K < p^{1/3}$)

Overall, *p* block products $A_{IJ} \cdot B_{JK} = C_{IK}^J$

 $C_{IK} = \sum_{0 \leq J < p^{1/2}} C^J_{IK}$ for all I, K

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Matrix multiplication

Parallel matrix multiplication (contd.)

 $\begin{array}{l} C \leftarrow 0 \\ \text{For all } I, \ J, \ K: \ C_{IK} \stackrel{+}{\leftarrow} C_{IK}^J \leftarrow A_{IJ} \cdot B_{JK} \end{array}$



Matrix multiplication

Parallel matrix multiplication (contd.) Initialise $C \leftarrow 0$ in external memory

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Matrix multiplication

Parallel matrix multiplication (contd.)

Initialise $C \leftarrow 0$ in external memory

Every processor

- is assigned I, J, K
- reads blocks A_{IJ} , B_{JK} , and computes $C_{IK}^J \leftarrow A_{IJ} \cdot B_{JK}$
- updates $C_{IK} \stackrel{+}{\leftarrow} C^J_{IK}$ in external memory
- concurrent writing resolved by operator + (recall concurrent block writing by array combining)

Matrix multiplication

Parallel matrix multiplication (contd.)

```
Initialise C \leftarrow 0 in external memory
```

Every processor

- is assigned I, J, K
- reads blocks A_{IJ} , B_{JK} , and computes $C_{IK}^J \leftarrow A_{IJ} \cdot B_{JK}$
- updates $C_{IK} \stackrel{+}{\leftarrow} C^J_{IK}$ in external memory
- concurrent writing resolved by operator + (recall concurrent block writing by array combining)

$$comp = O\left(\frac{n^3}{p}\right)$$

$$comm = O\left(\frac{n^2}{p^{2/3}}\right)$$

$$sync = O(1)$$

Slackness required $n \ge p^{2/3}$ (as $\frac{n}{p^{1/3}} \ge p^{1/3}$ needed for concurrent write)

Matrix multiplication

Theorem. Computing the matrix multiplication dag requires communication $\Omega\bigl(\frac{n^2}{p^{2/3}}\bigr)$ per processor

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Matrix multiplication

Theorem. Computing the matrix multiplication dag requires communication $\Omega(\frac{n^2}{p^{2/3}})$ per processor

Proof: (discrete) volume vs surface area

Let V be the subset of nodes computed by a certain processor

For at least one processor: $|V| \ge \frac{n^3}{p}$

Let A, B, C be projections of V onto coordinate planes

Matrix multiplication

Theorem. Computing the matrix multiplication dag requires communication $\Omega(\frac{n^2}{n^{2/3}})$ per processor

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Let V be the subset of nodes computed by a certain processor

For at least one processor: $|V| \ge \frac{n^3}{p}$

Let A, B, C be projections of V onto coordinate planes

Arithmetic vs geometric mean: $|A| + |B| + |C| \ge 3(|A| \cdot |B| \cdot |C|)^{1/3}$ Loomis–Whitney inequality: $|A| \cdot |B| \cdot |C| \ge |V|^2$ Matrix multiplication

Theorem. Computing the matrix multiplication dag requires communication $\Omega(\frac{n^2}{p^{2/3}})$ per processor

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Arithmetic vs geometric mean: $|A| + |B| + |C| \ge 3(|A| \cdot |B| \cdot |C|)^{1/3}$ Loomis–Whitney inequality: $|A| \cdot |B| \cdot |C| \ge |V|^2$

We have
$$comm \ge |A| + |B| + |C| \ge 3(|A| \cdot |B| \cdot |C|)^{1/3} \ge 3|V|^{2/3} \ge 3(\frac{n^3}{p})^{2/3} = \frac{3n^2}{p^{2/3}}$$
, hence $comm = \Omega(\frac{n^2}{p^{2/3}})$

Note that this is not conditioned on $comp = O(\frac{n^3}{p})$

The optimality theorem only applies to matrix multiplication by the specific $O(n^3)$ -node dag

Includes e.g. the following forms of matrix multiplication:

- numerical, with only operators $+, \cdot$ allowed (not operator -)
- Boolean, with only operators ∨, ∧ allowed (not if/then)

Parallel matrix algorithms Fast matrix multiplication

2-matrix multiplication: standard circuit

$$A \cdot B = C \qquad A = \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix} \quad B = \begin{bmatrix} B_{00} & B_{01} \\ B_{10} & B_{11} \end{bmatrix} \quad C = \begin{bmatrix} C_{00} & C_{01} \\ C_{10} & C_{11} \end{bmatrix}$$
$$C_{\underline{00}} = A_{\underline{00}} \cdot B_{\underline{00}} + A_{\underline{01}} \cdot B_{\underline{10}} \qquad C_{01} = A_{00} \cdot B_{01} + A_{01} \cdot B_{11}$$
$$C_{\underline{10}} = A_{\underline{10}} \cdot B_{\underline{00}} + A_{\underline{11}} \cdot B_{\underline{10}} \qquad C_{11} = A_{10} \cdot B_{01} + A_{11} \cdot B_{11}$$

 A_{00}, \ldots : either ordinary elements or blocks; 8 multiplications

Image: A matrix

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Fast matrix multiplication

2-matrix multiplication: Strassen's circuit

$$A \cdot B = C$$
 $A = \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$ $B = \begin{bmatrix} B_{00} & B_{01} \\ B_{10} & B_{11} \end{bmatrix}$ $C = \begin{bmatrix} C_{00} & C_{01} \\ C_{10} & C_{11} \end{bmatrix}$

Image: A matrix

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Parallel matrix algorithms Fast matrix multiplication

2-matrix multiplication: Strassen's circuit

$$A \cdot B = C \qquad A = \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix} \quad B = \begin{bmatrix} B_{00} & B_{01} \\ B_{10} & B_{11} \end{bmatrix} \quad C = \begin{bmatrix} C_{00} & C_{01} \\ C_{10} & C_{11} \end{bmatrix}$$

Let A, B, C be over a ring: operators +, -, · allowed on elements
$$D^{(0)} = (A_{00} + A_{11}) \cdot (B_{00} + B_{11})$$

$$D^{(1)} = (A_{10} + A_{11}) \cdot B_{00} \qquad D^{(2)} = A_{00} \cdot (B_{01} - B_{11})$$

$$D^{(3)} = A_{11} \cdot (B_{10} - B_{00}) \qquad D^{(4)} = (A_{00} + A_{01}) \cdot B_{11}$$

$$D^{(5)} = (A_{10} - A_{00}) \cdot (B_{00} + B_{01}) \qquad D^{(6)} = (A_{01} - A_{11}) \cdot (B_{10} + B_{11})$$

$$C_{\underline{00}} = D^{(0)} + D^{(3)} - D^{(4)} + D^{(6)} \qquad C_{\underline{01}} = D^{(2)} + D^{(4)}$$

$$C_{\underline{10}} = D^{(1)} + D^{(3)} \qquad C_{\underline{11}} = D^{(0)} - D^{(1)} + D^{(2)} + D^{(5)}$$

 A_{00} , ...: either ordinary elements or square blocks; 7 multiplications

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N-matrix multiplication: bilinear circuit

- certain R linear combinations of elements of A
- certain R linear combinations of elements of B
- R pairwise products of these combinations
- \bullet certain N^2 linear combinations of these products, each giving an element of C

Bilinear circuits for matrix multiplication:

- standard: N = 2, R = 8, combinations trivial
- Strassen: N = 2, R = 7, combinations highly surprising!
- sub-Strassen: N > 2, N² < R < N^{log_27} \approx N^{2.81}

Elements of A, B, C: either ordinary elements or square blocks

Block-recursive matrix multiplication

Given a scheme: bilinear circuit with fixed N, R

Let A, B, C be *n*-matrices, $n \ge N$ $A \cdot B = C$

Partition each of A, B, C into an $N \times N$ regular grid of n/N-blocks

Apply the scheme, treating

- each '+' as block '+', each '-' as block '-'
- each '·' as recursive call on blocks

Block-recursive matrix multiplication

Given a scheme: bilinear circuit with fixed N, R

Let A, B, C be *n*-matrices, $n \ge N$ $A \cdot B = C$

Partition each of A, B, C into an $N \times N$ regular grid of n/N-blocks Apply the scheme, treating

- each '+' as block '+', each '-' as block '-'
- each '·' as recursive call on blocks

Resulting recursive bilinear circuit:

• size
$$O(n^{\omega})$$
, where $\omega = \log_N R < \log_N N^3 = 3$

• depth $\approx 2 \log n$

Sequential work $O(n^{\omega})$

Block-recursive matrix multiplication (contd.)

Historical improvements in block-recursive matrix multiplication:

Ν	N ³	R	$\omega = \log_N R$	
2	8	8	3	standard
2	8	7	2.81	[Strassen: 1969]
3	27	23	2.85 > 2.81	
5	125	100	2.86 > 2.81	
48	110592	47216	2.78	[Pan: 1978]
HUGE	HUGE	HUGE	2.3755	[Coppersmith, Winograd: 1987]
HUGE	HUGE	HUGE	2.3737	[Stothers: 2010]
HUGE	HUGE	HUGE	2.3727	[Vassilevska–Williams: 2011]
?	?	?	?	

Image: A matrix

3 1 4 3 1

Block-recursive matrix multiplication (contd.)

Circuit size is determined by the scheme parameters N, R; the number of operations in scheme's linear combinations turns out to be irrelevant

Optimal circuit size unknown: only near-trivial lower bound $\Omega(n^2 \log n)$

At each level of the recursion tree, the R recursive calls are independent, hence the recursion tree can be computed breadth-first

At recursion level k:

• *R^k* independent block multiplication subproblems

In particular, at level $\log_R p$:

• *p* independent block multiplication subproblems, therefore each subproblem can be solved sequentially on an arbitrary processor

In recursion levels 0 to $\log_R p$, need to compute elementwise linear combinations on distributed matrices

Assigning matrix elements to processors:

- partition A into regular $\frac{n}{p^{1/\omega}}$ -blocks
- distribute each block evenly and identically across processors
- partition *B*, *C* analogously (distribution identical across all blocks of the same matrix, need not be identical across different matrices)

In recursion levels 0 to $\log_R p$, need to compute elementwise linear combinations on distributed matrices

Assigning matrix elements to processors:

- partition A into regular $\frac{n}{p^{1/\omega}}$ -blocks
- distribute each block evenly and identically across processors
- partition *B*, *C* analogously (distribution identical across all blocks of the same matrix, need not be identical across different matrices)

E.g. cyclic distribution

Linear combinations of matrix blocks in recursion levels 0 to $\log_R p$ can now be computed without communication Parallel block-recursive matrix multiplication (contd.) Each processor inputs its assigned elements of A, BDownsweep of recursion tree, levels 0 to $\log_R p$:

• linear combinations of blocks of A, B, no communication

Each processor inputs its assigned elements of A, B

Downsweep of recursion tree, levels 0 to $\log_R p$:

• linear combinations of blocks of A, B, no communication

Recursion levels below $\log_R p$: p block multiplication subproblems

- assign each subproblem to a different processor
- a processor collects its subproblem's two input blocks, solves it sequentially, then redistributes the subproblem's output block

Each processor inputs its assigned elements of A, B

Downsweep of recursion tree, levels 0 to $\log_R p$:

• linear combinations of blocks of A, B, no communication

Recursion levels below log_R p: p block multiplication subproblems

- assign each subproblem to a different processor
- a processor collects its subproblem's two input blocks, solves it sequentially, then redistributes the subproblem's output block

Upsweep of recursion tree, levels $\log_R p$ to 0:

• linear combinations giving blocks of C, no communication

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Each processor inputs its assigned elements of A, B

Downsweep of recursion tree, levels 0 to $\log_R p$:

• linear combinations of blocks of A, B, no communication

Recursion levels below log_R p: p block multiplication subproblems

- assign each subproblem to a different processor
- a processor collects its subproblem's two input blocks, solves it sequentially, then redistributes the subproblem's output block

Upsweep of recursion tree, levels $\log_R p$ to 0:

• linear combinations giving blocks of C, no communication

Each processor outputs its assigned elements of C

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Each processor inputs its assigned elements of A, B

Downsweep of recursion tree, levels 0 to $\log_R p$:

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Recursion levels below log_R p: p block multiplication subproblems

- assign each subproblem to a different processor
- a processor collects its subproblem's two input blocks, solves it sequentially, then redistributes the subproblem's output block

Upsweep of recursion tree, levels $\log_R p$ to 0:

• linear combinations giving blocks of C, no communication

Each processor outputs its assigned elements of C

$$comp = O\left(\frac{n^{\omega}}{p}\right)$$

$$comm = O(\frac{n^2}{p^{2/\omega}})$$

$$sync = O(1)$$

Alexander Tiskin (Warwick)

Fast matrix multiplication

Theorem. Computing the block-recursive matrix multiplication dag requires communication $\Omega(\frac{n^2}{p^{2/\omega}})$ per processor [Ballard+:2012]



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Fast matrix multiplication

Theorem. Computing the block-recursive matrix multiplication dag requires communication $\Omega(\frac{n^2}{p^{2/\omega}})$ per processor [Ballard+:2012] Proof: generalises the Loomis–Whitney inequality using graph expansion

(details omitted)

Boolean matrix multiplication

Boolean matrix multiplication

Let A, B, C be Boolean *n*-matrices: ' \lor ', ' \land ', 'if/then' allowed on elements

 $A \wedge B = C$

 $C_{ik} = \bigvee_{i} A_{ik} \wedge B_{jk} \qquad 0 \le i, j, k < n$

Overall, n^3 elementary products $A_{ij} \wedge B_{jk}$

Sequential work $O(n^3)$ bit operations

BSP costs in bit operations:

$$comp = O\left(\frac{n^3}{p}\right)$$
 $comm = O\left(\frac{n^2}{p^{2/3}}\right)$ $sync = O(1)$

Boolean matrix multiplication

Fast Boolean matrix multiplication

 $A \wedge B = C$

Convert A, B into integer matrices by treating 0, 1 as integers (requires if/then on elements)

Compute $A \cdot B = C$ modulo n + 1 using a Strassen-like algorithm

Convert C into a Boolean matrix by evaluating $C_{jk} \neq 0 \mod n+1$ Sequential work $O(n^{\omega})$

BSP costs:

$$comp = O\left(\frac{n^{\omega}}{p}\right)$$
 $comm = O\left(\frac{n^2}{p^{2/\omega}}\right)$ $sync = O(1)$

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Boolean matrix multiplication

Parallel Boolean matrix multiplication by regular decomposition

The following algorithm is impractical, but of theoretical interest, because it beats the generic Loomis–Whitney communication lower bound

Parallel Boolean matrix multiplication by regular decomposition

The following algorithm is impractical, but of theoretical interest, because it beats the generic Loomis–Whitney communication lower bound

Regularity Lemma: in a Boolean matrix, the rows and the columns can be
partitioned into K (almost) equal-sized subsets, so that K^2 resulting
submatrices are random-like (of various densities)[Szemerédi: 1978]

Parallel Boolean matrix multiplication by regular decomposition

The following algorithm is impractical, but of theoretical interest, because it beats the generic Loomis–Whitney communication lower bound

Regularity Lemma: in a Boolean matrix, the rows and the columns can be
partitioned into K (almost) equal-sized subsets, so that K^2 resulting
submatrices are random-like (of various densities)[Szemerédi: 1978]

 $K = K(\epsilon)$, where ϵ is the "degree of random-likeness"

Function $K(\epsilon)$ grows enormously as $\epsilon \rightarrow 0$, but is independent of *n*



We shall call this the regular decomposition of a Boolean matrix

Parallel Boolean matrix multiplication by regular decomposition (contd.) $A \wedge B = C$

If A, B, C random-like, then either A or B has few 1s, or C has few 0s Equivalently, at least one of A, B, \overline{C} has few 1s, i.e. is sparse Fix ϵ so that "sparse" means density < 1/p Parallel Boolean matrix multiplication by regular decomposition (contd.) By Regularity Lemma, we have the three-way regular decomposition

• $A^{(1)} \wedge B^{(1)} = C^{(1)}$, where $A^{(1)}$ is sparse

•
$$\mathcal{A}^{(2)} \wedge \mathcal{B}^{(2)} = \mathcal{C}^{(2)}$$
, where $\mathcal{B}^{(2)}$ is sparse

•
$$A^{(3)} \wedge B^{(3)} = C^{(3)}$$
, where $\overline{C^{(3)}}$ is sparse

•
$$C = C^{(1)} \vee C^{(2)} \vee C^{(3)}$$

 $A^{(1,2,3)}$, $B^{(1,2,3)}$, $C^{(1,2,3)}$ can be computed "efficiently" from A, B, C

$$B^{(1)} \bigcup_{C^{(1)}}^{A^{(1)}} B^{(2)} \bigcup_{C^{(2)}}^{A^{(2)}} B^{(3)} \bigcup_{C^{(3)}}^{A^{(3)}}$$

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Parallel Boolean matrix multiplication by regular decomposition (contd.) $A \wedge B = \overline{C}$

Partition *ijk*-cube into a regular grid of $p^3 = p \cdot p \cdot p$ cubic $\frac{n}{p}$ -blocks A, B, C each gets partitioned into p^2 square $\frac{n}{p}$ -blocks A_{IJ} , B_{JK} , C_{IK} $0 \le I, J, K \le p$

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Parallel Boolean matrix multiplication by regular decomposition (contd.) Consider *J*-layers of cubic blocks for a fixed *J* and all *I*, *K*

Every processor

- assigned a *J*-layer for fixed *J*
- reads A_{IJ}, B_{JK}
- computes $A_{IJ} \wedge B_{JK} = C^J_{IK}$ by fast Boolean multiplication for all I, K
- computes regular decomposition $A_{IJ}^{(1,2,3)} \wedge B_{JK}^{(1,2,3)} = C_{IK}^{J(1,2,3)}$ where $A_{IJ}^{(1)}$, $B_{JK}^{(2)}$, $\overline{C_{IK}^{J(3)}}$ sparse, for all *I*, *K*

 $0 \leq I, J, K < p$

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Parallel Boolean matrix multiplication by regular decomposition (contd.) Consider also *I*-layers for a fixed *I* and *K*-layers for a fixed *K* Recompute every block product $A_{IJ} \wedge B_{JK} = C_{IK}^{J}$ by computing

•
$$A_{IJ}^{(1)} \wedge B_{JK}^{(1)} = C_{IK}^{J(1)}$$
 in K-layers
• $A_{IJ}^{(2)} \wedge B_{JK}^{(2)} = C_{IK}^{J(2)}$ in I-layers
• $A_{IJ}^{(3)} \wedge B_{JK}^{(3)} = C_{IK}^{J(3)}$ in J-layers

Every layer depends on $\leq \frac{n^2}{p}$ nonzeros of A, B, contributes $\leq \frac{n^2}{p}$ nonzeros to \overline{C} due to sparsity

Communication saved by only sending the indices of nonzeros

$$comp = O\left(\frac{n^{\omega}}{p}\right)$$
 $comm = O\left(\frac{n^2}{p}\right)$ $sync = O(1)$ $n >>>> p :-/$

Triangular system solution

Let *L* be an *n*-matrix, *b*, *c* be *n*-vectors *L* is lower triangular: $L_{ij} = \begin{cases} 0 & 0 \le i < j < n \\ arbitrary & otherwise \end{cases}$

Triangular system solution

Let *L* be an *n*-matrix, *b*, *c* be *n*-vectors *L* is lower triangular: $L_{ij} = \begin{cases} 0 & 0 \le i < j < n \\ \text{arbitrary otherwise} \end{cases}$ L · *b* = *c* The triangular system problem: given *L*, *c*, find *b*

Triangular system solution

Forward substitution

 $L \cdot b = c$ $L_{00} \cdot b_{0} = c_{0}$ $L_{10} \cdot b_{0} + L_{11} \cdot b_{1} = c_{1}$ $L_{20} \cdot b_{0} + L_{21} \cdot b_{1} + L_{22} \cdot b_{2} = c_{2}$... $\sum_{j:j \le i} L_{ij} \cdot b_{j} = c_{i}$... $\sum_{j:j \le n-1} L_{n-1,j} \cdot b_{j} = c_{n-1}$

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Triangular system solution

Forward substitution

 $L \cdot b = c$ $L_{00} \cdot b_0 = c_0$ $L_{10} \cdot b_0 + L_{11} \cdot b_1 = c_1$ $L_{20} \cdot b_0 + L_{21} \cdot b_1 + L_{22} \cdot b_2 = c_2$... $\sum_{j:j \le i} L_{ij} \cdot b_j = c_i$... $\sum_{i:j \le n-1} L_{n-1,j} \cdot b_j = c_{n-1}$

$$b_{0} \leftarrow L_{00}^{-1} \cdot c_{0}$$

$$b_{1} \leftarrow L_{11}^{-1} \cdot (c_{1} - L_{10} \cdot b_{0})$$

$$b_{2} \leftarrow L_{22}^{-1} \cdot (c_{2} - L_{20} \cdot b_{0} - L_{21} \cdot b_{1})$$
...
$$b_{i} \leftarrow L_{ii}^{-1} \cdot (c_{i} - \sum_{j:j < i} L_{ij} \cdot b_{j})$$
...
$$b_{n-1} \leftarrow L_{n-1,n-1}^{-1} \cdot (c_{n-1} - \sum_{j:j < n-1} L_{n-1,j} \cdot b_{j})$$

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Triangular system solution

Forward substitution

 $L \cdot b = c$ $L_{00} \cdot b_0 = c_0$ $L_{10} \cdot b_0 + L_{11} \cdot b_1 = c_1$ $L_{20} \cdot b_0 + L_{21} \cdot b_1 + L_{22} \cdot b_2 = c_2$. . . $\sum_{i:i < i} L_{ij} \cdot b_j = c_i$. . . $\sum_{i:i < n-1} L_{n-1,i} \cdot b_i = c_{n-1}$ Sequential work $O(n^2)$

$$b_{0} \leftarrow L_{00}^{-1} \cdot c_{0}$$

$$b_{1} \leftarrow L_{11}^{-1} \cdot (c_{1} - L_{10} \cdot b_{0})$$

$$b_{2} \leftarrow L_{22}^{-1} \cdot (c_{2} - L_{20} \cdot b_{0} - L_{21} \cdot b_{1})$$
...
$$b_{i} \leftarrow L_{ii}^{-1} \cdot (c_{i} - \sum_{j:j < i} L_{ij} \cdot b_{j})$$
...
$$b_{n-1} \leftarrow L_{n-1,n-1}^{-1} \cdot (c_{n-1} - \sum_{j:j < n-1} L_{n-1,j} \cdot b_{j})$$

Triangular system solution

Forward substitution

$L \cdot b = c$	
$L_{00}\cdot b_0=c_0$	$b_0 \leftarrow L_{00}^{-1} \cdot c_0$
$L_{10} \cdot b_0 + L_{11} \cdot b_1 = c_1$	$b_1 \gets L_{11}^{-1} \cdot (c_1 - L_{10} \cdot b_0)$
$L_{20} \cdot b_0 + L_{21} \cdot b_1 + L_{22} \cdot b_2 = c_2$	$b_2 \leftarrow L_{22}^{-1} \cdot (c_2 - L_{20} \cdot b_0 - L_{21} \cdot b_1)$
$\sum_{j:j\leq i} L_{ij} \cdot b_j = c_i$	$b_i \leftarrow L_{ii}^{-1} \cdot (c_i - \sum_{j:j < i} L_{ij} \cdot b_j)$
$\sum_{j:j\leq n-1}L_{n-1,j}\cdot b_j=c_{n-1}$	$b_{n-1} \leftarrow L_{n-1,n-1}^{-1} \cdot (c_{n-1} - \sum_{j:j < n-1} L_{n-1,j} \cdot b_j)$
Sequential work $O(n^2)$	

Symmetrically, an upper triangular system solved by back substitution

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Triangular system solution

Parallel forward substitution by 2D grid

Assume L is predistributed as needed, does not count as input



Triangular system solution

Block-recursive forward substitution

$$\begin{bmatrix} L \cdot b = c \\ \\ L_{00} \\ \\ L_{10} \\ \\ L_{11} \end{bmatrix} \cdot \begin{bmatrix} b_0 \\ \\ b_1 \end{bmatrix} =$$

Recursion: two half-sized subproblems

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 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

 $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} - L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion



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Triangular system solution

Block-recursive forward substitution

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Triangular system solution

Block-recursive forward substitution

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Triangular system solution

Block-recursive forward substitution

$$L \cdot b = c$$

$$[L_{00}] [b_0]$$

$$\begin{bmatrix} L_{\underline{00}} \\ L_{\underline{10}} \\ L_{\underline{11}} \end{bmatrix} \cdot \begin{bmatrix} D_{\underline{0}} \\ b_{\underline{1}} \end{bmatrix} = \begin{bmatrix} C_{\underline{0}} \\ c_{\underline{1}} \end{bmatrix}$$

Recursion: two half-sized subproblems

 $\begin{bmatrix} \alpha \end{bmatrix}$

 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

 $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} - L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion



Triangular system solution

Block-recursive forward substitution

Recursion: two half-sized subproblems

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Triangular system solution

Block-recursive forward substitution

Recursion: two half-sized subproblems

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Triangular system solution

Block-recursive forward substitution

Recursion: two half-sized subproblems

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Triangular system solution

Block-recursive forward substitution

Recursion: two half-sized subproblems

 $\begin{bmatrix} c_{\underline{0}} \\ c_1 \end{bmatrix}$

 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

 $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} - L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion



3 K 4 3 K

Triangular system solution

Block-recursive forward substitution

$$L \cdot b = c \\ \begin{bmatrix} L_{\underline{00}} \\ L_{\underline{10}} \\ L_{\underline{11}} \end{bmatrix} \cdot \begin{bmatrix} b_{\underline{0}} \\ b_{\underline{1}} \end{bmatrix} = \begin{bmatrix} c_{\underline{0}} \\ c_{\underline{1}} \end{bmatrix}$$

Recursion: two half-sized subproblems

 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

 $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} - L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion



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Triangular system solution

Block-recursive forward substitution

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Recursion: two half-sized subproblems

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Triangular system solution

Block-recursive forward substitution

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 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

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Block-recursive forward substitution

Recursion: two half-sized subproblems

 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

 $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} - L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion



Triangular system solution

Block-recursive forward substitution

Recursion: two half-sized subproblems

 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

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.

Triangular system solution

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 $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion

 $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} - L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion



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Triangular system solution

Block-recursive forward substitution

Recursion: two half-sized subproblems

- $L_{\underline{00}} \cdot b_{\underline{0}} = c_{\underline{0}}$ by recursion
- $L_{\underline{11}} \cdot b_{\underline{1}} = c_{\underline{1}} L_{\underline{10}} \cdot b_{\underline{1}}$ by recursion

Sequential work $O(n^2)$



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Triangular system solution

Parallel block-recursive forward substitution

Assume L is predistributed as needed, does not count as input

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Parallel block-recursive forward substitution

Assume L is predistributed as needed, does not count as input

At each level, the two recursive subproblems are dependent, hence recursion tree must be computed depth-first

At recursion level k:

• sequence of 2^k triangular system subproblems, each on $n/2^k$ -blocks

In particular, at level log p:

- sequence of p triangular system subproblems, each on n/p-blocks
- total $p \cdot O((n/p)^2) = O(n^2/p)$ sequential work, therefore each subproblem can be solved sequentially on an arbitrary processor

Triangular system solution

Parallel block-recursive forward substitution (contd.)

Recursion levels 0 to log *p*: block forward substitution using parallel matrix-vector multiplication

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Parallel block-recursive forward substitution (contd.)

Recursion levels 0 to log *p*: block forward substitution using parallel matrix-vector multiplication

Recursion level log p: a designated processor reads the current task's input, performs the task sequentially, and writes back the task's output

Parallel block-recursive forward substitution (contd.)

Recursion levels 0 to log *p*: block forward substitution using parallel matrix-vector multiplication

Recursion level log p: a designated processor reads the current task's input, performs the task sequentially, and writes back the task's output

$$comp = O(n^{2}/p) \cdot (1 + 2 \cdot (\frac{1}{2})^{2} + 2^{2} \cdot (\frac{1}{2^{2}})^{2} + ...) + O((n/p)^{2}) \cdot p = O(n^{2}/p) + O(n^{2}/p) = O(n^{2}/p)$$

$$comm = O(n/p^{1/2}) \cdot (1 + 2 \cdot \frac{1}{2} + 2^{2} \cdot \frac{1}{2^{2}} + ...) + O(n/p) \cdot p = O(n/p^{1/2}) \cdot \log p + O(n) = O(n)$$

Parallel block-recursive forward substitution (contd.)

Recursion levels 0 to log *p*: block forward substitution using parallel matrix-vector multiplication

Recursion level log p: a designated processor reads the current task's input, performs the task sequentially, and writes back the task's output

$$comp = O(n^{2}/p) \cdot (1 + 2 \cdot (\frac{1}{2})^{2} + 2^{2} \cdot (\frac{1}{2^{2}})^{2} + ...) + O((n/p)^{2}) \cdot p = O(n^{2}/p) + O(n^{2}/p) = O(n^{2}/p)$$

$$comm = O(n/p^{1/2}) \cdot (1 + 2 \cdot \frac{1}{2} + 2^{2} \cdot \frac{1}{2^{2}} + ...) + O(n/p) \cdot p = O(n/p^{1/2}) \cdot \log p + O(n) = O(n)$$

$$comp = O(n^{2}/p) \quad comm = O(n) \quad sync = O(p)$$

Generic Gaussian elimination

Let A, L, U be n-matrices

LU decomposition of A: $A = L \cdot U$



Generic Gaussian elimination

Application: solving a linear system

Ax = b

If LU decomposition of A is known: Ax = LUx = b

Solve triangular systems Ly = b then Ux = y, obtaining x

LU decomposition of A can be reused for multiple right-hand sides b

Generic Gaussian elimination

Block generic Gaussian elimination

LU decomposition: $A = L \cdot U$, also returns L^{-1} , U^{-1}

$$\begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix} = \begin{bmatrix} L_{00} & \\ L_{10} & L_{11} \end{bmatrix} \begin{bmatrix} U_{00} & U_{01} \\ & U_{11} \end{bmatrix}$$

Image: A matrix

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Parallel matrix algorithms Generic Gaussian elimination

Block generic Gaussian elimination LU decomposition: $A = L \cdot U$, also returns L^{-1} , U^{-1} $\begin{vmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{vmatrix} = \begin{vmatrix} L_{00} \\ I_{10} & I_{11} \end{vmatrix} \begin{vmatrix} U_{00} & U_{01} \\ U_{11} \end{vmatrix}$ Compute $A_{00} = L_{00} \cdot U_{00}$, also L_{00}^{-1} , U_{00}^{-1} $L_{10} \leftarrow A_{10} \cdot U_{00}^{-1} \quad U_{01} \leftarrow L_{00}^{-1} \cdot A_{01}$ $\bar{A}_{11} = A_{11} - L_{10} \cdot U_{01} = A_{11} - A_{10}A_{00}^{-1}A_{01}$ (Schur complement of A_{11}) $\begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix} = \begin{bmatrix} L_{00} \\ L_{10} & \bar{A}_{11} \end{bmatrix} \begin{bmatrix} U_{00} & U_{01} \\ & I \end{bmatrix}$ Compute $\bar{A}_{11} = L_{11} \cdot U_{11}$, also L_{11}^{-1} , U_{11}^{-1} , then return L^{-1} , U^{-1} : $L^{-1} \leftarrow \begin{bmatrix} L_{00}^{-1} & & \\ -L_{00}^{-1} & L_{01}^{-1} & & \\ & & U^{-1} \end{bmatrix} \quad U^{-1} \leftarrow \begin{bmatrix} U_{00}^{-1} & -U_{00}^{-1} & U_{10}^{-1} \\ & & U_{01}^{-1} \end{bmatrix}$

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Block generic Gaussian elimination (contd.)

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A_{00}, \ldots: either ordinary elements or blocks, can be applied recursively
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Recursion base: 1×1 matrix $A = 1 \cdot A$

Assumption: pivot elements nonzero (respectively pivot blocks nonsingular):

- $A_{00} \neq 0$ (respectively det $A_{00} \neq 0$)
- $ar{A}_{11}
 eq 0$ (respectively det $ar{A}_{11}
 eq 0$)

Hence no pivoting required

In practice, pivots must be sufficiently large. Holds for some special classes of matrices: diagonally dominant; symmetric positive definite.

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Generic Gaussian elimination

Iterative generic Gaussian elimination

Let A be an $n \times n$ matrix

$$A = {(1) \ (n-1)} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$$

A = LU by block generic Gaussian elimination on A, then on \bar{A}_{11} Sequential work $O(n^3)$

Generic Gaussian elimination

Recursive generic Gaussian elimination

Let A be an $n \times n$ matrix

$$A = {(n/2) (n/2) \atop (n/2)} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$$

A = LU by block generic Gaussian elimination on A, treating

- each '+' ('-', '.') as block '+' ('-', '.')
- each LU decomposition as recursive call on blocks

Generic Gaussian elimination

Recursive generic Gaussian elimination

Let A be an $n \times n$ matrix

$$A = {(n/2) (n/2) \atop (n/2)} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$$

A = LU by block generic Gaussian elimination on A, treating

- each '+' ('-', '·') as block '+' ('-', '·')
- each LU decomposition as recursive call on blocks

Sequential work:

- $O(n^3)$ using standard matrix multiplication
- $O(n^{\omega})$ using fast (Strassen-like) matrix multiplication

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At each level, the two recursive subproblems are dependent, hence recursion tree must be computed depth-first

At recursion level k:

• sequence of 2^k LU decomposition subproblems, each on $\frac{n}{2^k}$ -blocks

In particular, at level $\frac{1}{2} \cdot \log p$:

- sequence of $p^{1/2}$ LU decomposition subproblems, each on $\frac{n}{p^{1/2}}$ -blocks
- total $p^{1/2} \cdot O((\frac{n}{p^{1/2}})^3) = O(\frac{n^3}{p})$ sequential work, therefore each subproblem can be solved sequentially on an arbitrary processor

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Level $\frac{1}{2} \cdot \log p$: threshold to switch from parallel to sequential computation Recursion levels 0 to $\frac{1}{2} \cdot \log p$:

• block generic LU decomposition using parallel matrix multiplication

Level $\frac{1}{2} \cdot \log p$: threshold to switch from parallel to sequential computation Recursion levels 0 to $\frac{1}{2} \cdot \log p$:

• block generic LU decomposition using parallel matrix multiplication

Threshold recursion level $\frac{1}{2} \cdot \log p$:

• a designated processor reads the subproblem's input block, solves it sequentially, and writes the output blocks

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Level $\frac{1}{2} \cdot \log p$: threshold to switch from parallel to sequential computation Recursion levels 0 to $\frac{1}{2} \cdot \log p$:

• block generic LU decomposition using parallel matrix multiplication

Threshold recursion level $\frac{1}{2} \cdot \log p$:

• a designated processor reads the subproblem's input block, solves it sequentially, and writes the output blocks

$$comp = O(n^{3}/p)$$
 $comm = O(n^{2}/p^{1/2})$ $sync = O(p^{1/2})$

Parallel recursive generic Gaussian elimination (contd.) More generally: threshold level $\alpha \log p$, $1/2 \le \alpha \le 2/3$ Recursion levels 0 to $\alpha \log p$:

• block generic LU decomposition using parallel matrix multiplication

Parallel recursive generic Gaussian elimination (contd.) More generally: threshold level $\alpha \log p$, $1/2 \le \alpha \le 2/3$ Recursion levels 0 to $\alpha \log p$:

• block generic LU decomposition using parallel matrix multiplication

Threshold recursion level $\alpha \log p$:

• a designated processor reads the subproblem's input block, solves it sequentially, and writes the output blocks

Parallel recursive generic Gaussian elimination (contd.) More generally: threshold level $\alpha \log p$, $1/2 \le \alpha \le 2/3$ Recursion levels 0 to $\alpha \log p$:

• block generic LU decomposition using parallel matrix multiplication

Threshold recursion level $\alpha \log p$:

• a designated processor reads the subproblem's input block, solves it sequentially, and writes the output blocks

$$comp = O(n^3/p)$$
 $comm = O(n^2/p^{\alpha})$ $sync = O(p^{\alpha})$

Continuous tradeoff between *comm* and *sync*

Controlled by parameter $\alpha,\,1/2\leq\alpha\leq2/3$

 $\alpha=1/2:~\textit{comm}$ and sync as for 3D grid

$$comp = O(n^3/p)$$
 co

$$comm = O(n^2/p^{1/2})$$

$$sync = O(p^{1/2})$$

Continuous tradeoff between *comm* and *sync*

Controlled by parameter $\alpha,\,1/2\leq\alpha\leq2/3$

 $\alpha = 1/2:~\textit{comm}$ and sync as for 3D grid

$$comm = O(n^2/p^{1/2})$$

$$sync = O(p^{1/2})$$

 $\alpha = 2/3$:

- comm goes down to that of matrix multiplication
- sync goes up accordingly

$$comp = O(n^3/p)$$

 $comp = O(n^3/p)$

$$comm = O(n^2/p^{2/3})$$
 syn

$$sync = O(p^{2/3})$$

Pivoting permutes rows/columns of input matrix to remove the assumptions of generic Gaussian elimination, ensuring that:

- pivot elements are always nonzero
- pivot blocks are always nonsingular

Gaussian elimination with pivoting

Let A, P, L, U be n-matrices

PLU decomposition of A: $P \cdot A = L \cdot U$



P is a permutation matrix:

- all elements 0 or 1
- exactly one 1 in every row and column
- L is unit lower triangular, U is upper triangular

The PLU decomposition problem: given A, find P, L, U

Block Gaussian elimination with column pivoting Generalise PLU decomposition to "tall" rectangular matrices Let A be an $m \times n$ matrix, m > n

$$A = \begin{pmatrix} n \\ (m-n) \end{pmatrix} \begin{bmatrix} A_{00} \\ A_{10} \end{bmatrix} P \cdot \begin{bmatrix} A_{00} \\ A_{10} \end{bmatrix} = \begin{bmatrix} L_{00} \\ L_{10} \end{bmatrix} \cdot \begin{bmatrix} U_{00} \\ \cdot \end{bmatrix}$$

P is an $m \times m$ permutation matrix

 L_{00} is $n \times n$ unit lower triangular, U_{00} is $n \times n$ upper triangular



Gaussian elimination with pivoting

Block Gaussian elimination with column pivoting (contd.)

$$\begin{bmatrix} P_{00} & P_{01} \\ P_{10} & P_{11} \end{bmatrix} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix} = \begin{bmatrix} L_{00} \\ L_{10} & L_{11} \end{bmatrix} \begin{bmatrix} U_{00} & U_{01} \\ U_{11} \end{bmatrix}$$

$$Compute \begin{bmatrix} P_{00} & P_{01} \\ P'_{10} & P'_{11} \end{bmatrix} \begin{bmatrix} A_{00} \\ A_{10} \end{bmatrix} = \begin{bmatrix} L_{00} \\ L'_{10} \end{bmatrix} \begin{bmatrix} U_{00} \\ \cdot \end{bmatrix}$$

$$U_{01} \leftarrow L_{00}^{-1} (P_{00}A_{01} + P_{01}A_{11})$$

$$\bar{A}'_{11} \leftarrow P'_{10}A_{01} + P'_{11}A_{11} - L'_{10}U_{01}$$

$$\begin{bmatrix} P_{00} & P_{01} \\ P'_{10} & P'_{11} \end{bmatrix} \begin{bmatrix} A_{00} & A_{01} \\ A_{01} & A_{11} \end{bmatrix} = \begin{bmatrix} L_{00} \\ L'_{10} & \bar{A}'_{11} \end{bmatrix} \begin{bmatrix} U_{00} & U_{01} \\ \cdot & I \end{bmatrix}$$

$$Compute P''_{11}\bar{A}'_{11} = L_{11}U_{11}$$

$$\begin{bmatrix} P_{00} & P_{01} \\ P''_{11}P'_{10} & P''_{11}P'_{11} \end{bmatrix} \begin{bmatrix} A_{00} & A_{01} \\ A_{01} & A_{11} \end{bmatrix} = \begin{bmatrix} L_{00} \\ P''_{11}L'_{10} & L_{11} \end{bmatrix} \begin{bmatrix} U_{00} & U_{01} \\ \cdot & U_{11} \end{bmatrix}$$

Block Gaussian elimination with column pivoting (contd.)

 A_{00}, \ldots : either ordinary elements or blocks, can be applied recursively Recursion base: $m \times 1$ matrix

$$A = {(1) \atop (m-1)} \begin{bmatrix} A_0 \\ A_1 \end{bmatrix} P \begin{bmatrix} A_0 \\ A_1 \end{bmatrix} = \begin{bmatrix} A'_0 \\ A'_1 \end{bmatrix} = \begin{bmatrix} 1 \\ L_1 \end{bmatrix} \begin{bmatrix} A'_0 \\ \cdot \end{bmatrix}$$

P is a permutation such that $|A'_0|$ is largest across A

Iterative Gaussian elimination with column pivoting

Let A be an $n \times n$ matrix

$$A = {(1) \\ (n-1)} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$$

 $\mathit{P\!A} = \mathit{LU}$ by block Gaussian elimination with column pivoting on $\mathit{A},$ then on $\bar{\mathit{A}}_{11}'$

Sequential work $O(n^3)$



Recursive Gaussian elimination with column pivoting

Let A be an $n \times n$ matrix

$$A = \frac{(n/2)}{(n/2)} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$$

PA = LU by block Gaussian elimination with column pivoting on A, treating

- each '+' ('-', '.') as block '+' ('-', '.')
- each PLU decomposition as recursive call on blocks

Recursive Gaussian elimination with column pivoting

Let A be an $n \times n$ matrix

$$A = \frac{(n/2)}{(n/2)} \begin{bmatrix} A_{00} & A_{01} \\ A_{10} & A_{11} \end{bmatrix}$$

PA = LU by block Gaussian elimination with column pivoting on A, treating

- each '+' ('-', '.') as block '+' ('-', '.')
- each PLU decomposition as recursive call on blocks

Sequential work:

- $O(n^3)$ using standard matrix multiplication
- $O(n^{\omega})$ using fast (Strassen-like) matrix multiplication

Parallel recursive Gaussian elimination with column pivoting

At each level, the two recursive subproblems are dependent, hence recursion tree must be computed depth-first

At recursion level k:

• sequence of 2^k PLU decomposition subproblems, each on $\frac{n}{2^k} \times n$ blocks

In particular, at level log p:

- sequence of p PLU decomposition subproblems, each on $\frac{n}{p} \times n$ blocks
- total $p \cdot O(\frac{n^3}{p^2}) = O(\frac{n^3}{p})$ sequential work, therefore each subproblem can be solved sequentially on an arbitrary processor

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Parallel recursive Gaussian elimination with column pivoting (contd.) Level log p: threshold to switch from parallel to sequential computation Recursion levels 0 to log p:

• block PLU decomposition using parallel matrix multiplication

Threshold recursion level log p:

• a designated processor reads the subproblem's input block, solves it sequentially, and writes the output blocks

$$comp = O(n^3/p)$$
 $comm = O(n^2)$ $sync = O(p)$

Parallel recursive Gaussian elimination with column pivoting (contd.) Alternative: no switching to sequential computation Level log p: threshold to switch to fine-grained parallel computation Recursion levels 0 to log p:

- block PLU decomposition using parallel matrix multiplication Recursion levels log p to log n:
 - block PLU decomposition on partitioned matrix, using broadcast of pivot subrows and *p* instances of sequential matrix multiplication

Recursion base at level log n:

• column PLU decomposition; pivot selected by balanced binary tree

$$comp = O(n^3/p)$$

$$comm = O(n^2/p^{2/3})$$

Parallel recursive Gaussian elimination with column pivoting (contd.)

Discontinuous tradeoff between *comm* and *sync*

Coarse-grained algorithm: *comm* and *sync* as for 2D grid with work and data size O(n) per node

$$comp = O(n^3/p)$$
 $comm = O(n^2)$ $sync = O(p)$

Fine-grained algorithm: *comm* as for matrix multiplication; *sync* becomes a function of n

$$comp = O(n^3/p)$$
 $comm = O(n^2/p^{2/3})$ $sync = O(n)$

Computation by circuits

- 2 Parallel computation models
- 3 Basic parallel algorithms
- 4 Further parallel algorithms
- 5 Parallel matrix algorithms
- 6 Parallel graph algorithms

Semiring: a set S with addition \oplus and multiplication \odot \oplus commutative, associative, has identity \square $a \oplus b = b \oplus a$ $a \oplus (b \oplus c) = (a \oplus b) \oplus c$ $a \oplus \square = \square \oplus a = a$ \odot associative, has annihilator \square and identity \square $a \odot (b \odot c) = (a \odot b) \odot c$ $a \odot \Box = \Box \odot a = \Box$ $a \odot \Box = \Box \odot a = a$ \odot distributes over \odot $a \odot (b \oplus c) = a \odot b \oplus a \odot c$ $(a \oplus b) \odot c = a \odot c \oplus b \odot c$ In general, no subtraction or division!

We will occasionally write ab for $a \odot b$, a^2 for $a \odot a$, etc.

Parallel graph algorithms

Algebraic path problem

Some specific semirings:

	5	\oplus	0	\odot	1			
real	\mathbb{R}	+	0	•	1			
Boolean	$\{0, 1\}$	V	0	\wedge	1			
tropical	\mathbb{R}^+	min	$+\infty$	+	0			
$\mathbb{R}^+ = \mathbb{R}_{>0} \cup \{+\infty\}$								

Alexander Tiskin (Warwick)

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Some specific semirings:

	S	\oplus	0	\odot	1			
real	\mathbb{R}	+	0	•	1			
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tropical	\mathbb{R}^+	min	$+\infty$	+	0			
$\mathbb{R}^+ = \mathbb{R}_{>0} \cup \{+\infty\}$								

Given a semiring S, square matrices of size n over S also form a semiring:

- \oplus given by matrix addition; \square by the zero matrix
- \odot given by matrix multiplication; \blacksquare by the identity matrix

Parallel graph algorithms

Algebraic path problem

The closure of *a*: $a^* = \square \oplus a \oplus a^2 \oplus a^3 \oplus \cdots$

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Parallel graph algorithms Algebraic path problem

The closure of a: $a^* = \square \oplus a \oplus a^2 \oplus a^3 \oplus \cdots$ Examples

• real:
$$a^* = 1 + a + a^2 + a^3 + \dots = \begin{cases} \frac{1}{1-a} & \text{if } |a| < 1\\ \text{undefined} & \text{otherwise} \end{cases}$$

• Boolean: $a^* = 1 \lor a \lor a \lor a \lor \dots = 1$
• tropical: $a^* = \min(0, a, 2a, 3a, \dots) = 0$

In matrix semirings, closures are more interesting

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- infinite $a_1 \oplus a_2 \oplus a_3 \oplus \cdots$ (e.g. a closure) always defined
- infinite \oplus commutative, associative
- ullet \odot distributive over infinite \oplus

In a closed semiring, every element and every square matrix have a closure

- infinite $a_1 \oplus a_2 \oplus a_3 \oplus \cdots$ (e.g. a closure) always defined
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In a closed semiring, every element and every square matrix have a closure Examples

• real semiring not closed: infinite + can be divergent

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In a closed semiring, every element and every square matrix have a closure Examples

- real semiring not closed: infinite + can be divergent
- Boolean semiring closed: infinite \lor is \exists

- infinite $a_1 \oplus a_2 \oplus a_3 \oplus \cdots$ (e.g. a closure) always defined
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- $\bullet \ \odot \ {\sf distributive \ over \ infinite \ } \oplus$

In a closed semiring, every element and every square matrix have a closure Examples

- real semiring not closed: infinite + can be divergent
- Boolean semiring closed: infinite \lor is \exists
- tropical semiring closed: infinite min is inf (greatest lower bound)

- infinite $a_1 \oplus a_2 \oplus a_3 \oplus \cdots$ (e.g. a closure) always defined
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In a closed semiring, every element and every square matrix have a closure Examples

- real semiring not closed: infinite + can be divergent
- Boolean semiring closed: infinite \lor is \exists
- tropical semiring closed: infinite min is inf (greatest lower bound)

Matrix closure problem, aka algebraic path problem

Given A: $n \times n$ matrix over a semiring

Compute $A^* = I \oplus A \oplus A^2 \oplus A^3 \oplus \cdots$

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Matrix closure problem, aka algebraic path problem

Given A: $n \times n$ matrix over a semiring

Compute $A^* = I \oplus A \oplus A^2 \oplus A^3 \oplus \cdots$

• real: $A^* = I + A + A^2 + \cdots = (I - A)^{-1}$, if nonsingular

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Matrix closure problem, aka algebraic path problem

Given A: $n \times n$ matrix over a semiring

Compute $A^* = I \oplus A \oplus A^2 \oplus A^3 \oplus \cdots$

• real:
$$A^* = I + A + A^2 + \cdots = (I - A)^{-1}$$
, if nonsingular

Weighted digraph on n nodes: define matrix as

$$A_{ij} = \begin{cases} \square = 0 & \text{if } i = j \\ \text{length of edge } i \to j & \text{if edge exists} \\ \square = +\infty & \text{otherwise} \end{cases}$$

• Boolean: A* gives transitive closure

• tropical: A* gives all-pairs shortest paths

Parallel graph algorithms

Algebraic path problem

$$A = \begin{bmatrix} 0 & 5 & 10 & \infty & 10 \\ \infty & 0 & 3 & 2 & 9 \\ \infty & 2 & 0 & \infty & 1 \\ 7 & \infty & \infty & 0 & 6 \\ \infty & \infty & \infty & 4 & 0 \end{bmatrix}$$



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Parallel graph algorithms

Algebraic path problem

$$A = \begin{bmatrix} 0 & 5 & 10 & \infty & 10 \\ \infty & 0 & 3 & 2 & 9 \\ \infty & 2 & 0 & \infty & 1 \\ 7 & \infty & \infty & 0 & 6 \\ \infty & \infty & \infty & 4 & 0 \end{bmatrix}$$
$$A^* = \begin{bmatrix} 0 & 5 & 8 & 7 & 9 \\ 9 & 0 & 3 & 2 & 4 \\ 11 & 2 & 0 & 4 & 1 \\ 7 & 12 & 15 & 0 & 6 \\ 11 & 16 & 19 & 4 & 0 \end{bmatrix}$$



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Floyd–Warshall algorithm

A: $n \times n$ matrix over closed semiring

First step of elimination: pivot $A_{00} = \square$

 $A'_{\underline{11}} \leftarrow A_{\underline{11}} \oplus A_{\underline{10}} \odot A_{0\underline{1}}$

(E.g. replace A_{ij} with $A_{i0} + A_{0j}$, if it gives a shortcut)

Continue elimination on reduced matrix A'_{11}

Generic Gaussian elimination in disguise

Works for any closed semiring

Sequential work $O(n^3)$

[Floyd, Warshall: 1962]



Parallel graph algorithms

Algebraic path problem

Block Floyd–Warshall algorithm

$$A = \begin{bmatrix} A_{\underline{00}} & A_{\underline{01}} \\ A_{\underline{10}} & A_{\underline{11}} \end{bmatrix} \qquad A^* = \begin{bmatrix} A_{\underline{00}}^{\prime\prime} & A_{\underline{01}}^{\prime\prime} \\ A_{\underline{10}}^{\prime\prime} & A_{\underline{11}}^{\prime\prime} \end{bmatrix}$$

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Parallel graph algorithms Algebraic path problem

Block Floyd–Warshall algorithm

$$A = \begin{bmatrix} A_{\underline{00}} & A_{\underline{01}} \\ A_{\underline{10}} & A_{\underline{11}} \end{bmatrix} \qquad A^* = \begin{bmatrix} A_{\underline{00}}'' & A_{\underline{01}}'' \\ A_{\underline{10}}'' & A_{\underline{11}}'' \end{bmatrix}$$

Recursion: two half-sized subproblems

$$\begin{array}{l} \mathcal{A}_{\underline{00}}' \leftarrow \mathcal{A}_{\underline{00}}^{*} \text{ by recursion} \\ \mathcal{A}_{\underline{01}}' \leftarrow \mathcal{A}_{\underline{00}}' \mathcal{A}_{\underline{01}} & \mathcal{A}_{\underline{10}}' \leftarrow \mathcal{A}_{\underline{10}} \mathcal{A}_{\underline{00}}' & \mathcal{A}_{\underline{11}}' \leftarrow \mathcal{A}_{\underline{11}} \oplus \mathcal{A}_{\underline{10}} \mathcal{A}_{\underline{00}}' \mathcal{A}_{\underline{01}} \\ \mathcal{A}_{\underline{11}}'' \leftarrow (\mathcal{A}_{\underline{11}}')^{*} \text{ by recursion} \\ \mathcal{A}_{\underline{10}}'' \leftarrow \mathcal{A}_{\underline{11}}' \mathcal{A}_{\underline{10}}' & \mathcal{A}_{\underline{01}}' \leftarrow \mathcal{A}_{\underline{01}}' \mathcal{A}_{\underline{11}}'' & \mathcal{A}_{\underline{00}}' \leftarrow \mathcal{A}_{\underline{00}}' \oplus \mathcal{A}_{\underline{01}}' \mathcal{A}_{\underline{11}}'' \mathcal{A}_{\underline{10}}' \end{array}$$



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Parallel graph algorithms Algebraic path problem

Block Floyd–Warshall algorithm

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Recursion: two half-sized subproblems

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Parallel algebraic path computation

Similar to LU decomposition by block generic Gaussian elimination

Recursion tree is unfolded depth-first

Recursion levels 0 to $\alpha \log p$: block Floyd–Warshall using parallel matrix multiplication

Recursion level $\alpha \log p$: on each visit, a designated processor reads the current task's input, performs the task sequentially, and writes back the task's output

Parallel algebraic path computation

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Recursion levels 0 to $\alpha \log p$: block Floyd–Warshall using parallel matrix multiplication

Recursion level $\alpha \log p$: on each visit, a designated processor reads the current task's input, performs the task sequentially, and writes back the task's output

Threshold level controlled by parameter α : $1/2 \le \alpha \le 2/3$

$$comm = O(n^2/p^{lpha})$$
 $sync = O(p^{lpha})$

 $comp = O(n^3/p)$

Parallel graph algorithms Algebraic path problem

Parallel algebraic path computation (contd.)

In particular:

 $\alpha = 1/2$

$$comp = O(n^3/p) \quad comm = O(n^2/p^{1/2}) \quad sync = O(p^{1/2})$$

Cf. 2D grid

Image: Image:

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Parallel graph algorithms Algebraic path problem

Parallel algebraic path computation (contd.)

In particular:

 $\alpha = 1/2$ $\boxed{comp = O(n^3/p)} \quad \boxed{comm = O(n^2/p^{1/2})} \quad \boxed{sync = O(p^{1/2})}$ Cf. 2D grid $\alpha = 2/3$ $\boxed{comp = O(n^3/p)} \quad \boxed{comm = O(n^2/p^{2/3})} \quad \boxed{sync = O(p^{2/3})}$

Cf. matrix multiplication

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All-pairs shortest paths (APSP) problem: matrix closure (algebraic path) problem over tropical semiring

We continue to use the generic notation: \oplus for min, \odot for +

All-pairs shortest paths (APSP) problem: matrix closure (algebraic path) problem over tropical semiring

$$\begin{array}{c|c} S & \oplus & \boxdot & \odot & \fbox{I} \\ \hline tropical & \mathbb{R}_{\geq 0} \cup \{+\infty\} & \min & +\infty & + & 0 \end{array}$$

We continue to use the generic notation: \oplus for min, \odot for +

Can be solved by Floyd-Warshall algorithm (ordinary or block)

Also works with negative weights, but no negative cycles

To improve on Floyd–Warshall, we must exploit the tropical semiring's idempotence: $a \oplus a = \min(a, a) = a$

All-pairs shortest paths

A: $n \times n$ matrix over the tropical semiring, defining a weighted digraph Path length: sum (\odot -product) of all its edge lengths Path size: its total number of edges A: $n \times n$ matrix over the tropical semiring, defining a weighted digraph Path length: sum (\odot -product) of all its edge lengths Path size: its total number of edges $(A^k)_{ij} =$ length of shortest path $i \rightsquigarrow j$ among those of size $\leq k$ $(A^*)_{ij} =$ length of the shortest path $i \rightsquigarrow j$ of any size

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A: $n \times n$ matrix over the tropical semiring, defining a weighted digraph Path length: sum (\odot -product) of all its edge lengths Path size: its total number of edges $(A^k)_{ij} =$ length of shortest path $i \rightsquigarrow j$ among those of size $\leq k$ $(A^*)_{ij} =$ length of the shortest path $i \rightsquigarrow j$ of any size The APSP problem:

$$A^* = I \oplus A \oplus A^2 \oplus \cdots = I \oplus A \oplus A^2 \oplus \cdots \oplus A^n = (I \oplus A)^n = A^n$$

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APSP by multi-Dijkstra

Dijkstra's algorithm

[Dijkstra: 1959]

Computes single-source shortest paths from fixed source (say, node 0)

Ranks all nodes by distance from node 0: nearest, second nearest, etc.

Every time a node *i* has been ranked:

 $A_{0j} \leftarrow A_{0j} \oplus A_{0i} \odot A_{ij}$ for all j not yet ranked

Assign the next rank to the unranked node closest to node 0 and repeat

APSP by multi-Dijkstra

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It is essential that the edge lengths are nonnegative

Sequential work $O(n^2)$

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It is essential that the edge lengths are nonnegative

Sequential work $O(n^2)$

APSP: run Dijkstra's algorithm independently from every node as a source, sequential work $O(n^3)$

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All-pairs shortest paths

Parallel APSP by multi-Dijkstra

Every processor

- reads matrix A and is assigned a subset of n/p nodes
- runs *n*/*p* independent instances of Dijkstra's algorithm from its assigned nodes
- writes back the resulting n^2/p shortest distances

All-pairs shortest paths

Parallel APSP by multi-Dijkstra

Every processor

 $comp = O(n^3/\mu)$

- reads matrix A and is assigned a subset of n/p nodes
- runs *n*/*p* independent instances of Dijkstra's algorithm from its assigned nodes
- writes back the resulting n^2/p shortest distances

$$o) \quad comm = O(n^2) \quad sync = O(1)$$

Parallel graph algorithms

All-pairs shortest paths

Parallel APSP: summary so far

 $comp = O(n^3/p)$

Floyd–Warshall, $\alpha = 2/3$

Floyd–Warshall, $\alpha = 1/2$

Multi-Dijkstra

$$comm = O(n^2/p^{2/3})$$
 $sync = O(p^{2/3})$ $comm = O(n^2/p^{1/2})$ $sync = O(p^{1/2})$ $comm = O(n^2)$ $sync = O(1)$

Parallel graph algorithms

All-pairs shortest paths

Parallel APSP: summary so far

 $comp = O(n^3/p)$

Floyd–Warshall, $lpha=2/3$	$comm = O(n^2/p^{2/3})$	sync = $O(p^{2/3})$
Floyd–Warshall, $lpha=1/2$	$\boxed{\textit{comm} = O(n^2/p^{1/2})}$	$sync = O(p^{1/2})$
Multi-Dijkstra	$\boxed{comm = O(n^2)}$	sync = O(1)
Coming next	$comm = O(n^2/p^{2/3})$	$sync = O(\log p)$

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All-pairs shortest paths

Path doubling

Compute A,
$$A^2$$
, $A^4 = (A^2)^2$, $A^8 = (A^4)^2$, ..., $A^n = A^*$

Overall, log n rounds of matrix \odot -multiplication: looks promising...

... but not work-optimal: sequential time $O(n^3 \log n)$

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All-pairs shortest paths

Sparsified path doubling

[Alon+: 1997]

Idea: remove redundancy in path doubling by keeping track of path sizes

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Sparsified path doubling

[Alon+: 1997]

Idea: remove redundancy in path doubling by keeping track of path sizes Lex-tropical semiring (aka lexicographic semiring)

- elements are pairs (a,k) $a\in \mathbb{R}^+$ $k\in \mathbb{Z}^+$
- \oplus is lexicographic min $\square = (+\infty, +\infty)$
- \odot is numerical + $\mathbb{1} = (0,0)$

Weighted digraph on n nodes: define matrix as

$$egin{aligned} \mathsf{A}_{ij} = egin{cases} \mathbb{I} = (0,0) & ext{if } i = j \ (ext{length of edge } i o j,1) & ext{if edge exists} \ \mathbb{O} = (+\infty,+\infty) & ext{otherwise} \end{aligned}$$

Sparsified path doubling (contd.) $A_{ij}^{k} = \text{length of shortest path } i \rightsquigarrow j \text{ among those of size } \leq k$ Let $(a, k)|_{t} = \begin{cases} (a, k) & \text{if } k = t \\ \hline \Box & \text{otherwise} \end{cases}$ $A_{ij}^{k}|_{\ell} = \begin{cases} A_{ij}^{k} & \text{if realised by a path of size exactly } \ell \leq k \\ \hline \Box & \text{otherwise} \end{cases}$

 $A^k|_{\ell}$ contains all lengths of shortest paths of size exactly ℓ . May also contain some non-shortest path lengths (where the shortest path is of size $\geq k$), but that does no harm.

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Sparsified path doubling (contd.) We have $A^k = A^k|_0 \oplus \cdots \oplus A^k|_{\frac{k}{2}} \oplus \cdots \oplus A^k|_k$ Consider matrices in \oplus -sum $A^k|_{\frac{k}{2}} \oplus \cdots \oplus A^k|_k$ Total density of these $\frac{k}{2}$ matrices is ≤ 1 . This is $\leq \frac{2}{k}$ per matrix on average, and hence also for some specific $A^k|_{\frac{k}{2}+\ell}$, $0 \le \ell \le \frac{k}{2}$ We have $(I \oplus A^k|_{\frac{k}{2}+\ell}) \odot A^k = A^{\frac{3k}{2}+\ell}$ This is because a shortest path of size $\leq \frac{3k}{2} + \ell$ is either

- of size $\leq k$, or
- (shortest path of size exactly $\frac{k}{2} + \ell$) \odot (one of size $\leq k$)

Sparse-by-dense matrix \odot -product: $\leq \frac{2n^2}{k} \cdot n = \frac{2n^3}{k}$ elementary \odot -products

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Parallel graph algorithms All-pairs shortest paths

Sparsified path doubling (contd.) Compute matrices A, $A^{\frac{3}{2}+\ell}$, $A^{(\frac{3}{2})^2+\ell'}$, ..., $A^n = A^*$ Overall, $\leq \log_{3/2} n$ rounds of sparsified path doubling Sequential work $O(n^3) \cdot \left(1 + \left(\frac{3}{2}\right)^{-1} + \left(\frac{3}{2}\right)^{-2} + \cdots\right) = O(n^3)$

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Parallel APSP by sparsified path doubling

All processors collectively

- compute $B = A^{p+\ell}$ by $\leq \log_{3/2} p$ rounds of sparsified path doubling
- select $B|_p$ from B

 $B|_p$ is dense, but can be decomposed into a \odot -product of sparse matrices $B|_p = B|_q \odot B|_{p-q}$ $0 \le q \le \frac{p}{2}$

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Such a q is found sequentially by a designated processor

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Parallel APSP by sparsified path doubling (contd.) Every processor

- selects and writes its shares of $B|_q$, $B|_{p-q}$ from B
- reads whole $B|_q$, $B|_{p-q}$ and combines them to $B|_p = B|_q \odot B|_{p-q}$

All processors collectively

- compute $(B|_p)^*$ by parallel multi-Dijkstra
- compute $(B|_p)^* \odot B = A^*$ by parallel matrix \odot -multiplication

Use of multi-Dijkstra requires that all edge lengths in A are nonnegative

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Parallel APSP by sparsified path doubling (contd.) Every processor

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Use of multi-Dijkstra requires that all edge lengths in A are nonnegative

$$comp = O(n^3/p)$$

$$comm = O(n^2/p^{2/3})$$
 $sync = O(\log p^{2/3})$

All-pairs shortest paths

Parallel APSP by sparsified path doubling (contd.)

Now let A have arbitrary (nonnegative or negative) edge lengths. We still assume there are no negative-length cycles.

Parallel APSP by sparsified path doubling (contd.)

Now let A have arbitrary (nonnegative or negative) edge lengths. We still assume there are no negative-length cycles.

All processors collectively

• compute $B = A^{p^2 + \ell}$ by $\leq 2 \log_{3/2} p$ rounds of sparsified path doubling

Let $P = \{p, 2p, \dots, p^2\}$, $P - q = \{p - q, 2p - q, \dots, p^2 - q\}$ for any q $B|_P = B|_p \oplus B|_{2p} \oplus \dots \oplus B|_{p^2}$

All processors collectively

• select $B|_P$ from B
Parallel APSP by sparsified path doubling (contd.)

 $B|_P$ is dense, but can be decomposed into a \odot -product of sparse matrices

$$B|_P = B|_q \odot B|_{P-q} \quad 0 \le q \le \frac{p}{2}$$

Consider matrix pair $B|_q$, $B|_{P-q}$ for each q

Total density of these $\frac{p}{2}$ pairs is ≤ 1 . This is $\leq \frac{2}{p}$ per pair on average, and hence also for some specific pair with a fixed q

Such a q is found sequentially by a designated processor

Parallel APSP by sparsified path doubling (contd.)

Every processor

- selects and writes its shares of $B|_q$, $B|_{P-q}$ from B
- reads whole $B|_q$, $B|_{P-q}$ and combines them to $B|_P = B|_q \odot B|_{P-q}$
- computes (B|_P)* by ≤ log_{3/2} n rounds of sparsified path doubling (with path sizes multiples of p)

All processors collectively

• compute $(B|_P)^* \odot B = A^*$ by parallel matrix \odot -multiplication

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Parallel APSP by sparsified path doubling (contd.)

Every processor

- selects and writes its shares of $B|_q$, $B|_{P-q}$ from B
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All processors collectively

• compute $(B|_P)^* \odot B = A^*$ by parallel matrix \odot -multiplication

$$comm = O(n^2/p^{2/3}) \quad sync = O(\log p)$$

 $comp = O(n^3/p)$

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