

Introduction to Parallel Computation

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A Graph-Theoretical Problem

“All Pairs Shortest Paths”

- Given: graph $G = (V, A)$

– Directed acyclic graph, $|V| = n$

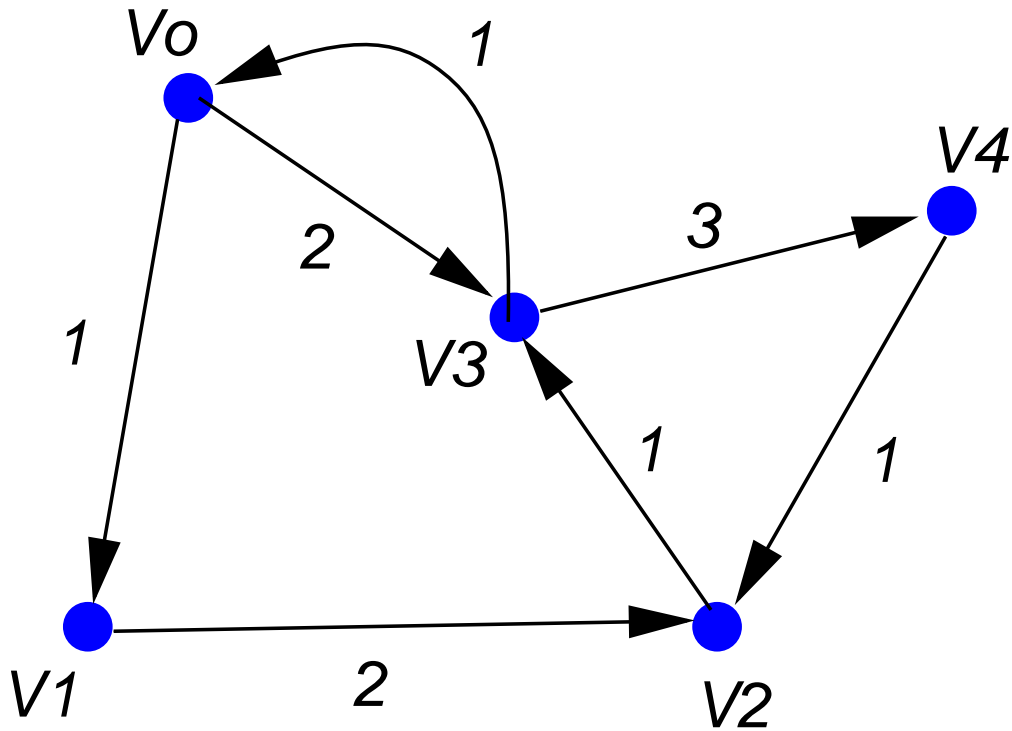
- Representation: weight matrix W

$$W(i, j) = \begin{cases} 0 & \text{if } i = j \\ w > 0 & \text{if edge of length } w \\ & \text{between nodes } i \text{ and } j \\ \infty & \text{if no edge between } i \text{ and } j \end{cases}$$

- Wanted: distance matrix D

$$D(i, j) = \begin{cases} 0 & \text{if } i = j \\ d > 0 & \text{if } i \neq j \text{ where } d \text{ length of} \\ & \text{shortest path between } i \text{ and } j \\ \infty & \text{if no edge between } i \text{ and } j \end{cases}$$

Example



W	v_0	v_1	v_2	v_3	v_4
v_0	0	1	∞	2	∞
v_1	∞	0	2	∞	∞
v_2	∞	∞	0	1	∞
v_3	1	∞	∞	0	3
v_4	∞	∞	1	∞	0

 \Rightarrow

D	v_0	v_1	v_2	v_3	v_4
v_0	0	1	3	2	5
v_1	4	0	2	3	6
v_2	2	3	0	1	4
v_3	1	2	4	0	3
v_4	3	4	1	2	0

Solution Idea

- Construct sequence of matrices
 - D_0, D_1, \dots, D_{n-1}
 - D_i describes all shortest paths with not more than i edges.
- Consequence: $D_{n-1} = D$
- Proof
 - Assume shortest path p with more than $n - 1$ edges. Then there is some node v twice in this path i.e. $p = \langle i, \dots, \underline{v}, \dots, \underline{v}, \dots, j \rangle$. But then path $p' = \langle i, \dots, v, \dots, j \rangle$ is shorter!

Construction

$$D_0(i, j) = \begin{cases} 0 & \text{if } i = j \\ \infty & \text{if } i \neq j \end{cases}$$

$$D_1(i, j) = W(i, j)$$

$$D_{r+1}(i, j) = ?$$

Two Cases

Let l be the number of edges of path captured by $D_r(i, j)$.

1. $|p| = l$

- $p = \langle i, \dots, j \rangle$
 l edges

$$D_{r+1}(i, j) = D_r(i, j)$$

2. $|p| = n + 1$

- $p = \langle i, \dots, k, j \rangle$
 l edges

$$D_{r+1}(i, j) = D_r(i, k) + W(k, j)$$

$$D_{r+1}(i, j) = \min\{D_r(i, j), \min_k\{D_r(i, k) + W(k, j)\}\}$$

Sequential Algorithm

AllPairsShortestPaths(W) :

```
D = W
for r=1 to n-1 do
  D = MatMin(D, W)
return D
```

MatMin(D, W) :

```
for i=1 to n do
  for j=1 to n do
    E[i,j] = D[i,j]
    for k=1 to n do
      E[i,j] = min(E[i,j], D[i,k]+W[k,j])
return E
```

Observation

MatMin has same structure as matrix multiplication ($+ \rightarrow \min, * \rightarrow +$).

- Define $D \times W = \text{MatMult}(D, W)$
- Begin: $D_1 = W$
- General: $D_i = D^{i-1} \times W = W^i$
- End: $D = D_{n-1} = W^{n-1}$

Problem solution is essentially repeated matrix multiplication!

Optimization

- Instead of computing

- $W^1, W^2, W^3, \dots, W^{n-1}$

- we can compute

- $W^1, W^2, W^4, \dots, W^{2^s}$
(where $2^s \geq n - 1$).

True for matrix multiplication as well as for MatMin (since both are associative).

AllPairsShortestPaths(W) :

```
D = W
for r=1 to s do
  D = MatMin(D, D)
return D
```

We can reduce n matrix multiplications to $\log n$ square computations!

Time Analysis

- n nodes.
- $\log n$ square computations.
- n^3 (min,+) operations for each square computation.

*Sequential time complexity $O(\log n * n^3)$*

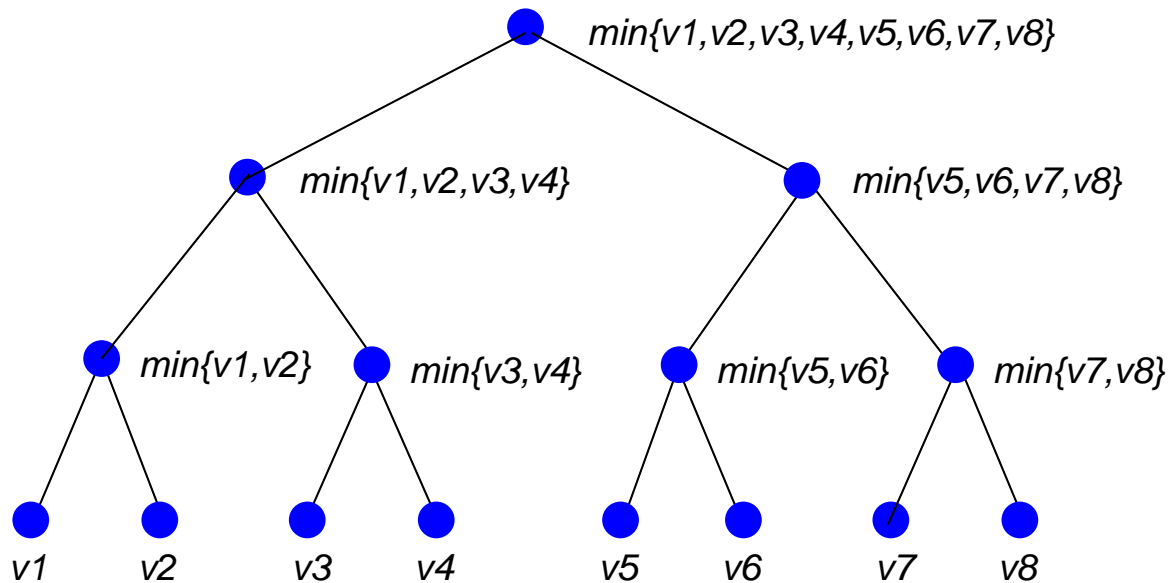
Parallel Algorithm

- Sequence of square operations.
- Each square op. contains n^3 independent (min,+) operations that can be performed in parallel yielding n^3 results.
- Each of the n^2 entries $D(i, j)$ is a minimum of n values.
- Time complexity:
 - $\log n$ square computations.
 - 1 time step for all (min,+) operations.
 - $\log n$ time to compute each of the n^2 minimums.

Parallel time complexity $O(\log^2 n)$

Minimum of n Values

Tree-like minimum construction



Depth of tree = computation time = $O(\log n)$

Comparison

- General

- Sequential: $O(\log n * n^3)$
- Parallel: $O(\log^2 n)$
- Processors: $O(n^3)$

- Time/processor product

- Sequential product: $O(\log n * n^3)$
- Parallel product: $O(\log^2 n * n^3)$

Parallel algorithm is in some sense less efficient than sequential one!

Parallel Machine Models

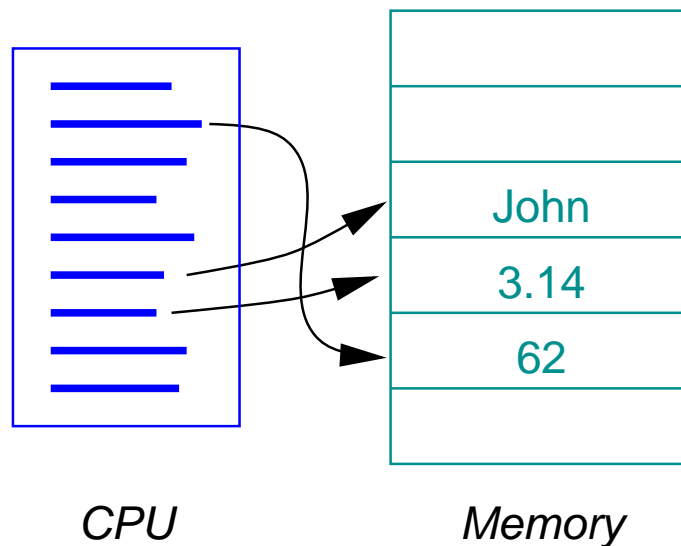
- How to define algorithm?
- How to implement algorithm?
- How to analyze algorithm?

We need a parallel machine and programming model!

Sequential Machine Model

Von Neumann Computer, Random Access Machine (RAM).

- Central Processing Unit (CPU)
 - Executes a stored program.
- Storage Unit (Memory)
 - Random access to any memory cell.

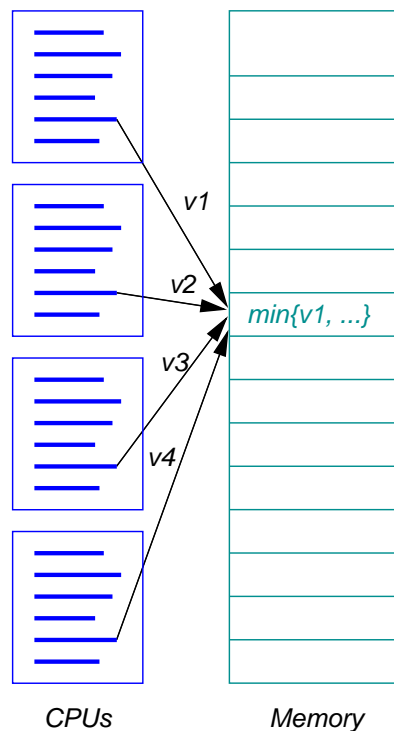


Program execution is sequence of read/write operations on memory.

PRAM Model

Parallel Random Access Machine.

- Set of CPUs.
- CPUs operate on same memory.
- CPUs execute same program lock-step.



Theoretical model for designing and analyzing parallel algorithms.

PRAM Variants

Restrictions of access to memory cells.

- EREW (exclusive read, exclusive write)
 - Only one access to individual memory cell at a time.
- CREW (concurrent read, exclusive write).
 - Multiple concurrent reads to a memory cell, but writes are exclusive.
- CRCW (concurrent read, concurrent write).
 - Multiple concurrent writes allowed, random value (maximum value, sum, ...) is written.

Different variants may yield different complexities of parallel algorithms.

PRAM Program

AllPairsShortestPaths(W):

```
D = W
for r=1 to s do
  D = MatMin(D, D)
return D
```

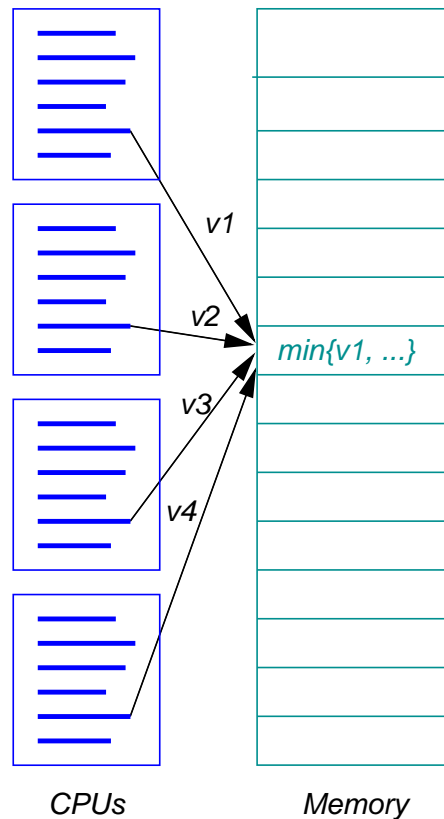
MatMin(D, W):

```
for i=1 to n do in parallel
  for j=1 to n do in parallel
    for k=1 to n do in parallel
      M[i,j,k] =
        min(D[i,j], D[i,k]+W[k,j])
    E[i,j] = MIN(M[i,j])
return E
```

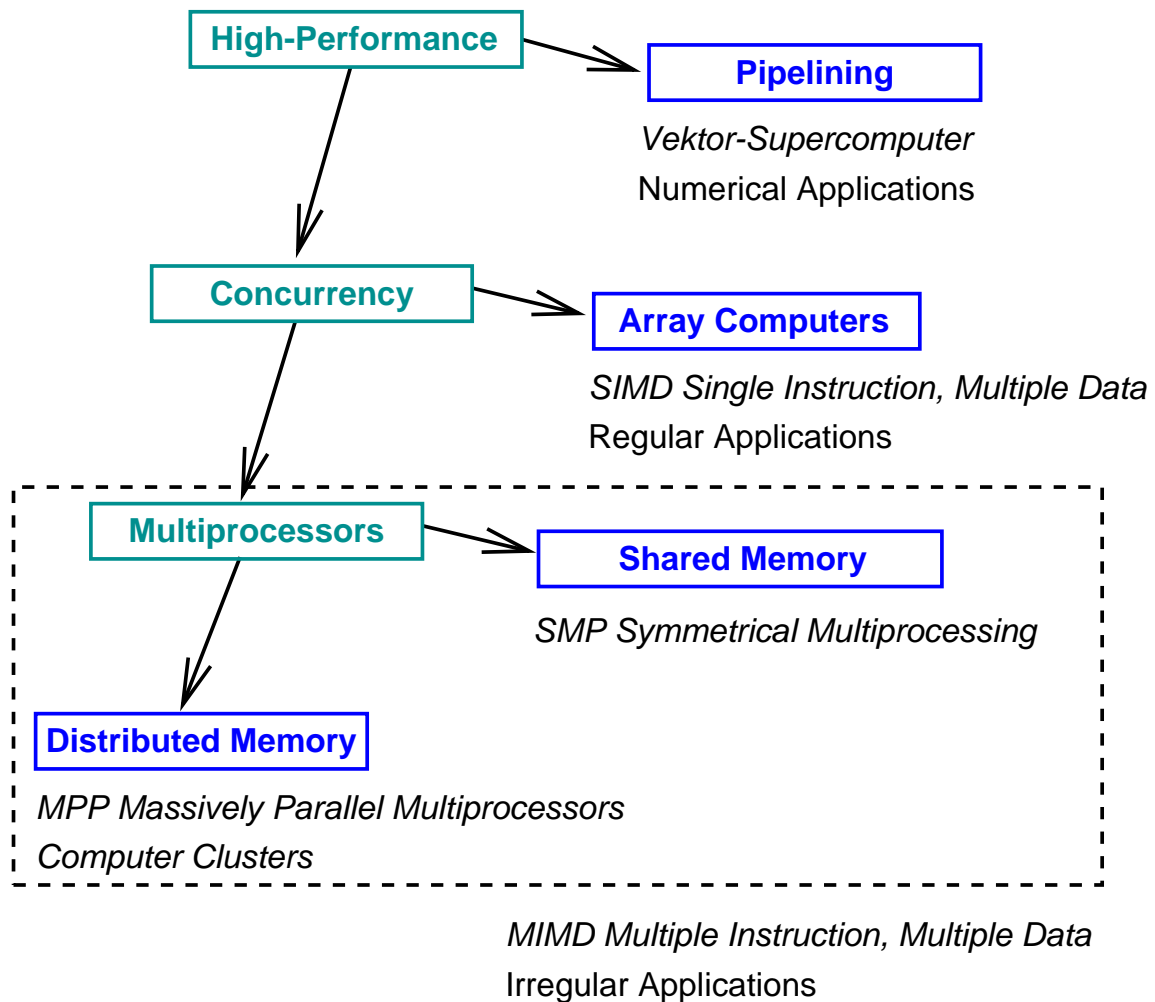
MIN computes minimum of n values.

Complexity of MIN

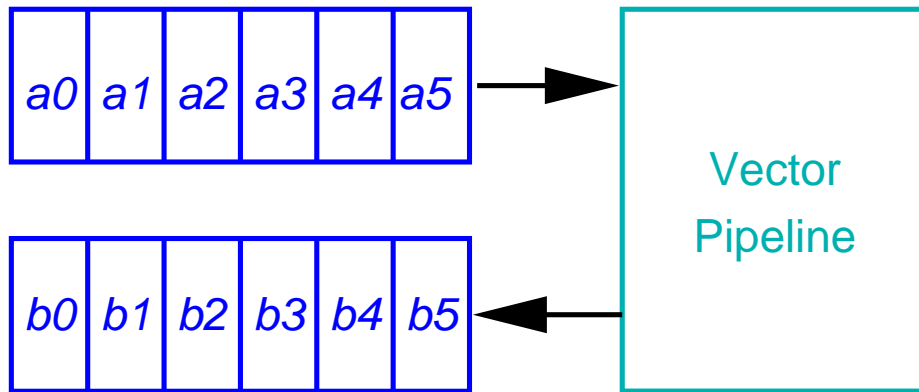
- EREW, CREW: $O(\log n)$
 - Tree-like combination of values.
- CRCW: $O(1)$
 - All values written simultaneously into same cell, minimum value remains.



High-Performance Architectures



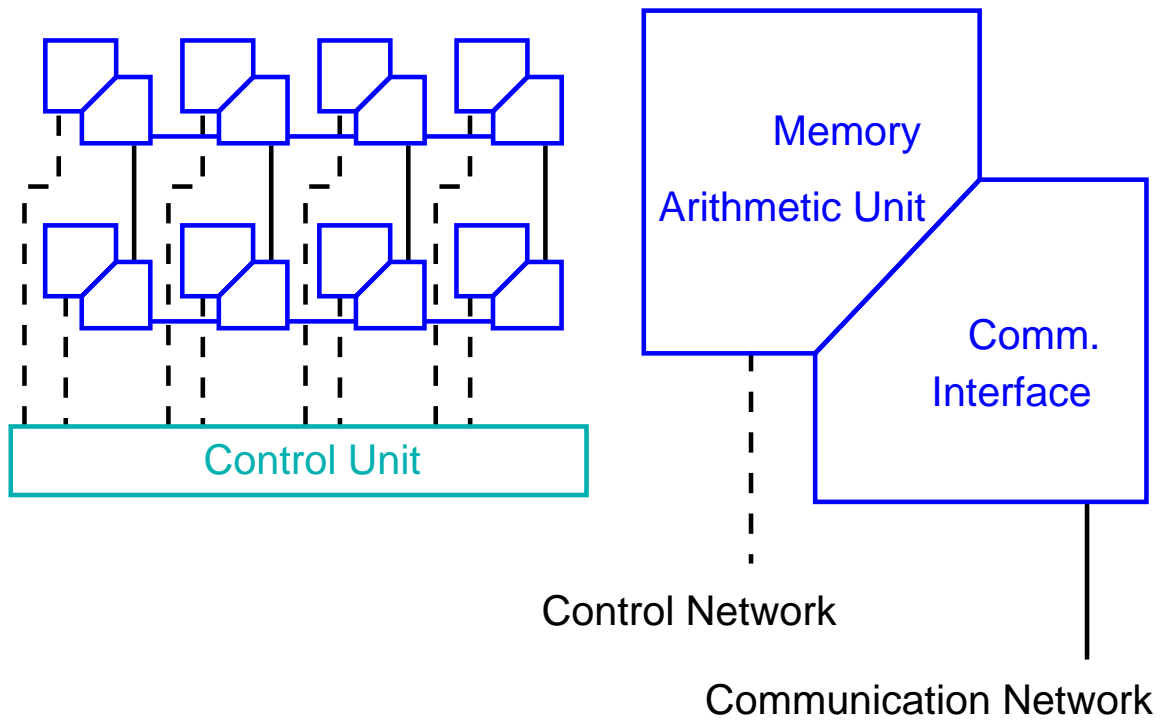
Vector Supercomputers



Vector Registers

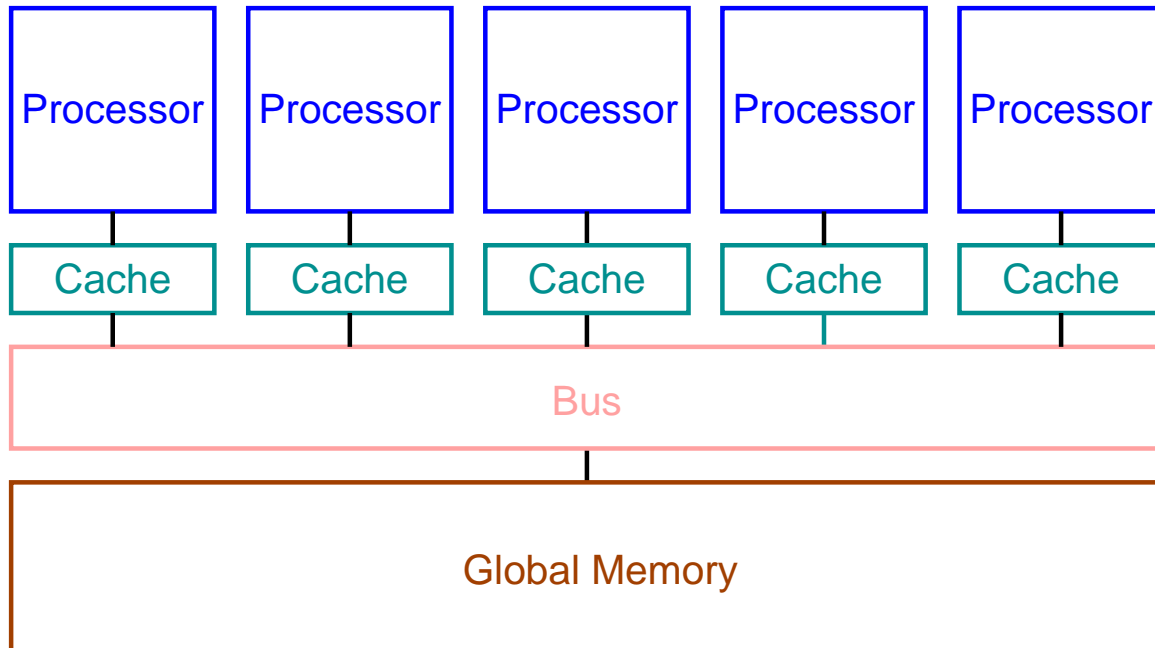
Vectors of numbers are passed through arithmetic pipeline.

SIMD Array Computers



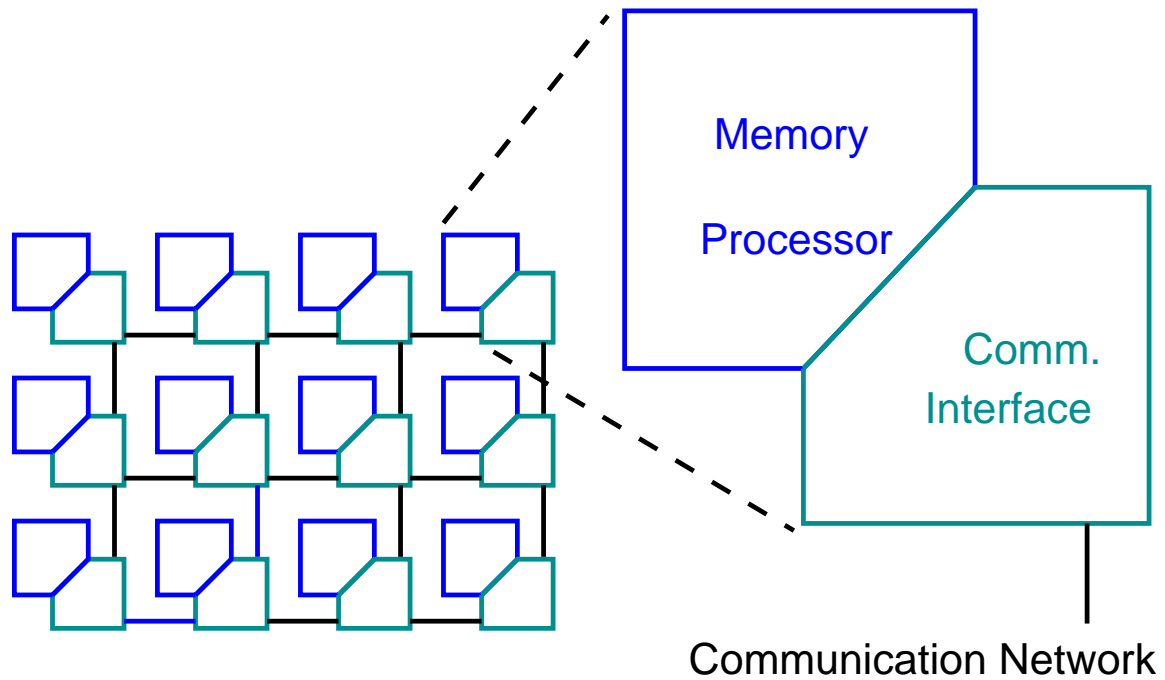
Array of arithmetic units operates and communicates in lock-step.

Shared Memory Multiprocessors



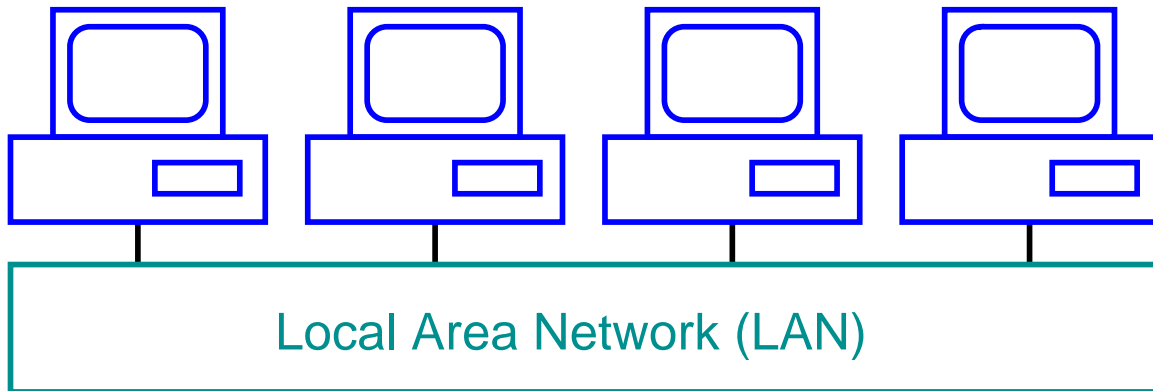
Processors operate asynchronously on shared memory.

Distributed Memory Multiprocessors



Processors operate asynchronously on local memory and communicate by point-to-point network.

Computer Clusters



Cluster of independent computers cooperate via local network.

A Parallel Program

```

cccccccccccccccccccccccccccccccccccccccccccccccccccccccccccc
c      Matrix Multiplication MPI Program      c
c      For this simple version, # of procssors c
c      equals # of columns in matrix         c
c                                             c
c      To run, mpirun -np 4 a.out            c
cccccccccccccccccccccccccccccccccccccccccccccccccccccccccccc

      include 'mpif.h'

      parameter (ncols=4, nrows=4)
      integer a(ncols,nrows), b(ncols,nrows), c(ncols,nrows)
      integer buf(ncols),ans(nrows)
      integer myid, root, numprocs, ierr, status(MPI_STATUS_SIZE)
      integer sender, count

      call MPI_INIT(ierr)
      call MPI_COMM_RANK( MPI_COMM_WORLD, myid, ierr )
      call MPI_COMM_SIZE( MPI_COMM_WORLD, numprocs, ierr )

      if(numprocs.ne.4) then
        print *, "Please run this exercise on 4 processors"
        call MPI_FINALIZE(ierr)
        stop
      endif

      root = 0
      tag = 100
      count = nrows*ncols

c      Master initializes and then dispatches to others
      IF ( myid .eq. root ) then

        do j=1,ncols
          do i=1,nrows
            a(i,j) = 1
            b(i,j) = j
          enddo
        enddo
      enddo

```

A Parallel Program (Contd)

```

c      send a to all other process
      call MPI_BCAST(a,count,MPI_INTEGER,root,MPI_COMM_WORLD,ierr)

c      send one column of b to each other process
      do j=1,numprocs-1
        do i = 1,nrows
          buf(i) = b(i,j+1)
        enddo
      call MPI_SEND(buf,nrows,MPI_INTEGER,j,tag,MPI_COMM_WORLD,ierr)
      enddo

c      Master does his own part here
      do i=1,nrows
        ans(i) = 0
        do j=1,ncols
          ans(i) = ans(i) + a(i,j) * b(i,1)
        enddo
      c(i,1) = ans(i)
      enddo

c      then receives answers from others

      do j=1,numprocs-1
        call MPI_RECV(ans, nrows, MPI_INTEGER, MPI_ANY_SOURCE,
$          MPI_ANY_TAG, MPI_COMM_WORLD, status, ierr)

        sender = status(MPI_SOURCE)
        do i=1,nrows
          c(i,sender+1) = ans(i)
        enddo

      enddo

      do i=1,nrows
        write(6,*)(c(i,j),j=1,ncols)
      enddo

```

A Parallel Program (Contd)

```
ELSE

c      slaves receive a, and one column of b, then compute dot product
      call MPI_BCAST(a,count,MPI_INTEGER,root,MPI_COMM_WORLD,ierr)

      call MPI_RECV(buf, nrows, MPI_INTEGER, root,
$         MPI_ANY_TAG, MPI_COMM_WORLD, status, ierr)

      do i=1,nrows
        ans(i) = 0
        do j=1,ncols
          ans(i) = ans(i) + a(i,j) * buf(j)
        enddo
      enddo

      call MPI_SEND(ans,nrows,MPI_INTEGER,root,0,MPI_COMM_WORLD,ierr)

ENDIF

call MPI_FINALIZE(ierr)

stop
end
```

Literature

- Ian T. Foster, *Designing and Building Parallel Programs — Concepts and Tools for Parallel Software Engineering*, Addison Wesley, Reading, Massachusetts, 1995. Online version: <http://www.mcs.anl.gov/dbpp>
- Michael J. Quinn, *Parallel Computing — Theory and Practice*, 2nd edition, McGraw-Hill, New York, 1994.
- Kai Hwang, *Advanced Computer Architecture: Parallelism, Scalability, Programmability*, McGraw-Hill, New York, 1993.