#### **CSE 613: Parallel Programming**

# Lectures 22 & 23 ( Distributed Memory Algorithms: Dense Matrices )

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## **2D Heat Diffusion**

Let  $h_t(x, y)$  be the heat at point (x, y) at time t.

#### **Heat Equation**

$$\frac{\partial h}{\partial t} = \alpha \left( \frac{\partial^2 h}{\partial x^2} + \frac{\partial^2 h}{\partial y^2} \right), \ \alpha = \text{thermal diffusivity}$$

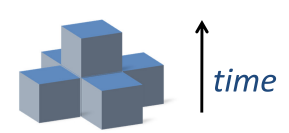
#### Update Equation (on a discrete grid)

$$h_{t+1}(x,y) = h_t(x,y)$$

$$+c_x (h_t(x+1,y) - 2h_t(x,y) + h_t(x-1,y))$$

$$+c_y (h_t(x,y+1) - 2h_t(x,y) + h_t(x,y-1))$$

#### **2D 5-point Stencil**

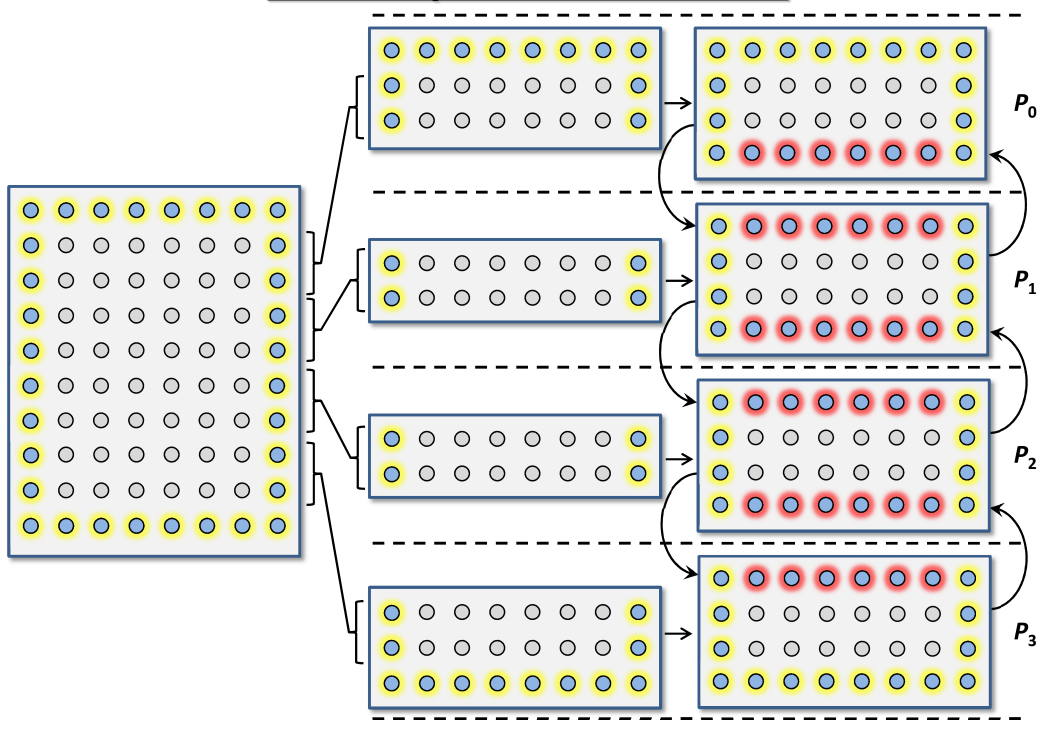


#### **Standard Serial Implementation**

#### **Implementation Tricks**

- Reuse storage for odd and even time steps
- Keep a halo of ghost cells around the array with boundary values

#### One Way of Parallelization



```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreg[0]); }
  if ( myrank > 0 )
                    { MPI Isend( h[ 1 ], Y, MPI FLOAT, myrank - 1, 2 * t + 1, MPI COMM WORLD , & sendreg[ 1 ] );
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
  if ( myrank MPI_Wait( &recvreq[ 0 ], &stat );
  if ( myrank > 0 ) MPI_Wait( &recvreq[ 1 ], &stat );
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
  if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
  if (myrank > 0) MPI_Wait(&sendreg[1], &stat);
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
                                                                      leave enough space
MPI_FLOAT h[ XX + 2 ][ Y + 2 ], g[ XX + 2 ][ Y + 2 ];
                                                                          for ghost cells
MPI Status stat;
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreg[0]); }
 if ( myrank > 0 )
                    { MPI Isend( h[ 1 ], Y, MPI FLOAT, myrank - 1, 2 * t + 1, MPI COMM WORLD , & sendreg[ 1 ] );
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
  if ( myrank MPI_Wait( &recvreq[ 0 ], &stat );
 if ( myrank > 0 ) MPI_Wait( &recvreq[ 1 ], &stat );
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
 if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
 if (myrank > 0) MPI_Wait( &sendreg[ 1], &stat );
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
                                                                    downward send and
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
                                                                      upward receive
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
for ( int t = 0; t < T; ++t )
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                       MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreg[0]); }
  if ( myrank > 0 )
                     \{ MPI | Isend(h[1], Y, MPI | FLOAT, myrank - 1, 2 * t + 1, MPI | COMM | WORLD |, & sendreg[1] \};
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
  if ( mvrank 
  if ( myrank > 0 ) MPI_Wait( &recvreq[ 1 ], &stat );
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
  if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
  if (myrank > 0) MPI_Wait( &sendreg[ 1], &stat );
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
                                                               upward send and
                                                              downward receive
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2/* t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI_Irecv( h[ XX + 1 ], Y, MPI_FLOAT, myrank + 1, 2 * t + 1, MPI_COMM_WORLD , & recvreq[ 0 ] ); }
  if (myrank > 0)
                     { MPI_Isend( h[ 1 ], Y, MPI_FLOAT, myrank - 1, 2 * t + 1, MPI_COMM_WORLD , & sendreg[ 1 ] );
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
  if ( mvrank 
  if ( myrank > 0 ) MPI_Wait( &recvreq[ 1 ], &stat );
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
  if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
  if (myrank > 0) MPI_Wait( &sendreg[ 1], &stat );
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreq[0]); }
 if ( myrank > 0 )
                   \{MPI | Isend(h[1], Y, MPI | FLOAT, myrank - 1, 2 * t + 1, MPI | COMM | WORLD |, & sendreg[1]);
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
                                                                       in addition to the ghost rows exclude
 for ( int x = 2; x < XX; ++x )
                                                                          the two outermost interior rows
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
 if ( myrank MPI Wait ( &recvreg[ 0 ], &stat );
 if ( myrank > 0 ) MPI_Wait( &recvreq[ 1 ], &stat );
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
 if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
 if (myrank > 0) MPI_Wait(&sendreg[1], &stat);
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreg[ 2 ], recvreg[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreq[0]); }
 if ( myrank > 0 )
                    { MPI Isend(h[1], Y, MPI FLOAT, myrank - 1, 2 * t + 1, MPI COMM WORLD, & sendreg[1]);
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
                                                                          wait until data is received
   for ( int y = 1; y <= Y; ++y)
                                                                              for the ghost rows
      g[x][y] = UPDATE(x, y);
  if ( myrank < p - 1 ) MPI_Wait( &recvreg[ 0 ], &stat );
                    MPI_Wait( &recvreq[ 1 ], &stat );
 if ( myrank > 0 )
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
 if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
 if (myrank > 0) MPI_Wait( &sendreg[ 1], &stat );
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreg[0]); }
  if ( myrank > 0 )
                    { MPI Isend( h[ 1 ], Y, MPI FLOAT, myrank - 1, 2 * t + 1, MPI COMM WORLD , & sendreg[ 1 ] );
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
                                                                             update the two
      g[x][y] = UPDATE(x, y);
                                                                        outermost interior rows
  if ( myrank MPI Wait( &recvreg[ 0 ], &stat );
  if (myrank > 0) MPI_Wait(&recvreq[1], &stat);
 for ( int y = 1; y <= Y; ++y ) \{g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); \}
  if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
  if (myrank > 0) MPI_Wait(&sendreg[1], &stat);
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreg[ 2 ], recvreg[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreq[0]); }
 if ( myrank > 0 )
                    { MPI Isend(h[1], Y, MPI FLOAT, myrank - 1, 2 * t + 1, MPI COMM WORLD, & sendreg[1]);
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
                                                                            wait until sending data is complete
  if ( myrank MPI_Wait( &recvreq[ 0 ], &stat );
                                                                                so that h can be overwritten
 if (myrank > 0) MPI_Wait(&recvreq[1], &stat);
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
 if ( myrank < p - 1 ) MPI_Wait( &sendreg[ 0 ], &stat );
                    MPI_Wait( &sendreg[ 1 ], &stat );
 if ( myrank > 0 )
 for ( int x = 1; x \le XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

```
#define UPDATE( u, v ) ( h[u][v] + cx * (h[u+1][v] - 2* h[u][v] + h[u-1][v] ) + cy * (h[u][v+1] - 2* h[u][v] + h[u][v-1] ) )
MPI FLOAT h[XX + 2][Y + 2], g[XX + 2][Y + 2];
MPI Status stat;
MPI_Request sendreq[ 2 ], recvreq[ 2 ];
for ( int t = 0; t < T; ++t)
 if (myrank < p - 1) { MPI_Isend(h[XX], Y, MPI_FLOAT, myrank + 1, 2 * t, MPI_COMM_WORLD, & sendreq[0]);
                        MPI Irecv(h[XX + 1], Y, MPI FLOAT, myrank + 1, 2 * t + 1, MPI COMM WORLD, & recvreq[0]); }
  if ( myrank > 0 )
                    { MPI Isend( h[ 1 ], Y, MPI FLOAT, myrank - 1, 2 * t + 1, MPI COMM WORLD , & sendreg[ 1 ] );
                        MPI_Irecv( h[ 0 ], Y, MPI_FLOAT, myrank - 1, 2 * t , MPI_COMM_WORLD , & recvreq[ 1] ); }
 for ( int x = 2; x < XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      g[x][y] = UPDATE(x, y);
  if ( myrank MPI_Wait( &recvreq[ 0 ], &stat );
  if (myrank > 0) MPI_Wait(&recvreq[1], &stat);
 for (int y = 1; y <= Y; ++y) { g[1][y] = UPDATE(1, y); g[XX][y] = UPDATE(XX, y); }
  if ( myrank MPI_Wait( &sendreg[ 0 ], &stat );
  if (myrank > 0) MPI_Wait( &sendreg[ 1], &stat );
                                                                         now overwrite h
  for ( int x = 1; x <= XX; ++x )
   for ( int y = 1; y <= Y; ++y)
      h[x][y] = g[x][y];
```

#### **Analysis of the MPI Implementation of Heat Diffusion**

Let the dimension of the 2D grid be  $n_X \times n_Y$ , and suppose we execute  $n_T$  time steps. Let p be the number of processors, and suppose the grid is decomposed along X direction.

The computation cost in each time step is clearly  $\frac{n_X n_Y}{p}$ . Hence, the total computation cost,  $t_{comp} = \frac{n_T n_X n_Y}{p}$ .

All processors except processors 0 and p-1 send two rows and receive two rows each in every time step. Processors 0 and p-1 send and receive only one row each. Hence, the total communication  $\cos t$ ,  $t_{comm} = 4n_T(t_S + n_Y t_W)$ , where  $t_S$  is the startup time of a message and  $t_W$  is the per-word transfer time.

Thus 
$$T_p=t_{comp}+t_{comm}=\frac{n_Tn_Xn_Y}{p}+4n_T(t_S+n_Yt_W)$$
, and  $T_1=n_Tn_Xn_Y$ .

#### Naïve Matrix Multiplication

$$\boldsymbol{z}_{ij} = \sum_{k=1}^{n} \boldsymbol{x}_{ik} \boldsymbol{y}_{kj}$$

$$\mathbf{y}_{11}$$
  $\mathbf{y}_{12}$   $\cdots$   $\mathbf{y}_{1n}$ 
 $\mathbf{y}_{21}$   $\mathbf{y}_{22}$   $\cdots$   $\mathbf{y}_{2n}$ 
 $\vdots$   $\vdots$   $\ddots$   $\vdots$ 
 $\mathbf{y}_{n1}$   $\mathbf{y}_{n2}$   $\cdots$   $\mathbf{y}_{nn}$ 

$$Iter-MM(X, Y, Z, n)$$

- 1. for  $i \leftarrow 1$  to n do
- 2. for  $j \leftarrow 1$  to n do
- 3. for  $k \leftarrow 1$  to n do
- 4.  $z_{ij} \leftarrow z_{ij} + x_{ik} \times y_{kj}$

# Naïve Matrix Multiplication

$$\mathbf{z}_{ij} = \sum_{k=1}^{n} \mathbf{x}_{ik} \mathbf{y}_{kj}$$

Suppose we have  $p = n \times n$  processors, and processor  $P_{ij}$  is responsible for computing  $z_{ij}$ .

One master processor initially holds both X and Y, and sends all  $x_{ik}$ and  $y_{kj}$  for k = 1, 2, ..., n to each processor  $P_{ij}$ . One-to-all Broadcast is a bad idea as each processor requires a different part of the input.

Each  $P_{ij}$  computes  $z_{ij}$  and sends back to master.

Thus 
$$t_{comp} = 2n$$
, and  $t_{comm} = n^2(t_s + 2nt_w) + n^2(t_s + t_w)$ .

Hence, 
$$T_p = t_{comp} + t_{comm} = 2n + n^2(2t_s + t_w + 2nt_w)$$
.

Total work,  $T_1 = 2n^3$ .

# Naïve Matrix Multiplication

$$\mathbf{z}_{ij} = \sum_{k=1}^{n} \mathbf{x}_{ik} \mathbf{y}_{kj}$$

Observe that row i of X will be required by all  $P_{i,j}$ ,  $1 \le j \le n$ . So that row can be broadcast to the group  $\{P_{i,1}, P_{i,2}, \dots, P_{i,n}\}$  of size n.

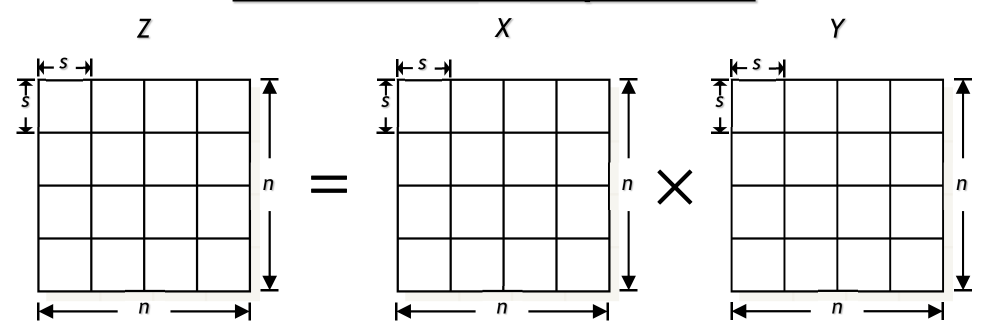
Similarly, for other rows of X, and all columns of Y.

The communication complexity of broadcasting m units of data to a group of size n is  $(t_s + mt_w) \log n$ .

As before, each  $P_{ij}$  computes  $z_{ij}$  and sends back to master.

Hence,  $t_{comm} = 2n(t_s + nt_w) \log n + n^2(t_s + t_w)$ .

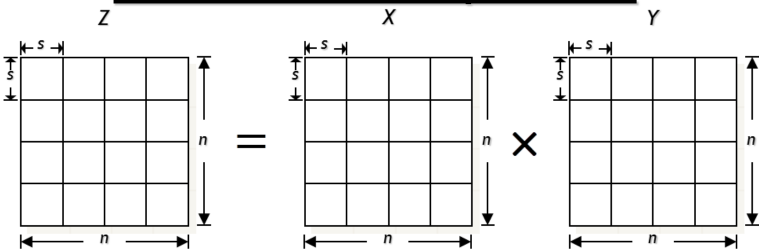
#### **Block Matrix Multiplication**



$$Block-MM(X, Y, Z, n)$$

- 1. for  $i \leftarrow 1$  to n / s do
- 2. for  $j \leftarrow 1$  to n / s do
- 3. for  $k \leftarrow 1$  to n / s do
  - 4. Iter-MM( $X_{ik}$ ,  $Y_{kj}$ ,  $Z_{ij}$ , s)

# **Block Matrix Multiplication**



Suppose  $p = \frac{n}{s} \times \frac{n}{s}$ , and processor  $P_{ij}$  computes block  $Z_{ij}$ .

One master processor initially holds both X and Y, and sends all blocks  $X_{ik}$  and  $Y_{kj}$  for  $k=1,2,\ldots,\frac{n}{s}$  to each processor  $P_{ij}$ .

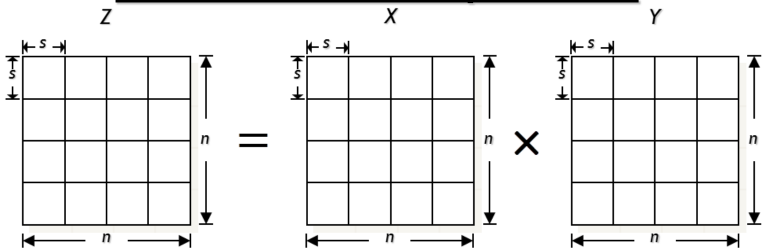
Thus 
$$t_{comp} = \frac{n}{s}(2s^3 + s^2) = O(ns^2)$$
,

and 
$$t_{comm} = \left(\frac{n}{s}\right)^2 \left(2(t_s + nst_w) + (t_s + s^2t_w)\right)$$
. (w/o broadcast)

For 
$$s = \sqrt{n}$$
,  $t_{comp} = O(n^2)$ , and  $t_{comm} = O(nt_s + n^{2.5}t_w)$ 

For 
$$s = n^{\frac{2}{3}}$$
,  $t_{comp} = O(n^{2 + \frac{1}{3}})$ , and  $t_{comm} = O(n^{\frac{2}{3}}t_s + n^{2 + \frac{1}{3}}t_w)$ 

### **Block Matrix Multiplication**



Now consider one-to-group broadcasting.

Block row i of X, i.e., blocks  $X_{ik}$  for  $k=1,2,\ldots,\frac{n}{s}$ , will be required by  $\frac{n}{s}$  different processors, i.e., processors  $P_{ij}$  for  $j=1,2,\ldots,\frac{n}{s}$ .

Similarly, for other block rows of X, and all block columns of Y.

As before, each  $P_{ij}$  computes block  $Z_{ij}$  and sends back to master.

Hence, 
$$t_{comm} = \frac{n}{s}(t_s + nst_w)\log\left(\frac{n}{s}\right) + \left(\frac{n}{s}\right)^2(t_s + s^2t_w).$$

#### **Recursive Matrix Multiplication**

#### Par-Rec-MM(X, Y, Z, n)

- 1. if n = 1 then  $Z \leftarrow Z + X \cdot Y$
- 2. else
- 3. in parallel do

$$Par-Rec-MM (X_{11}, Y_{11}, Z_{11}, n/2)$$

Par-Rec-MM ( 
$$X_{11}$$
,  $Y_{12}$ ,  $Z_{12}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{21}$$
,  $Y_{11}$ ,  $Z_{21}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{21}$$
,  $Y_{12}$ ,  $Z_{22}$ ,  $n / 2$  )

end do

4. in parallel do

Par-Rec-MM ( 
$$X_{12}$$
,  $Y_{21}$ ,  $Z_{11}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{12}$$
,  $Y_{22}$ ,  $Z_{12}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{22}$$
,  $Y_{21}$ ,  $Z_{21}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{22}$$
,  $Y_{22}$ ,  $Z_{22}$ ,  $n / 2$  )

end do

Assuming  $t_s$  and  $t_w$  are constants,

$$t_{comm}(n) = \begin{cases} \Theta(1), & if \ n = 1, \\ 8t_{comm}\left(\frac{n}{2}\right) + \Theta(n^2), & otherwise. \end{cases}$$
$$= \Theta(n^3) \qquad [MT Case 1]$$

Communication cost is too high!

#### **Recursive Matrix Multiplication**

#### Par-Rec-MM(X, Y, Z, n)

- 1. if n = 1 then  $Z \leftarrow Z + X \cdot Y$
- 2. else
- 3. in parallel do

$$Par-Rec-MM (X_{11}, Y_{11}, Z_{11}, n/2)$$

Par-Rec-MM ( 
$$X_{11}$$
,  $Y_{12}$ ,  $Z_{12}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{21}$$
,  $Y_{11}$ ,  $Z_{21}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{21}$$
,  $Y_{12}$ ,  $Z_{22}$ ,  $n / 2$  )

end do

in parallel do 4.

Par-Rec-MM ( 
$$X_{12}$$
,  $Y_{21}$ ,  $Z_{11}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{12}$$
,  $Y_{22}$ ,  $Z_{12}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{22}$$
,  $Y_{21}$ ,  $Z_{21}$ ,  $n / 2$  )

Par-Rec-MM ( 
$$X_{22}$$
,  $Y_{22}$ ,  $Z_{22}$ ,  $n / 2$  )

end do

But with a  $s \times s$  base case,

$$t_{comm}(n) = \begin{cases} \Theta(1), & if \ n \leq s, \\ 8t_{comm}\left(\frac{n}{2}\right) + \Theta(n^2), & otherwise. \end{cases}$$

$$=\Theta\left(\frac{n^3}{s}\right)$$

Parallel running time,

$$t_{comp}(n) = \Theta\left(\frac{n^3}{p} + ns^2\right)$$
 (how?)

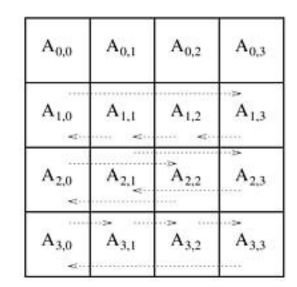
For 
$$s=n^{\frac{2}{3}}$$
,

$$t_{comp} = O\left(\frac{n^3}{p} + n^{2 + \frac{1}{3}}\right),$$
  
and  $t_{comm} = O\left(n^{2 + \frac{1}{3}}\right)$ 

and 
$$t_{comm} = O(n^{2 + \frac{1}{3}})$$

We decompose each matrix into  $\sqrt{p} \times \sqrt{p}$  blocks of size  $\frac{n}{\sqrt{p}} \times \frac{n}{\sqrt{p}}$  each.

We number the processors from  $P_{0,0}$  to  $P_{\sqrt{p}-1,\sqrt{p}-1}$ .

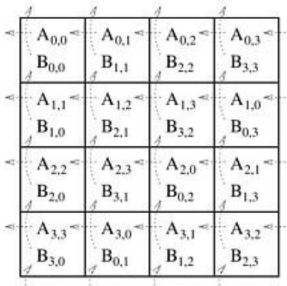


 $\mathbf{B}_{0,1}$  $B_{0,3}$ B<sub>0,2</sub>  $B_{0,0}$  $B_{1,3}$  $\mathbf{B}_{1,1}$  $B_{1,0}$  $B_{1,2}$  $B_{2,1}$  $B_{2,3}$  $B_{2.0}$  $B_{2,2}$  $B_{3,1}$  $B_{3,2}$  $B_{3,3}$  $B_{3.0}$ 

- Initially,  $P_{ij}$  holds  $A_{ij}$  and  $B_{ij}$ .
- We rotate block row i of A to the left by i positions, and block column j of B upward by j positions.

So,  $P_{ij}$  now holds  $A_{i,j+i}$  and  $B_{i+j,j}$ .

(a) Initial alignment of A



(c) A and B after initial alignment

(b) Initial alignment of B

1	.1	.4	1
A <sub>0,1</sub> B <sub>1,0</sub>	A <sub>0,2</sub>	A <sub>0,3</sub>	A <sub>0,0</sub> < B <sub>0,3</sub>
< A <sub>1,2</sub>	< A <sub>1,3</sub> <	B <sub>3,2</sub>	A <sub>1,1</sub> <
B <sub>2,0</sub>	B <sub>3,1</sub>	B <sub>0,2</sub>	B <sub>1,3</sub>
B <sub>3,0</sub>	B <sub>0,1</sub>	B <sub>1,2</sub>	B <sub>2,3</sub>
A <sub>3,0</sub>	A <sub>3,1</sub> < B <sub>1,1</sub>	A <sub>3,2</sub> < B <sub>2,2</sub>	A <sub>3,3</sub> < B <sub>3,3</sub>

(d) Submatrix locations after first shift

 $P_{ij}$  now holds  $A_{i,j+i}$  and  $B_{i+j,j}$ .

 $P_{ij}$  multiplies these two submatrices, and adds the result to  $C_{i,j}$ .

Then in each of the next  $\sqrt{p}-1$  steps, each block row of A is rotated to the left by 1 position, and each block column of B is rotated upward by 1 position. Each  $P_{ij}$  adds the product of its current submatrices to  $C_{i,j}$ .

1	1	1	1
A <sub>0,0</sub>	549 55550	A <sub>0,2</sub> <	A <sub>0,3</sub> B <sub>3,3</sub>
A <sub>1,1</sub>	< A <sub>1,2</sub>	A <sub>1,3</sub> = B <sub>3,2</sub>	A <sub>1,0</sub> <-
A <sub>2,2</sub>	Gent (100 th)	A <sub>2,0</sub> -	A <sub>2,1</sub> <
A <sub>3,3</sub>	A <sub>3,0</sub>	A <sub>3,1</sub> < B <sub>1,2</sub>	A <sub>3,2</sub> <-

(c) A and B after initial alignment

	4	8	d	A
~	A <sub>0,2</sub> <	A <sub>0,3</sub> <	A <sub>0,0</sub> <	A <sub>0,1</sub> <
V	A <sub>1,3</sub> <	B <sub>3,1</sub> A <sub>1,0</sub> <	A <sub>1,1</sub> <	B <sub>1,3</sub>
V	B <sub>3,0</sub>	B <sub>0,1</sub>	B <sub>1,2</sub>	B <sub>2,3</sub>
4	$B_{0,0}$ $A_{3,1} <$	B <sub>1,1</sub> A <sub>3,2</sub> <	B <sub>2,2</sub>	B <sub>3,3</sub>
	B <sub>1,0</sub>	$\mathbf{B}_{2,1}$	B <sub>3,2</sub>	B <sub>0,3</sub>

A <sub>0,1</sub>	A <sub>0,2</sub>	A <sub>0,3</sub> <	A <sub>0,0</sub> <
B <sub>1,0</sub>		B <sub>3,2</sub>	B <sub>0,3</sub>
A <sub>1,2</sub>		A <sub>1,0</sub> < B <sub>0,2</sub>	$A_{1,1} < B_{1,3}$
A <sub>2,3</sub>	A <sub>2,0</sub> A <sub>0,1</sub>	A <sub>2,1</sub> < B <sub>1,2</sub>	A <sub>2,2</sub> ~
A <sub>3,0</sub>	100	A <sub>3,2</sub> < B <sub>2,2</sub>	A <sub>3,3</sub> <

(d) Submatrix locations after first shift

A <sub>0,3</sub>	$A_{0,0} \\ B_{0,1}$	A <sub>0,1</sub>	A <sub>0,2</sub>
B <sub>3,0</sub>		B <sub>1,2</sub>	B <sub>2,3</sub>
A <sub>1,0</sub>	A <sub>1,1</sub>	A <sub>1,2</sub>	A <sub>1,3</sub>
B <sub>0,0</sub>	B <sub>1,1</sub>	B <sub>2,2</sub>	B <sub>3,3</sub>
A <sub>2,1</sub>	A <sub>2,2</sub>	A <sub>2,3</sub>	A <sub>2,0</sub>
B <sub>1,0</sub>	B <sub>2,1</sub>	B <sub>3,2</sub>	B <sub>0,3</sub>
A <sub>3,2</sub>	A <sub>3,3</sub>	A <sub>3,0</sub>	A <sub>3,1</sub> B <sub>1,3</sub>
B <sub>2,0</sub>	B <sub>3,1</sub>	B <sub>0,2</sub>	

(e) Submatrix locations after second shift (f) Submatrix locations after third shift

Initial arrangement makes

 $\sqrt{p}-1$  block rotations of A and B, and one block matrix multiplication per processor.

In each of the next  $\sqrt{p}-1$  steps, each processor performs one block matrix

multiplication, and sends and receives one block each.

$$t_{comp} = 2\sqrt{p} \left(\frac{n}{\sqrt{p}}\right)^3 = O\left(\frac{n^3}{p}\right)^3$$
$$t_{comm} = 4(\sqrt{p} - 1)$$

$$\times \left(t_S + \left(\frac{n}{\sqrt{p}}\right)^2 t_W\right).$$

A	- A	1 A <	A
B <sub>0,0</sub>	349 33000	B <sub>2,2</sub>	B <sub>3,3</sub>
A <sub>1,1</sub>	4.0	A <sub>1,3</sub>	A <sub>1,0</sub> <-
A <sub>2,2</sub> B <sub>2,0</sub>	A2.3	A <sub>2,0</sub> = B <sub>0,2</sub>	A <sub>2,1</sub> < B <sub>1,3</sub>
A <sub>3,3</sub> B <sub>3,0</sub>	18.1	A <sub>3,1</sub> « B <sub>1,2</sub>	A <sub>3,2</sub>

(c) A and B after initial alignment

(d) Submatrix locations after first shift

A <sub>0,3</sub>	$A_{0,0} \\ B_{0,1}$	A <sub>0,1</sub>	A <sub>0,2</sub>
B <sub>3,0</sub>		B <sub>1,2</sub>	B <sub>2,3</sub>
A <sub>1,0</sub>	A <sub>1,1</sub>	A <sub>1,2</sub>	A <sub>1,3</sub>
B <sub>0,0</sub>	B <sub>1,1</sub>	B <sub>2,2</sub>	B <sub>3,3</sub>
A <sub>2,1</sub>	A <sub>2,2</sub>	A <sub>2,3</sub>	A <sub>2,0</sub>
B <sub>1,0</sub>	B <sub>2,1</sub>	B <sub>3,2</sub>	B <sub>0,3</sub>
A <sub>3,2</sub>	A <sub>3,3</sub>	A <sub>3,0</sub>	A <sub>3,1</sub> B <sub>1,3</sub>
B <sub>2,0</sub>	B <sub>3,1</sub>	B <sub>0,2</sub>	

ubmatrix locations after second shift (f) Submatrix locations after third shift

What if initially, one master processor (say,  $P_{0,0}$ ) holds all data (i.e., matrices A and B), and the same processor wants to collect the entire output matrix (i.e., C) at the end?

Processor  $P_{0,0}$  initially sends  $A_{i,j}$  and  $B_{i,j}$  to processor  $P_{i,j}$ , and at the end processor  $P_{i,j}$  sends back  $C_{i,j}$  to  $P_{0,0}$ .

Since there are p processors, and each submatrix has size  $\frac{n}{\sqrt{p}} \times \frac{n}{\sqrt{p}}$ , the additional communication complexity:

$$3p \times \left(t_S + \left(\frac{n}{\sqrt{p}}\right)^2 t_W\right) = 3(pt_S + n^2 t_W).$$

So, the communication complexity increases by a factor of  $\sqrt{p}$ .

### Floyd-Warshall's All-Pairs Shortest Paths

Let G = (V, E, w) be a weighted directed graph with vertex set  $V = \{v_1, v_2, ..., v_n\}$ , edge set E, and weight function w.

The weight of edge  $(v_i, v_j) \in E$  is given by  $w(v_i, v_j)$ .

We construct an  $n \times n$  matrix A as follows:

$$A(i,j) = a_{ij} = \begin{cases} 0, & if \ i = j, \\ \infty, & if (v_i, v_j) \notin E, \\ w(v_i, v_j), & otherwise. \end{cases}$$

Floyd-Warshall's algorithm takes matrix A as input, and returns another  $n \times n$  matrix D as output with

 $D(i,j) = d_{ij} = \text{shortest distance from } v_i \text{ to } v_j \text{ in } G.$ 

#### Floyd-Warshall's All-Pairs Shortest Paths

```
FW-APSP(A, n)

1. D^{(0)} \leftarrow A

2. for k \leftarrow 1 to n do

3. for i \leftarrow 1 to n do

4. for j \leftarrow 1 to n do

5. d_{i,j}^{(k)} \leftarrow min \left\{ d_{i,j}^{(k-1)}, d_{i,k}^{(k-1)} + d_{k,j}^{(k-1)} \right\}

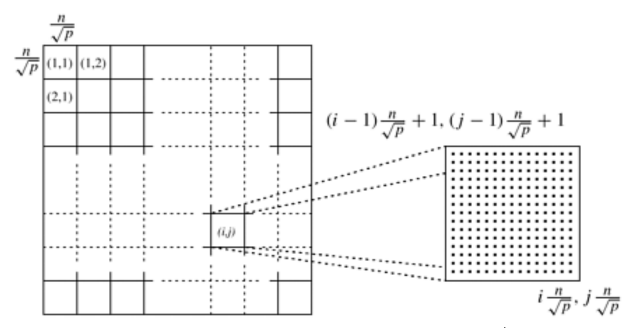
6. return D^{(n)}
```

- can be solved using only  $\Theta(n^2)$  extra space, e.g., using only two  $n \times n$  matrices for storing the values of D
- can be solved in-place in A
- serial running time is  $\Theta(n^3)$

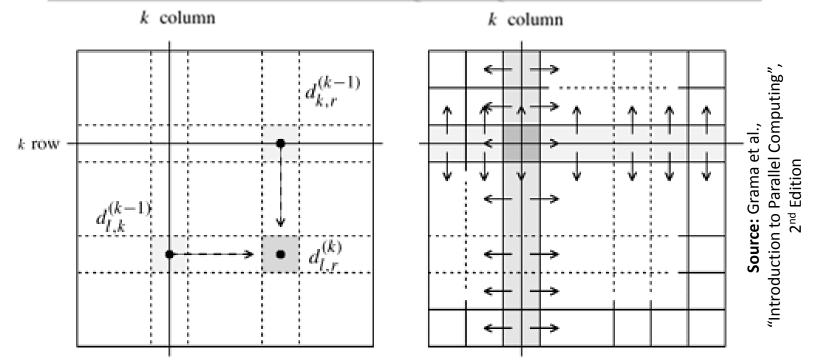
Let p be the number of processing nodes.

We divide  $D^{(k)}$  into  $\sqrt{p} \times \sqrt{p}$  blocks of size  $\frac{n}{\sqrt{p}} \times \frac{n}{\sqrt{p}}$  each.

We assign block (i,j) to processor  $P_{i,j}$  for  $1 \le i,j \le \sqrt{p}$ .

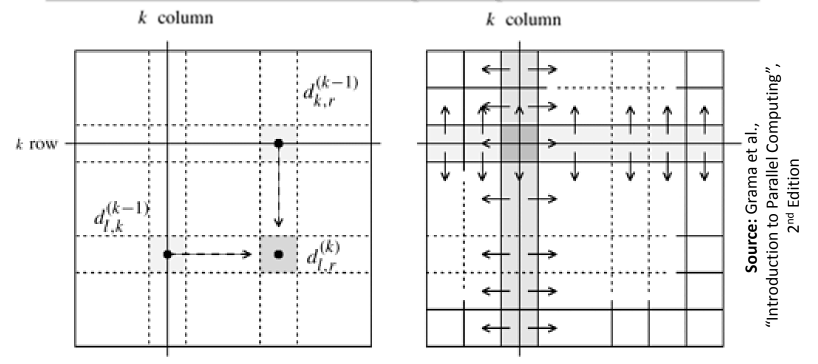


Source: Grama et al., "Introduction to Parallel Computing", 2<sup>nd</sup> Edition



During the computation of  $D^{(k)}$  each processor  $P_{i,j}$  requires

- a segment ( of length  $\frac{n}{\sqrt{p}}$  ) from row k of  $D^{(k-1)}$  which belongs to a processor in block column j
- a segment ( of length  $\frac{n}{\sqrt{p}}$  ) from column k of  $D^{(k-1)}$  which belongs to a processor in block row i



After the computation of  $D^{(k-1)}$  if processor  $P_{i,j}$ 

- contains a segment from row k of  $D^{(k-1)}$ , it broadcasts that segment to all processors in block column j
- contains a segment from column k of  $D^{(k-1)}$ , it broadcasts that segment to all processors in block row i

```
FW-APSP-2D-Block ( <math>D^{(0)} )
```

- 1. for  $k \leftarrow 1$  to n do
- 2. parallel: each node  $P_{i,j}$  does the following:
- 3. if it contains a segment of row k of  $D^{(k-1)}$ , broadcasts that segment to nodes  $P_{*,j}$
- 4. if it contains a segment of column k of  $D^{(k-1)}$ , broadcasts that segment to nodes  $P_{i,*}$
- 5. waits until all nodes receive the needed segments (global sync)
- 6. computes its part of the  $D^{(k)}$  matrix

In each iteration of the for loop ( assuming  $t_{\mathcal{S}}$  and  $t_{\mathcal{W}}$  to be constants )

- Line 3: communication complexity =  $\Theta\left(\frac{n}{\sqrt{p}}\log\sqrt{p}\right)$  (why?)
- Line 4: communication complexity =  $\Theta\left(\frac{n}{\sqrt{p}}\log\sqrt{p}\right)$  (why?)
- **Line 5:** communication complexity =  $\Theta(\log p)$  (sync)
- **Line 6:** computation complexity =  $\Theta(n^2/p)$

#### FW-APSP-2D- $Block(D^{(0)})$

- 1. for  $k \leftarrow 1$  to n do
- 2. parallel: each node  $P_{i,j}$  does the following:
- 3. if it contains a segment of row k of  $D^{(k-1)}$ , broadcasts that segment to nodes  $P_{*,j}$
- 4. if it contains a segment of column k of  $D^{(k-1)}$ , broadcasts that segment to nodes  $P_{i,*}$
- 5. waits until all nodes receive the needed segments (global sync)
- 6. computes its part of the  $D^{(k)}$  matrix

#### Overall:

$$t_{comm} = \Theta\left(n \times \frac{n}{\sqrt{p}}\log p\right) = \Theta\left(\frac{n^2}{\sqrt{p}}\log p\right)$$

and 
$$t_{comp} = \Theta\left(n \times \frac{n^2}{p}\right) = \Theta\left(\frac{n^3}{p}\right)$$

Hence, 
$$T_p = t_{comp} + t_{comm} = \Theta\left(\frac{n^3}{p} + \frac{n^2}{\sqrt{p}}\log p\right)$$

#### Improved Distributed Memory Implementation

```
    for k ← 1 to n do
    parallel: each node P<sub>i,j</sub> does the following:
    if it contains a segment of row k of D<sup>(k-1)</sup>, broadcasts that segment to nodes P<sub>*,j</sub>
    if it contains a segment of column k of D<sup>(k-1)</sup>, broadcasts that segment to nodes P<sub>i,*</sub>
    waits until all nodes receive the needed segments ( global sync )
    computes its part of the D<sup>(k)</sup> matrix
```

The global synchronization in line 5 can be removed without affecting the correctness of the algorithm.

The trick is to use pipelining.

### Pipelined 2D Block Mapping FW-APSP

#### FW-APSP-Pipelined-2D- $Block(<math>D^{(0)}$ )

- 1. for  $k \leftarrow 1$  to n do
- 2. parallel: each node  $P_{i,j}$  does the following:
- 3. if it contains a segment of row k of  $D^{(k-1)}$ , sends that segment to nodes  $P_{i-1,j}$  ( if i>1 ) and  $P_{i+1,j}$  ( if  $i<\sqrt{p}$  )
- 4. if it contains a segment of column k of  $D^{(k-1)}$ , sends that segment to nodes  $P_{i,j-1}$  ( if j>1 ) and  $P_{i,j+1}$  ( if  $j<\sqrt{p}$  )
- 5. waits only until it receives the two segments it needs
- 6. computes its part of the  $D^{(k)}$  matrix, and at any point if it receives data from any direction it stores them locally, and forwards them in the opposite direction

After the computation of row 1 & col 1, all relevant segments of

$$D^{(1)}$$
 reach  $P_{\sqrt{p},\sqrt{p}}$  after  $\Theta\left(\left(n/\sqrt{p}\right)\times\sqrt{p}\right)=\Theta(n)$  time units. (how?)

Successive rows & cols follow after time  $\Theta(n^2/p)$  in pipelined mode.

Hence,  $P_{\sqrt{p},\sqrt{p}}$  completes computation in time  $\Theta(n^3/p) + \Theta(n)$ .

#### Pipelined 2D Block Mapping FW-APSP

#### FW-APSP-Pipelined-2D- $Block(<math>D^{(0)}$ )

- 1. for  $k \leftarrow 1$  to n do
- 2. parallel: each node  $P_{i,j}$  does the following:
- 3. if it contains a segment of row k of  $D^{(k-1)}$ , sends that segment to nodes  $P_{i-1,j}$  ( if i>1 ) and  $P_{i+1,j}$  ( if  $i<\sqrt{p}$  )
- 4. if it contains a segment of column k of  $D^{(k-1)}$ , sends that segment to nodes  $P_{i,j-1}$  ( if j>1 ) and  $P_{i,j+1}$  ( if  $j<\sqrt{p}$  )
- 5. waits only until it receives the two segments it needs
- 6. computes its part of the  $D^{(k)}$  matrix, and at any point if it receives data from any direction it stores them locally, and forwards them in the opposite direction

When  $P_{\sqrt{p},\sqrt{p}}$  completes iteration n-1, it sends the relevant values of row n and column n to other nodes.

These values reach  $P_{1,1}$  in time  $\Theta(n)$ .

Hence, 
$$T_p = t_{comp} + t_{comm} = \Theta\left(\frac{n^3}{p}\right) + \Theta(n) = \Theta\left(\frac{n^3}{p} + n\right)$$