

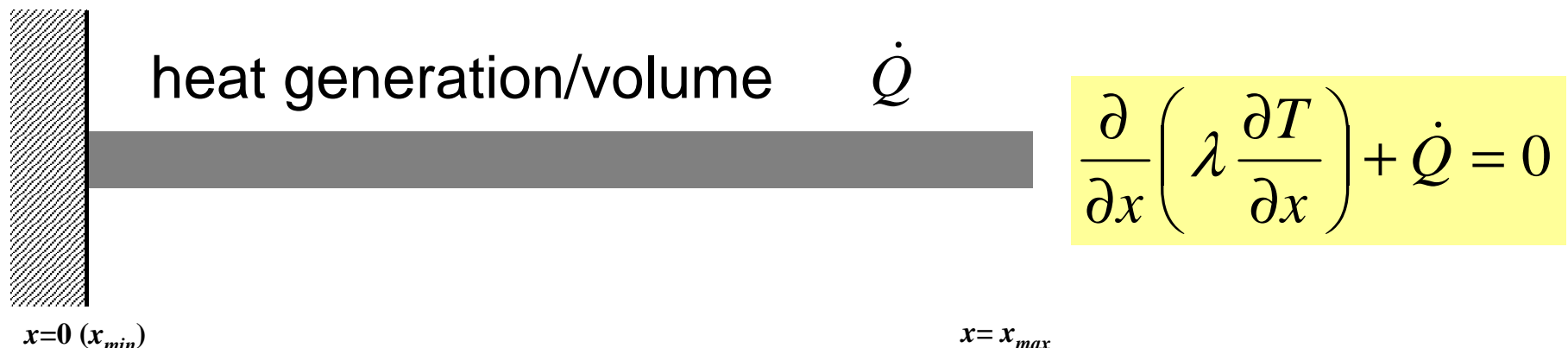
Report S2

Fortran

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Information Technology Center
The University of Tokyo

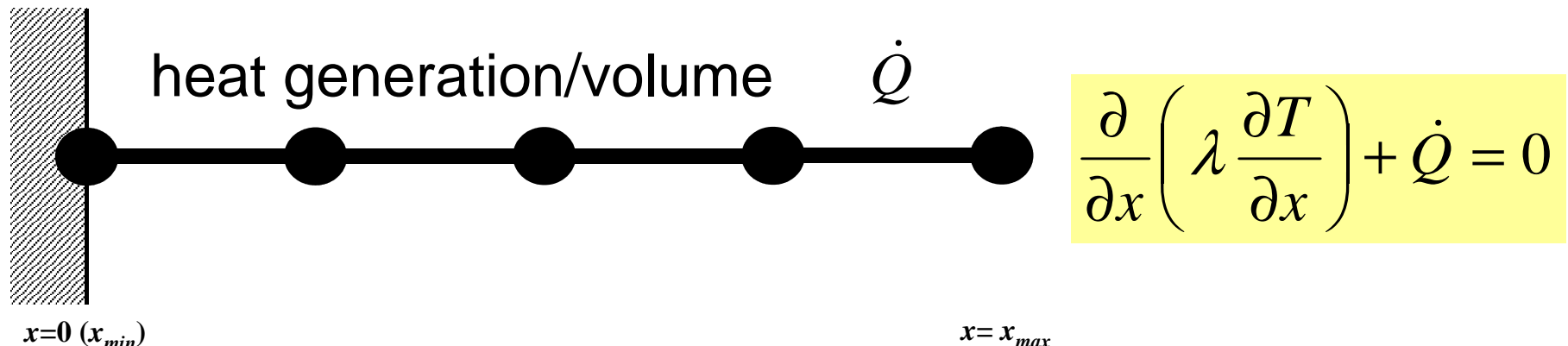
- Overview
- Distributed Local Data
- Program
- Results

1D Steady State Heat Conduction



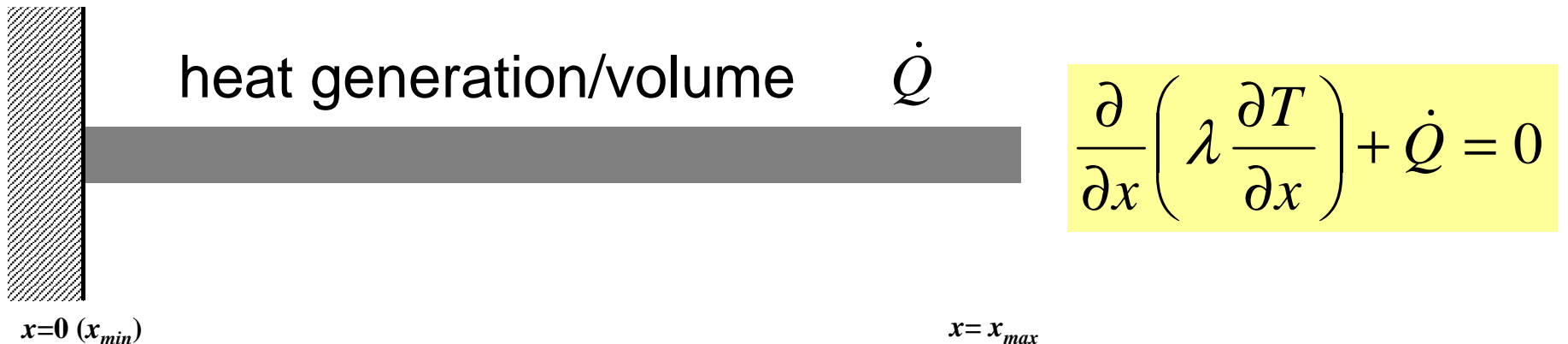
- Uniform: Sectional Area: A , Thermal Conductivity: λ
- Heat Generation Rate/Volume/Time [QL⁻³T⁻¹] \dot{Q}
- Boundary Conditions
 - $x=0$: $T=0$ (Fixed Temperature)
 - $x=x_{max}$: $\frac{\partial T}{\partial x} = 0$ (Insulated)

1D Steady State Heat Conduction



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Analytical Solution



$$T = 0 @ x = 0$$

$$\frac{\partial T}{\partial x} = 0 @ x = x_{max}$$

$$\lambda T'' = -\dot{Q}$$

$$\lambda T' = -\dot{Q}x + C_1 \Rightarrow C_1 = \dot{Q}x_{max}, \quad T' = 0 @ x = x_{max}$$

$$\lambda T = -\frac{1}{2}\dot{Q}x^2 + C_1x + C_2 \Rightarrow C_2 = 0, \quad T = 0 @ x = 0$$

$$\therefore T = -\frac{1}{2\lambda}\dot{Q}x^2 + \frac{\dot{Q}x_{max}}{\lambda}x$$

Copy and Compile

Fortran

```
>$ cd /lustre/gt14/t14XXX/pFEM  
>$ cp /lustre/gt00/z30088/class_eps/F/s2r-f.tar .  
>$ tar xvf s2r-f.tar
```

C

```
>$ cd /lustre/gt14/t14XXX/pFEM  
>$ cp /lustre/gt00/z30088/class_eps/C/s2r-c.tar .  
>$ tar xvf s2r-c.tar
```

Confirm/Compile

```
>$ cd mpi/S2-ref  
>$ mpiifort -O3 -xCORE-AVX2 -align array32byte 1d.f  
>$ mpicc -O3 -xCORE-AVX2 -align 1d.c
```

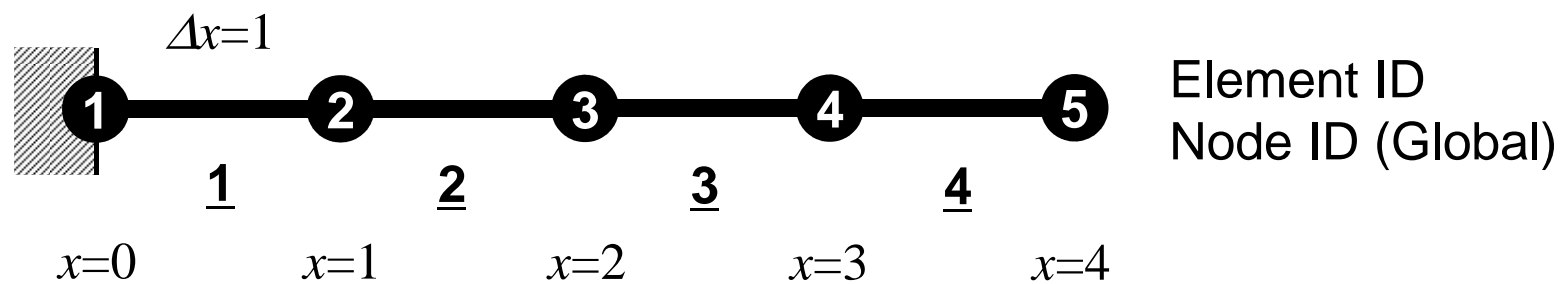
<\$O-S2r> = <\$O-TOP>/mpi/S2-ref

Control File: input.dat

Control Data input.dat

```
4  
1.0  1.0  1.0  1.0  
100  
1.e-8
```

NE (Number of Elements)
 Δx (Length of Each Elem.: L), Q, A, λ
Number of MAX. Iterations for CG Solver
Convergence Criteria for CG Solver



go.sh

```

#!/bin/sh
#PBS -q u-lecture4
#PBS -N test
#PBS -l select=4:mpiprocs=32
#PBS -Wgroup_list=gt14
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o test.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=32
mpirun ./impimap.sh ./a.out

```

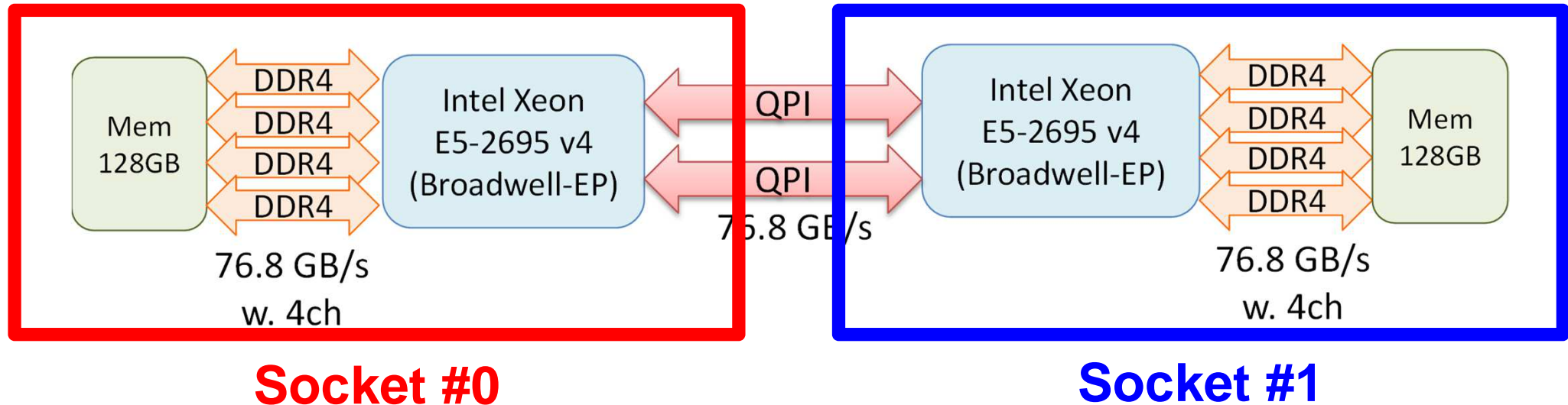
Name of "QUEUE"
Job Name
node#, proc#/node
Group Name (Wallet)
Computation Time
Standard Error
Standard Outpt

go to current dir
(ESSENTIAL)

Execution on each socket
MPI proc#/node (=mpiprocs)
Exec's

#PBS -l select=1:mpiprocs=4	1-node, 4-proc's
#PBS -l select=1:mpiprocs=16	1-node, 16-proc's
#PBS -l select=1:mpiprocs=36	1-node, 36-proc's
#PBS -l select=2:mpiprocs=32	2-nodes, 32x2=64-proc's
#PBS -l select=8:mpiprocs=36	8-nodes, 36x8=288-proc's

export I_MPI_PIN_DOMAIN=socket



- Each Node of Reedbush-U
 - 2 Sockets (CPU's) of Intel Broadwell-EP
 - Each socket has 18 cores
- Each core of a socket can access to the memory on the other socket : NUMA (Non-Uniform Memory Access)
 - `I_MPI_PIN_DOMAIN=socket`, `impimap.sh`: local memory to be used

Procedures for Parallel FEM

- Reading control file, entire element number etc.
 - Creating “distributed local data” in the program
 - Assembling local and global matrices for linear solvers
 - Solving linear equations by CG
-
- Not so different from those of original code

- Overview
- **Distributed Local Data**
- Program
- Results

Finite Element Procedures

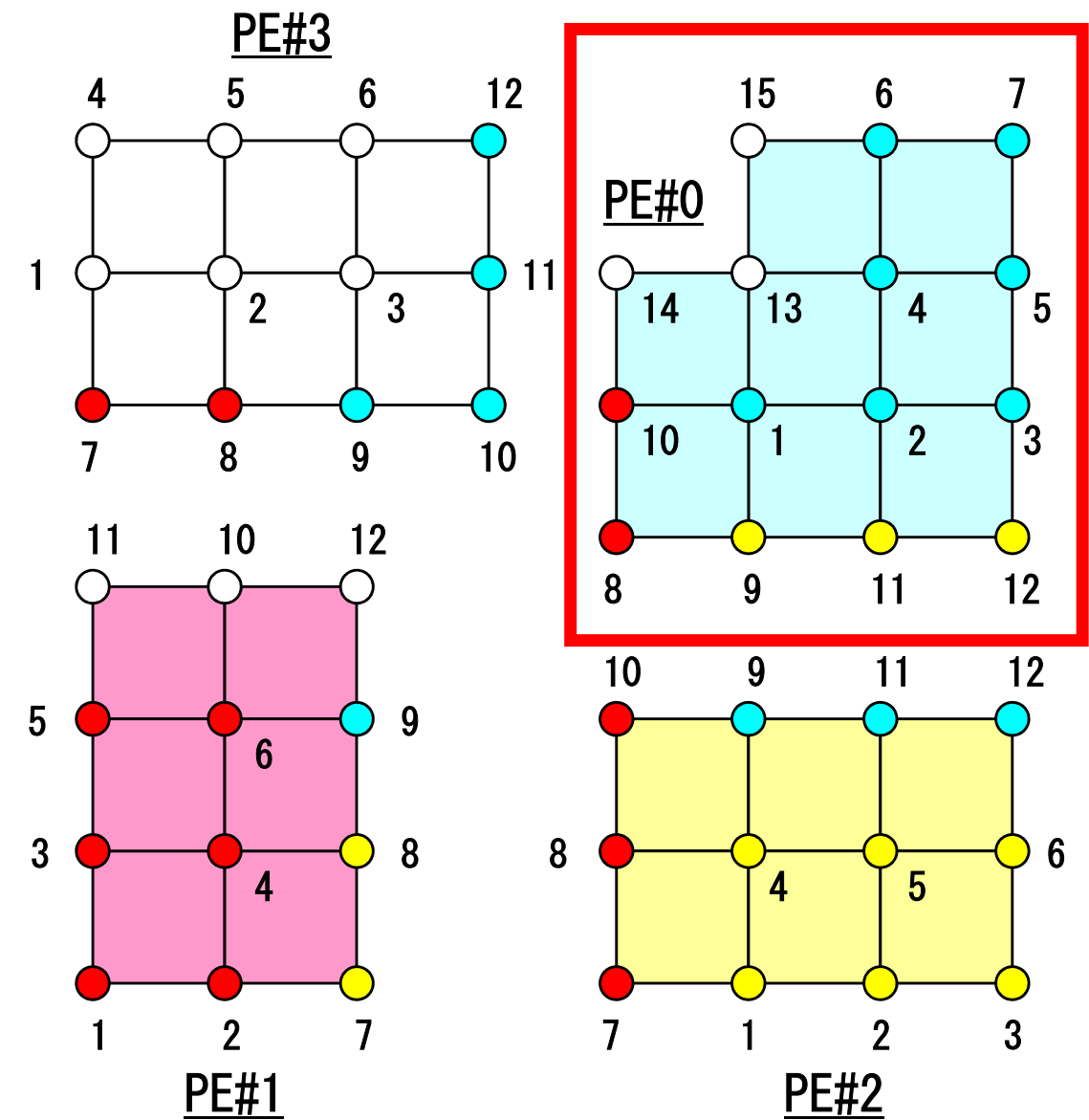
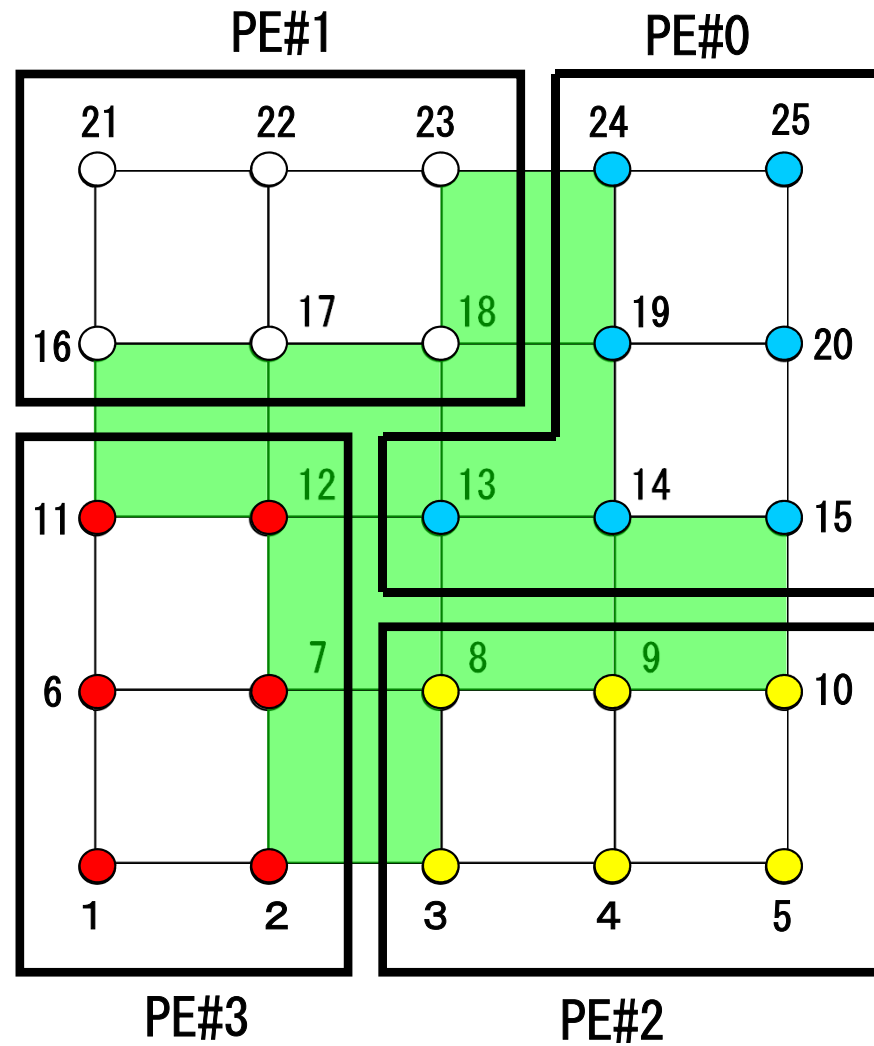
- Initialization
 - Control Data
 - Node, Connectivity of Elements (N: Node#, NE: Elem#)
 - Initialization of Arrays (Global/Element Matrices)
 - Element-Global Matrix Mapping (Index, Item)
- Generation of Matrix
 - Element-by-Element Operations (do $icel= 1, NE$)
 - Element matrices
 - Accumulation to global matrix
 - Boundary Conditions
- Linear Solver
 - Conjugate Gradient Method

Distributed Local Data Structure for Parallel FEM

- Node-based partitioning
- Local data includes:
 - Nodes originally assigned to the domain/PE/partition
 - Elements which include above nodes
 - Nodes which are included above elements, and originally NOT-assigned to the domain/PE/partition
- 3 categories for nodes
 - Internal nodes Nodes originally assigned to the domain/PE/partition
 - External nodes Nodes originally NOT-assigned to the domain/PE/partition
 - Boundary nodes External nodes of other domains/PE's/partitions
- Communication tables
- Global info. is not needed except relationship between domains
 - Property of FEM: local element-by-element operations

Node-based Partitioning

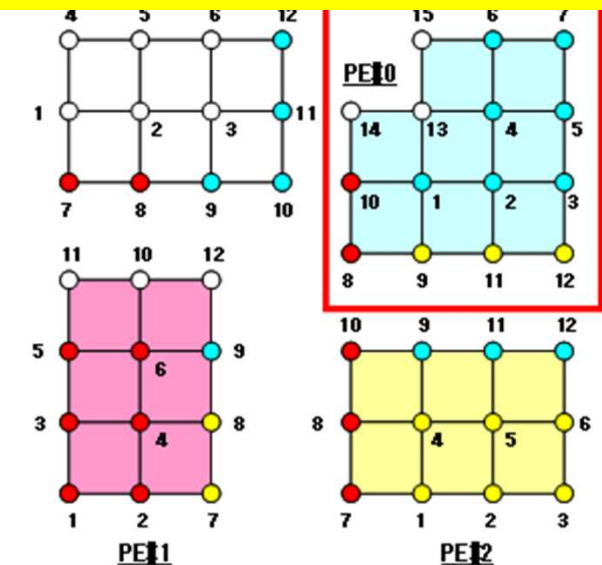
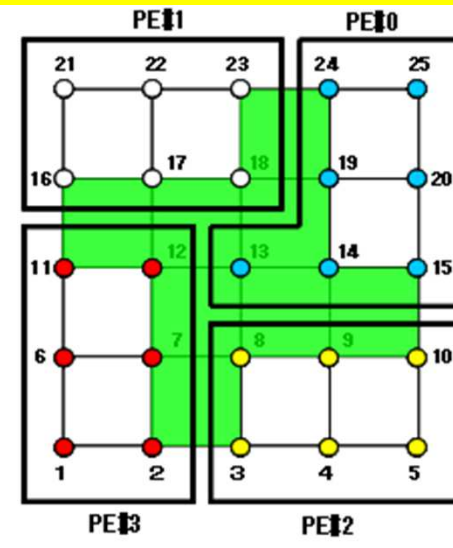
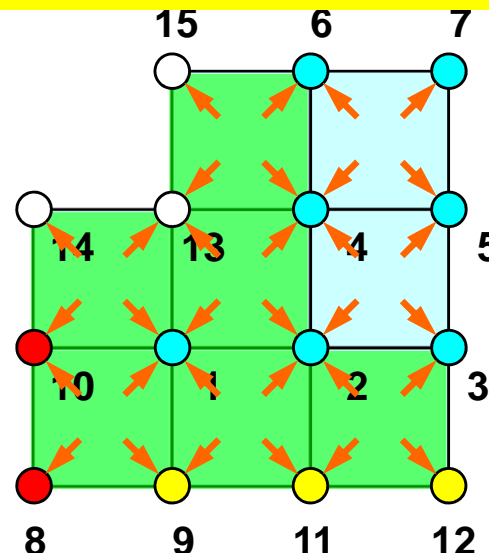
internal nodes - elements - external nodes



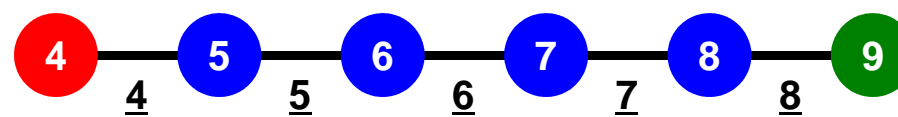
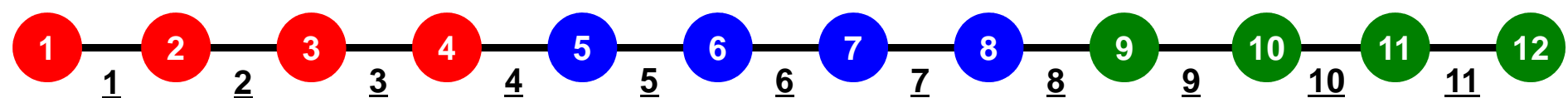
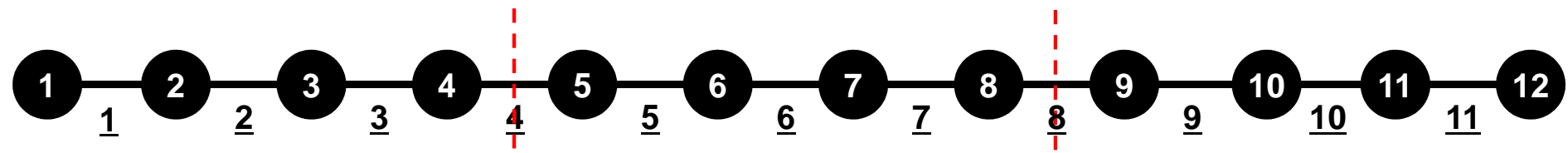
Node-based Partitioning

internal nodes - elements - external nodes

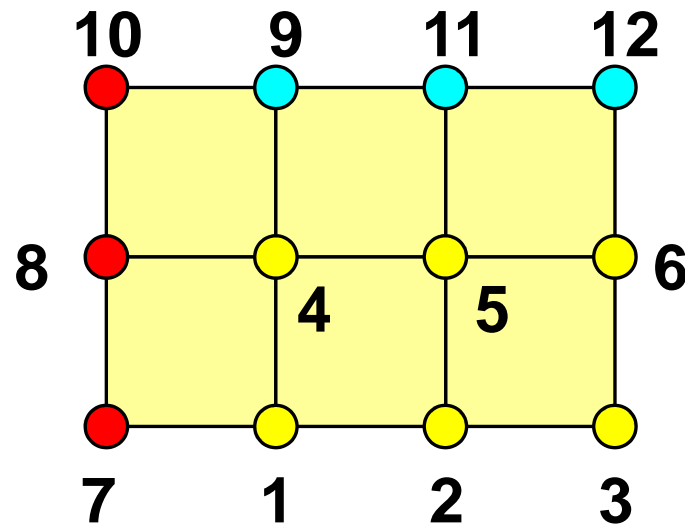
- Partitioned nodes themselves (Internal Nodes) 内点
- Elements which include Internal Nodes 内点を含む要素
- External Nodes included in the Elements 外点
in overlapped region among partitions.
- Info of External Nodes are required for completely local element-based operations on each processor.



1D FEM: 12 nodes/11 elem's/3 domains



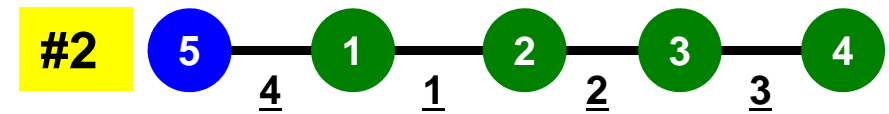
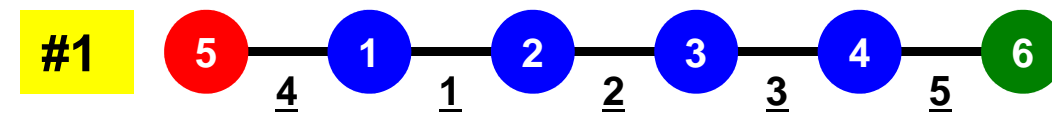
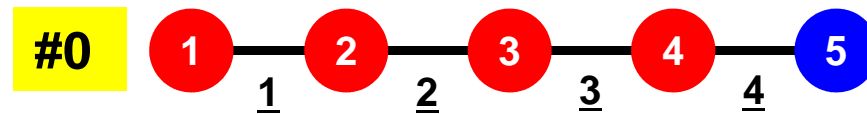
Description of Distributed Local Data



- Internal/External Points
 - Numbering: Starting from internal pts, then external pts after that
- Neighbors
 - Shares overlapped meshes
 - Number and ID of neighbors
- External Points
 - From where, how many, and which external points are received/imported ?
- Boundary Points
 - To where, how many and which boundary points are sent/exported ?

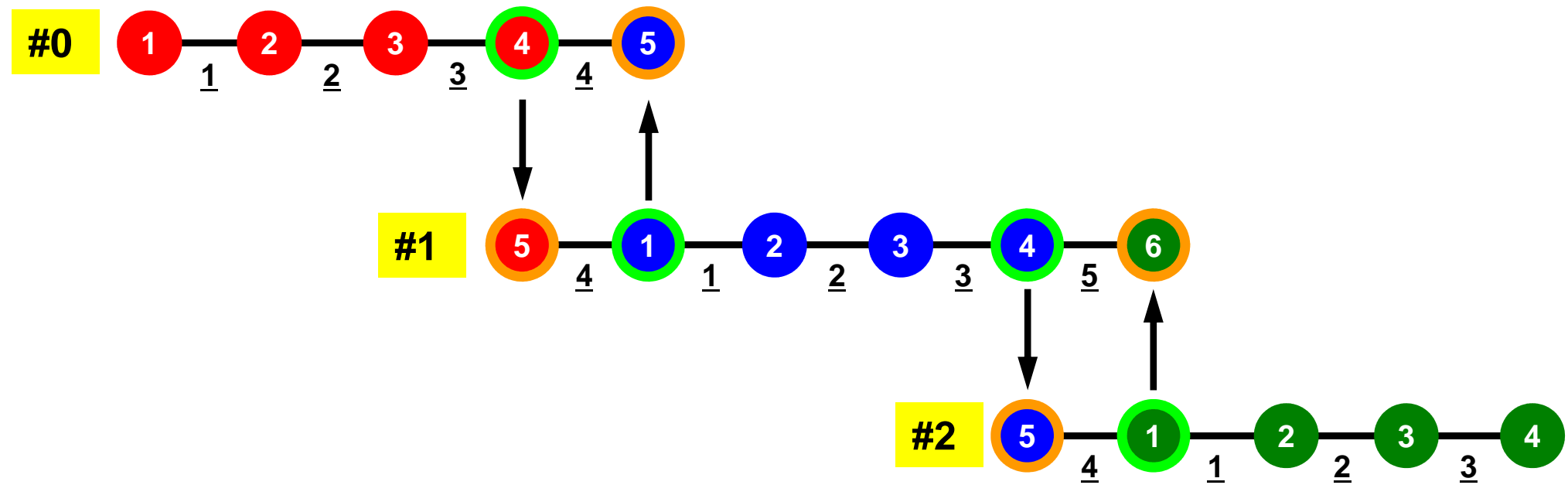
1D FEM: 12 nodes/11 elem's/3 domains

Local ID: Starting from 1 for node and elem at each domain

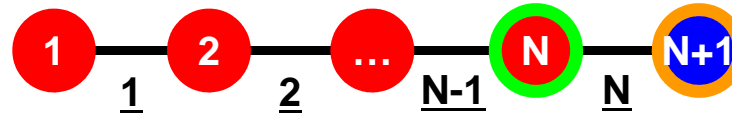


1D FEM: 12 nodes/11 elem's/3 domains

Internal/External Nodes



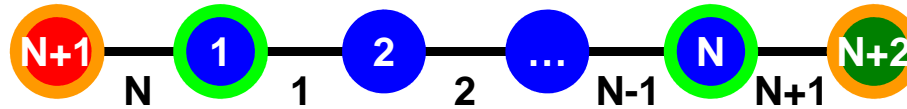
1D FEM: Numbering of Local ID



#0:
N+1 nodes
N elements



#PETot-1:
N+1 nodes
N elements

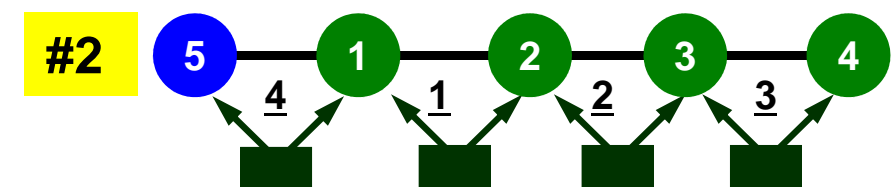
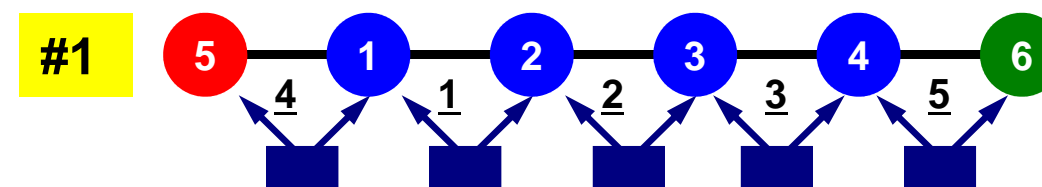
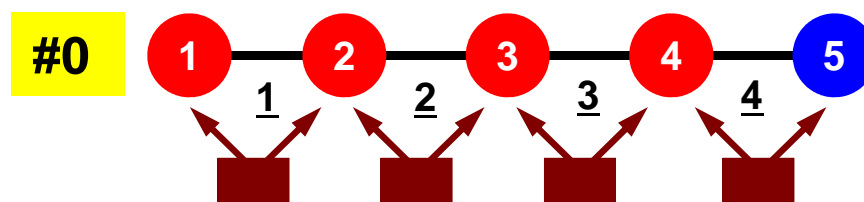


Others (General):
N+2 nodes
N+1 elements

1D FEM: 12 nodes/11 elem's/3 domains

Integration on each element, element matrix -> global matrix

Operations can be done by info. of internal/external nodes
and elements which include these nodes



Preconditioned Conjugate Gradient Method (CG)

```

Compute  $\mathbf{r}^{(0)} = \mathbf{b} - [\mathbf{A}] \mathbf{x}^{(0)}$ 
for i= 1, 2, ...
    solve  $[\mathbf{M}] \mathbf{z}^{(i-1)} = \mathbf{r}^{(i-1)}$ 
     $\rho_{i-1} = \mathbf{r}^{(i-1)} \cdot \mathbf{z}^{(i-1)}$ 
    if i=1
         $\mathbf{p}^{(1)} = \mathbf{z}^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $\mathbf{p}^{(i)} = \mathbf{z}^{(i-1)} + \beta_{i-1} \mathbf{p}^{(i-1)}$ 
    endif
     $\mathbf{q}^{(i)} = [\mathbf{A}] \mathbf{p}^{(i)}$ 
     $\alpha_i = \rho_{i-1} / \mathbf{p}^{(i)} \cdot \mathbf{q}^{(i)}$ 
     $\mathbf{x}^{(i)} = \mathbf{x}^{(i-1)} + \alpha_i \mathbf{p}^{(i)}$ 
     $\mathbf{r}^{(i)} = \mathbf{r}^{(i-1)} - \alpha_i \mathbf{q}^{(i)}$ 
    check convergence  $|\mathbf{r}|$ 
end

```

Preconditioning:
 Diagonal Scaling
 (or Point Jacobi)

Preconditioning, DAXPY

Local Operations by Only Internal Points: Parallel Processing is possible

```

!C
!C-- {z} = [Minv] {r}

do i= 1, N
    W(i, Z)= W(i, DD) * W(i, R)
enddo

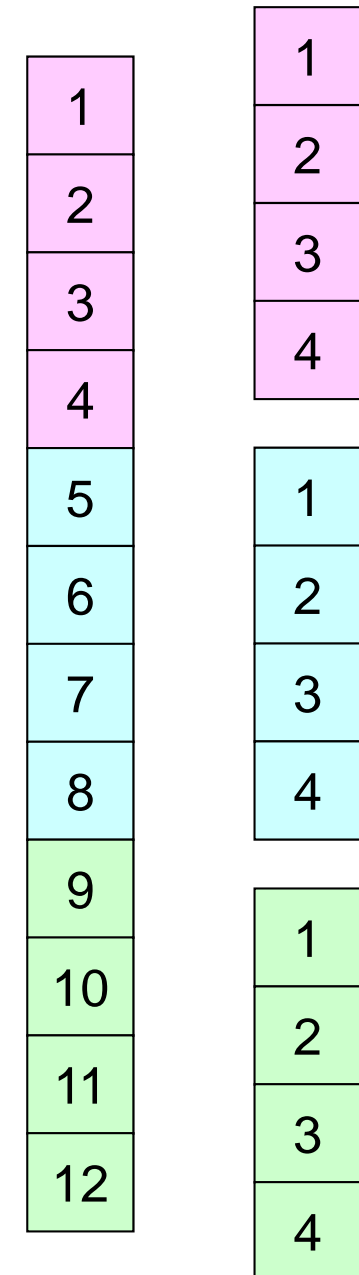
```

```

!C
!C-- {x} = {x} + ALPHA*{p}
!C   {r} = {r} - ALPHA*{q}

do i= 1, N
    PHI(i)= PHI(i) + ALPHA * W(i, P)
    W(i, R)= W(i, R) - ALPHA * W(i, Q)
enddo

```

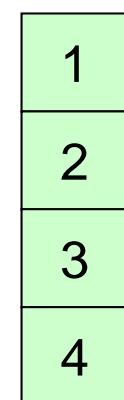
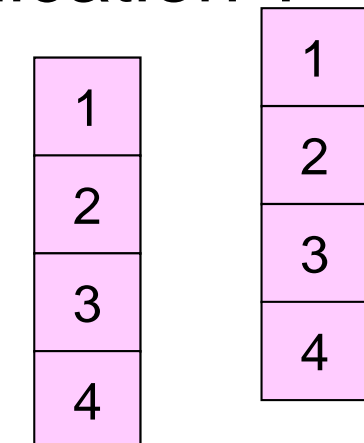


Dot Products

Global Summation needed: Communication ?

```
!C  
!C-- ALPHA= RHO / {p} {q}
```

```
C1= 0. d0  
do i= 1, N  
  C1= C1 + W(i, P)*W(i, Q)  
enddo  
ALPHA= RHO / C1
```

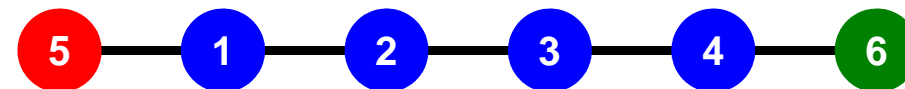


Matrix-Vector Products

Values at External Points: P-to-P Communication

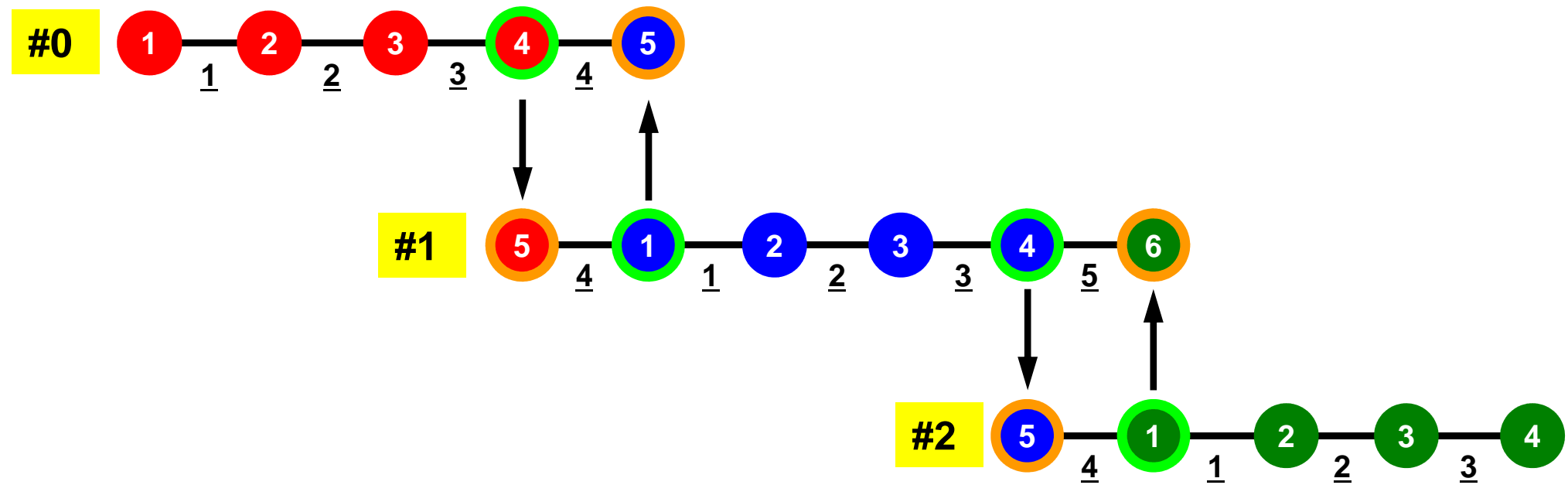
```
!C  
!C-- {q} = [A] {p}
```

```
do i= 1, N  
  W(i, Q) = DIAG(i)*W(i, P)  
  do j= INDEX(i-1)+1, INDEX(i)  
    W(i, Q) = W(i, Q) + AMAT(j)*W(ITEM(j), P)  
  enddo  
enddo
```



1D FEM: 12 nodes/11 elem's/3 domains

Internal/External Nodes



Mat-Vec Products: Local Op. Possible

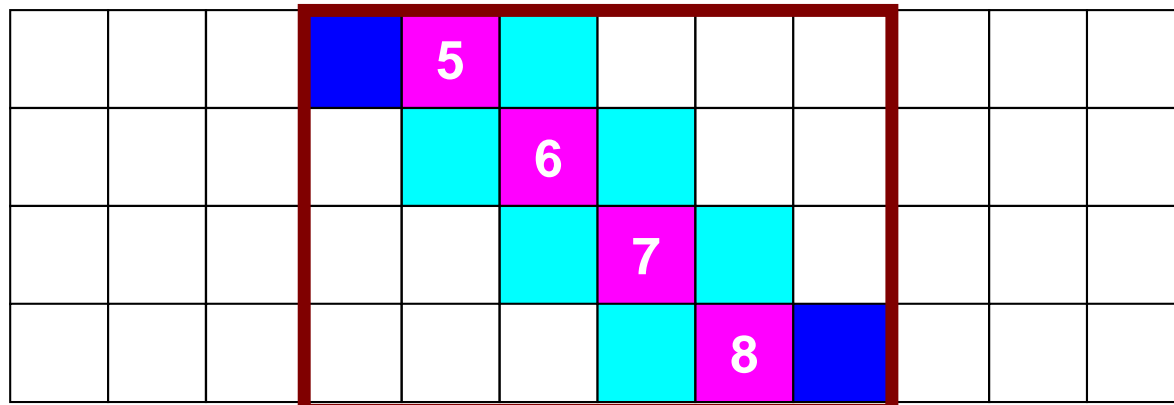
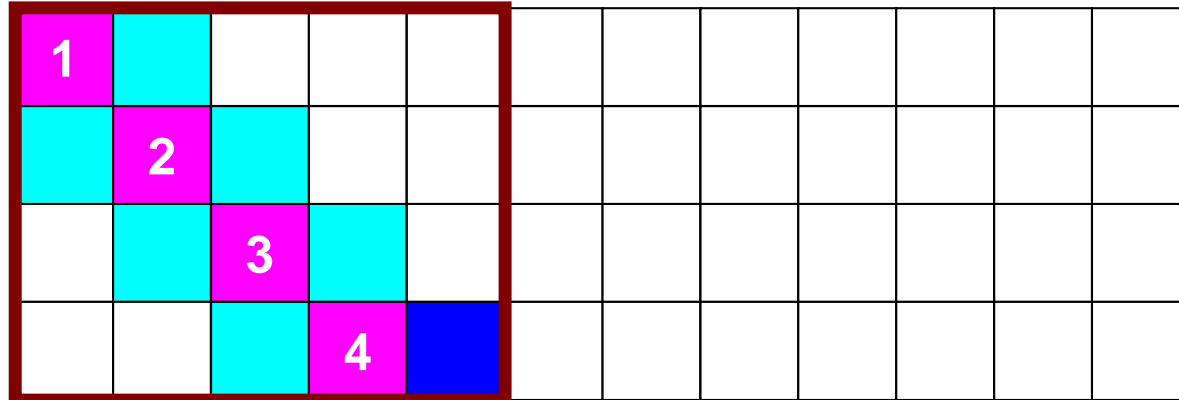
1												
	2											
		3										
			4									
				5								
					6							
						7						
							7					
								9				
									10			
										11		
											12	

1
2
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8
9
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11
12

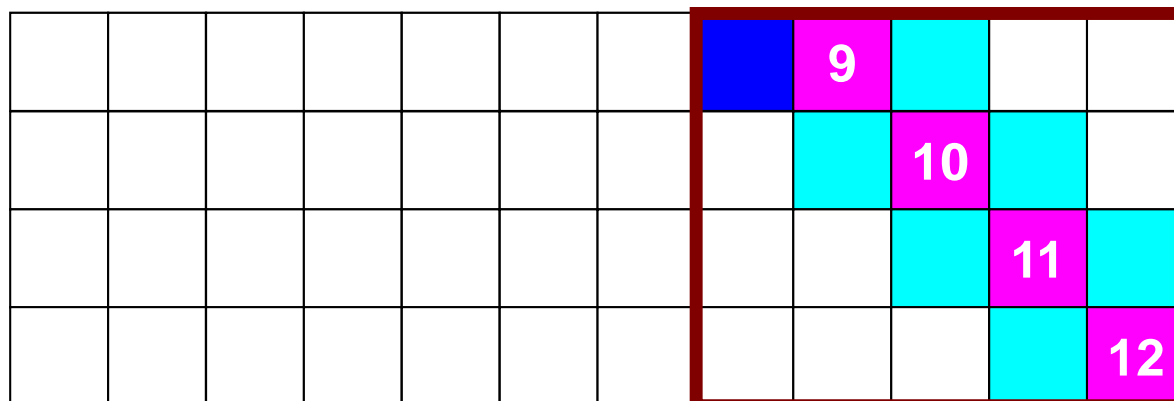
=

1
2
3
4
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10
11
12

Mat-Vec Products: Local Op. Possible



=



Mat-Vec Products: Local Op. Possible

1				
	2			
		3		
			4	

1
2
3
4

1
2
3
4

	5					
		6				
			7			
				8		

5
6
7
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5
6
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8

=

	9				
		10			
			11		

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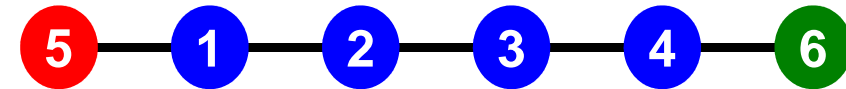
Mat-Vec Products: Local Op. #1

$$\begin{array}{c} \begin{array}{|c|c|c|c|c|c|} \hline & \textcolor{blue}{\boxed{}} & \textcolor{magenta}{\boxed{1}} & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline & \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{2}} & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline & \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{3}} & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{4}} & \textcolor{magenta}{\boxed{}} & \textcolor{blue}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline \end{array} \quad \begin{array}{|c|} \hline 1 \\ \hline 2 \\ \hline 3 \\ \hline 4 \\ \hline \end{array} \end{array} = \begin{array}{|c|} \hline 1 \\ \hline 2 \\ \hline 3 \\ \hline 4 \\ \hline \end{array}$$

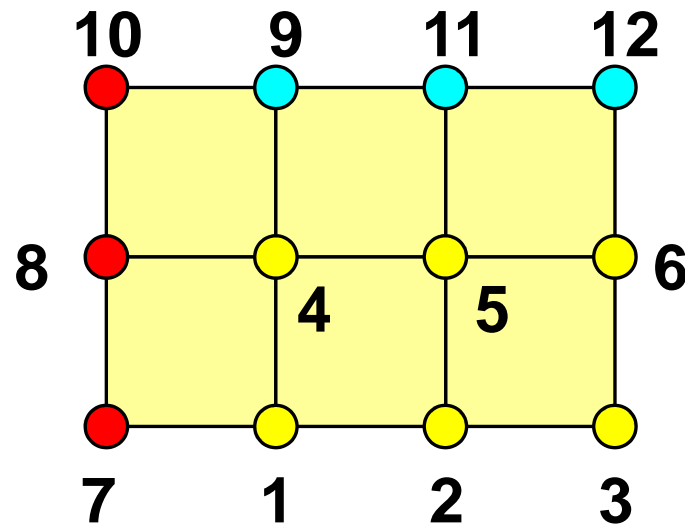
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$$\begin{array}{c} \begin{array}{|c|c|c|c|c|c|} \hline \textcolor{magenta}{\boxed{1}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{blue}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{2}} & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline & \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{3}} & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline & \textcolor{cyan}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{4}} & \textcolor{magenta}{\boxed{}} & \textcolor{blue}{\boxed{}} \\ \hline & \textcolor{white}{\boxed{}} & \textcolor{white}{\boxed{}} & \textcolor{cyan}{\boxed{}} & \textcolor{magenta}{\boxed{}} & \textcolor{white}{\boxed{}} \\ \hline \end{array} \quad \begin{array}{|c|c|c|c|} \hline 1 \\ \hline 2 \\ \hline 3 \\ \hline 4 \\ \hline 5 \\ \hline 6 \\ \hline \end{array} \end{array} = \begin{array}{|c|c|c|c|} \hline 1 \\ \hline 2 \\ \hline 3 \\ \hline 4 \\ \hline \end{array}$$

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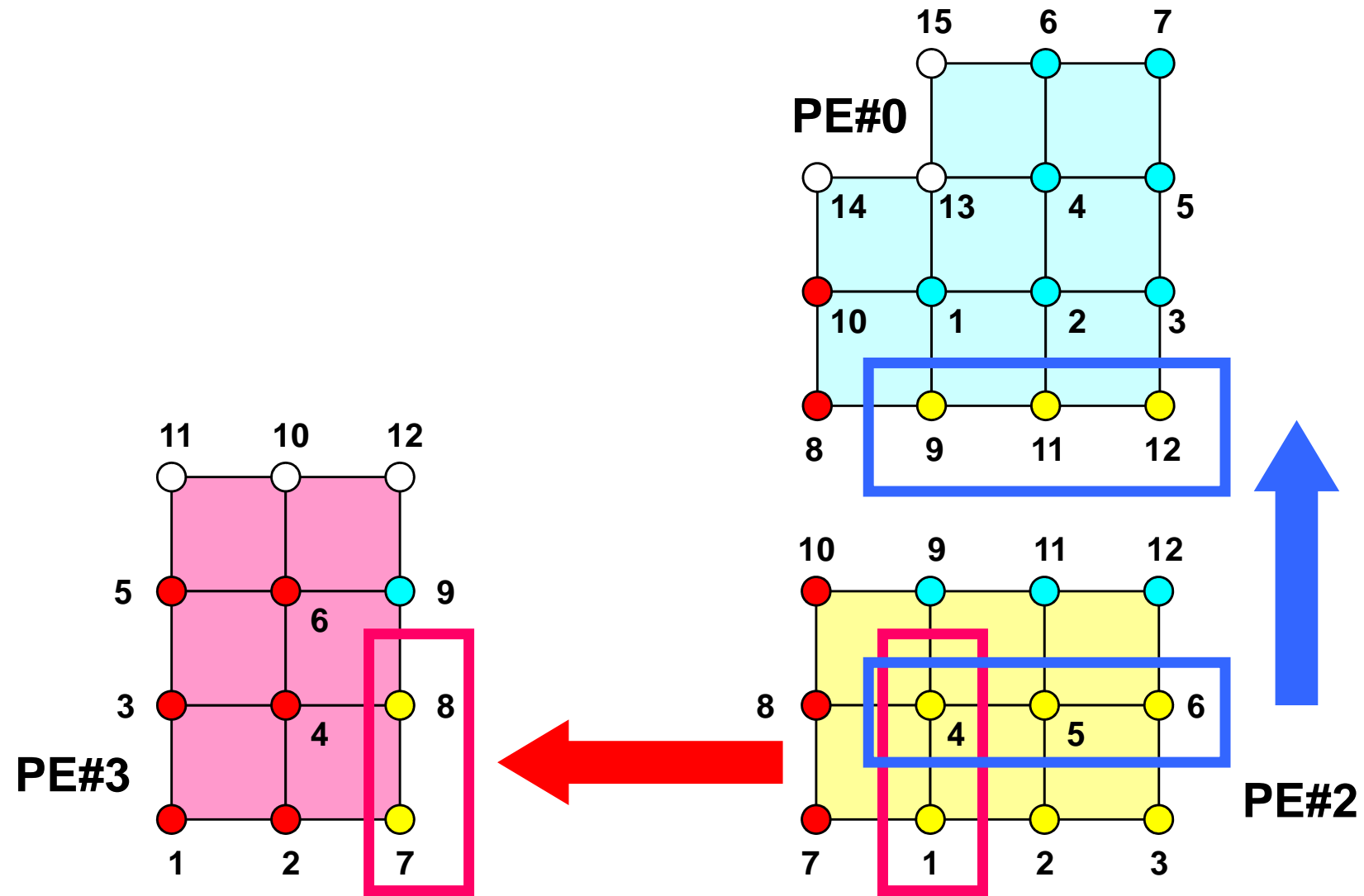
Description of Distributed Local Data



- Internal/External Points
 - Numbering: Starting from internal pts, then external pts after that
- Neighbors
 - Shares overlapped meshes
 - Number and ID of neighbors
- External Points
 - From where, how many, and which external points are received/imported ?
- Boundary Points
 - To where, how many and which boundary points are sent/exported ?

Boundary Nodes (境界点) : SEND

PE#2 : send information on “boundary nodes”



SEND: MPI_Isend/Irecv/Waitall

SENDbuf



```

do neib= 1, NEIBPETOT
  do k= export_index(neib-1)+1, export_index(neib)
    kk= export_item(k)
    SENDbuf (k) = VAL (kk)
  enddo
enddo

do neib= 1, NEIBPETOT
  iS_e= export_index(neib-1) + 1
  iE_e= export_index(neib   )
  BUFlength_e= iE_e + 1 - iS_e

  call MPI_ISEND
  &           (SENDbuf(iS_e), BUFlength_e, MPI_INTEGER, NEIBPE(neib), 0, &
  &           MPI_COMM_WORLD, request_send(neib), ierr)
enddo

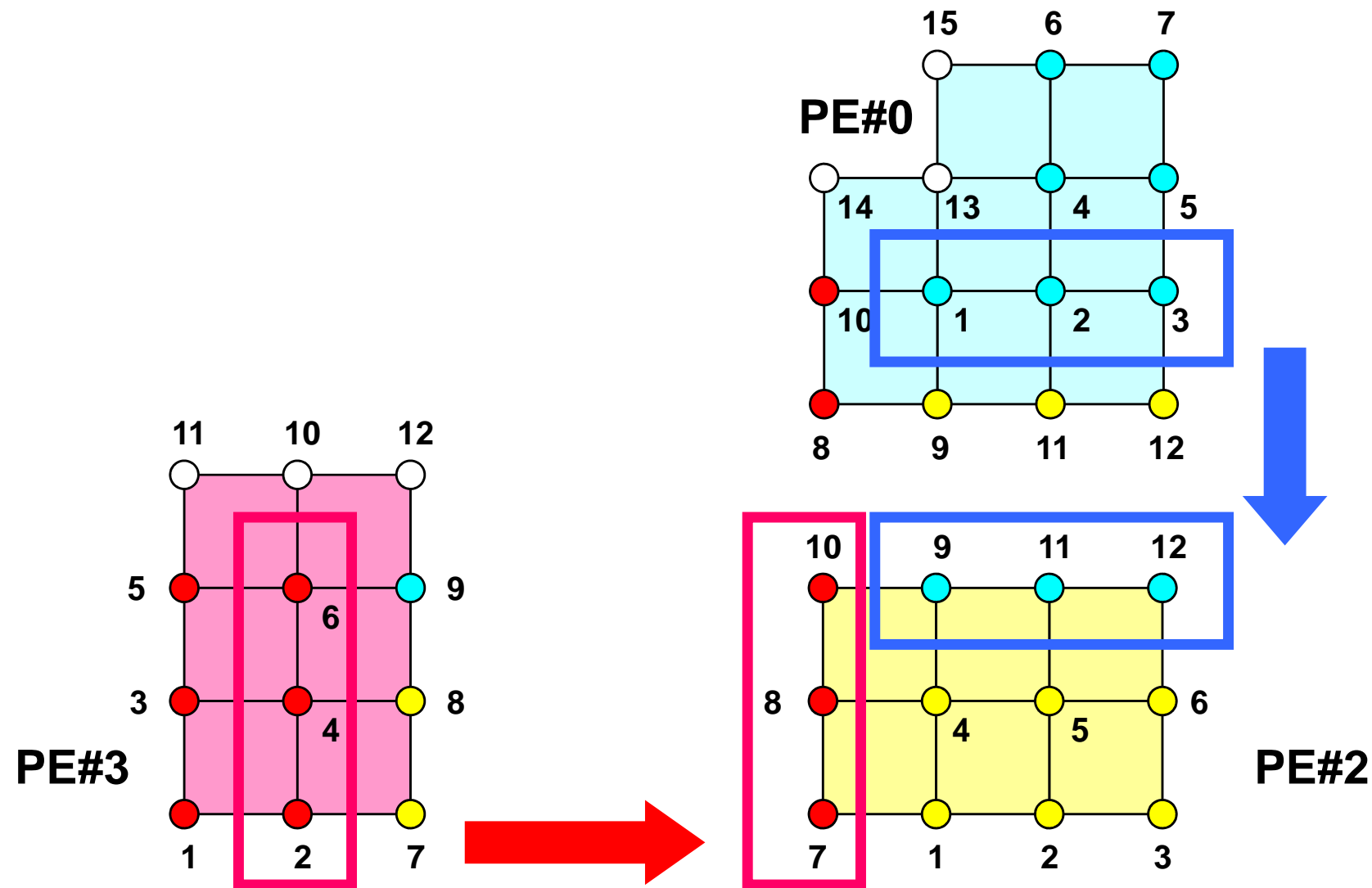
call MPI_WAITALL (NEIBPETOT, request_send, stat_recv, ierr)

```

Copied to sending buffers

External Nodes (外点) : RECEIVE

PE#2 : receive information for “external nodes”



RECV: MPI_Isend/Irecv/Waitall

```

do neib= 1, NEIBPETOT
    iS_i= import_index(neib-1) + 1
    iE_i= import_index(neib )
    BUFlength_i= iE_i + 1 - iS_i

    call MPI_IRecv
&          (RECVbuf(iS_i), BUFlength_i, MPI_INTEGER, NEIBPE(neib), 0, &
&           MPI_COMM_WORLD, request_recv(neib), ierr)
enddo

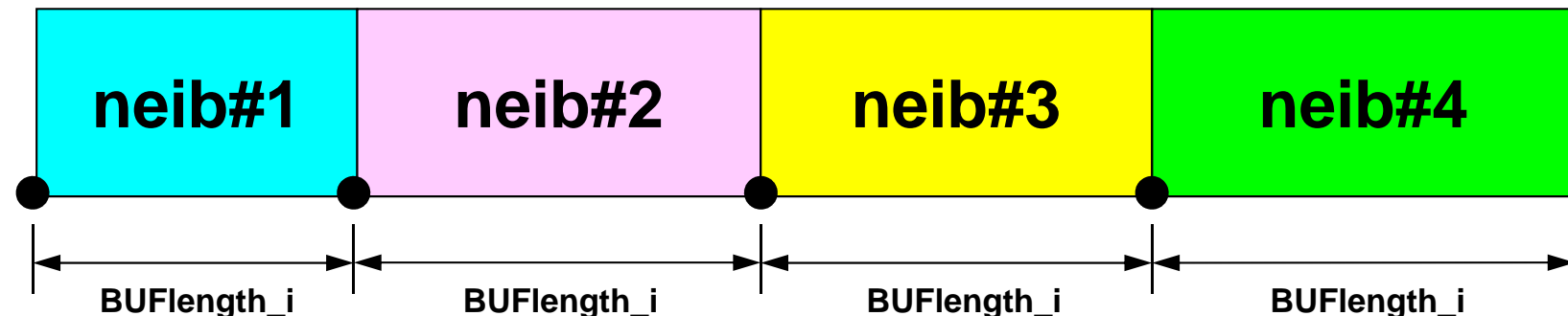
call MPI_WAITALL (NEIBPETOT, request_recv, stat_recv, ierr)

do neib= 1, NEIBPETOT
    do k= import_index(neib-1)+1, import_index(neib)
        kk= import_item(k)
        VAL(kk)= RECVbuf (k)
    enddo
enddo

```

Copied from receiving buffer

RECVbuf



`import_index(0)+1 import_index(1)+1 import_index(2)+1 import_index(3)+1 import_index(4)`

- Overview
- Distributed Local Data
- **Program**
- Results

Program: 1d.f (1/11)

Variables

```
program heat1Dp
implicit REAL*8 (A-H, O-Z)
include 'mpif.h'

integer :: N, NPLU, ITERmax
integer :: R, Z, P, Q, DD

real(kind=8) :: dX, RESID, EPS
real(kind=8) :: AREA, QV, COND
real(kind=8), dimension(:), allocatable :: PHI, RHS
real(kind=8), dimension(:,:), allocatable :: DIAG, AMAT
real(kind=8), dimension(:, :), allocatable :: W

real(kind=8), dimension(2, 2) :: KMAT, EMAT

integer, dimension(:), allocatable :: ICELNOD
integer, dimension(:), allocatable :: INDEX, ITEM
integer(kind=4) :: NEIBPETOT, BUFlength, PETOT
integer(kind=4), dimension(2) :: NEIBPE

integer(kind=4), dimension(0:2) :: import_index, export_index
integer(kind=4), dimension( 2) :: import_item , export_item

real(kind=8), dimension(2) :: SENDbuf, RECVbuf

integer(kind=4), dimension(:, :, ), allocatable :: stat_send
integer(kind=4), dimension(:, :, ), allocatable :: stat_recv
integer(kind=4), dimension(:, ), allocatable :: request_send
integer(kind=4), dimension(:, ), allocatable :: request_recv
```

Program: 1d.f (2/11)

Control Data

```

!C
!C +-----+
!C | INIT. |
!C +-----+
!C===
!C
!C-- MPI init.

    call MPI_Init      (ierr)
    call MPI_Comm_size (MPI_COMM_WORLD, PETOT, ierr )
    call MPI_Comm_rank (MPI_COMM_WORLD, my_rank, ierr )          Initialization
                                                               Entire Process #: PETOT
                                                               Rank ID (0-PETot-1): my_rank

!C
!C-- CTRL data
  if (my_rank.eq.0) thenn
    open (11, file='input.dat', status='unknown')
    read (11,*) NEg
    read (11,*) dX, QV, AREA, COND
    read (11,*) ITERmax
    read (11,*) EPS
    close (11)
  endif

  call MPI_Bcast (NEg      , 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr)
  call MPI_Bcast (ITERmax, 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr)
  call MPI_Bcast (dX       , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
  call MPI_Bcast (QV       , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
  call MPI_Bcast (AREA     , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
  call MPI_Bcast (COND     , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
  call MPI_Bcast (EPS      , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)

```

Program: 1d.f (2/11)

Control Data

```

!C
!C +-----+
!C | INIT. |
!C +-----+
!C===
!C
!C-- MPI init.

    call MPI_Init      (ierr)                      Initialization
    call MPI_Comm_size (MPI_COMM_WORLD, PETOT, ierr )  Entire Process #: PETOT
    call MPI_Comm_rank (MPI_COMM_WORLD, my_rank, ierr )  Rank ID (0-PETot-1): my_rank

!C
!C-- CTRL data
    if (my_rank.eq.0) then
        open (11, file='input.dat', status='unknown')
        read (11,*) Neg
        read (11,*) dX, QV, AREA, COND
        read (11,*) ITERmax
        read (11,*) EPS
        close (11)
    endif

    call MPI_Bcast (NEg      , 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr)
    call MPI_Bcast (ITERmax, 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr)
    call MPI_Bcast (dX       , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
    call MPI_Bcast (QV       , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
    call MPI_Bcast (AREA     , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
    call MPI_Bcast (COND     , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
    call MPI_Bcast (EPS      , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)


```

Reading control file if my_rank=0

Neg: Global Number of Elements

Program: 1d.f (2/11)

Control Data

```

!C
!C +-----+
!C | INIT. |
!C +-----+
!C===
!C
!C-- MPI init.

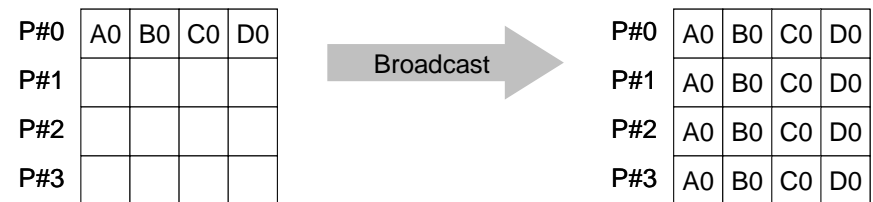
call MPI_Init      (ierr)
call MPI_Comm_size (MPI_COMM_WORLD, PETOT, ierr )
call MPI_Comm_rank (MPI_COMM_WORLD, my_rank, ierr )           Initialization
                                                               Entire Process #: PETOT
                                                               Rank ID (0-PETot-1): my_rank

!C
!C-- CTRL data
if (my_rank.eq.0) then
  open (11, file='input.dat', status='unknown')
  read (11,*) Neg
  read (11,*) dX, QV, AREA, COND
  read (11,*) ITERmax
  read (11,*) EPS
  close (11)
endif                                         Reading control file if my_rank=0
                                                               Neg: Global Number of Elements

call MPI_Bcast (NEg      , 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr) Parameters are sent to each proces
call MPI_Bcast (ITERmax, 1, MPI_INTEGER, 0, MPI_COMM_WORLD, ierr) from Process #0.
call MPI_Bcast (dX       , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
call MPI_Bcast (QV       , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
call MPI_Bcast (AREA     , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
call MPI_Bcast (COND     , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)
call MPI_Bcast (EPS      , 1, MPI_DOUBLE_PRECISION, 0, MPI_COMM_WORLD, ierr)

```

MPI_BCAST



- Broadcasts a message from the process with rank "root" to all other processes of the communicator
- **call MPI_BCAST (buffer, count, datatype, root, comm, ierr)**
 - **buffer** choice I/O starting address of buffer
type is defined by "datatype"
 - **count** I I number of elements in send/recv buffer
 - **datatype** I I data type of elements of send/recv buffer
FORTRAN: MPI_INTEGER, MPI_REAL, MPI_DOUBLE_PRECISION, MPI_CHARACTER etc.
C: MPI_INT, MPI_FLOAT, MPI_DOUBLE, MPI_CHAR etc.
 - **root** I I **rank of root process**
 - **comm** I I communicator
 - **ierr** I O completion code

Program: 1d.f (3/11)

Distributed Local Mesh

```
!C
!C-- Local Mesh Size
```

```
Ng= NEg + 1
N = Ng / PETOT
```

Global Number of Nodes
Local Number of Nodes

```
nr = Ng - N*PETOT
if (my_rank. lt. nr) N= N+1
```

mod(Ng, PETOT) .ne. 0

```
NE= N - 1 + 2
NP= N + 2
```

```
if (my_rank. eq. 0) NE= N - 1 + 1
if (my_rank. eq. 0) NP= N + 1
```

```
if (my_rank. eq. PETOT-1) NE= N - 1 + 1
if (my_rank. eq. PETOT-1) NP= N + 1
```

```
if (PETOT. eq. 1) NE= N-1
if (PETOT. eq. 1) NP= N
```

```
!C
!C- ARRAYS
```

```
allocate (PHI(NP), DIAG(NP), AMAT(2*NP-2), RHS(NP))
allocate (ICELNOD(2*NE))
allocate (INDEX(0:NP), ITEM(2*NP-2), W(NP, 4))
PHI= 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0
```

Program: 1d.f (3/11)

Distributed Local Mesh, Uniform Elements

```
!C
!C-- Local Mesh Size
```

```
Ng= NEg + 1
N = Ng / PETOT
```

Global Number of Nodes
Local Number of Nodes

```
nr = Ng - N*PETOT
if (my_rank. lt. nr) N= N+1
```

mod(Ng, PETOT) . ne. 0

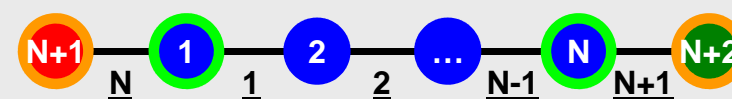
```
NE= N - 1 + 2
NP= N + 2
```

Number of Elements (Local)
Total Number of Nodes (Local) (Internal + External Nodes)

```
if (my_rank. eq. 0) NE= N - 1 + 1
if (my_rank. eq. 0) NP= N + 1
```

```
if (my_rank. eq. PETOT-1) NE= N - 1 + 1
if (my_rank. eq. PETOT-1) NP= N + 1
```

```
if (PETOT. eq. 1) NE= N-1
if (PETOT. eq. 1) NP= N
```



Others (General):
N+2 nodes
N+1 elements

```
!C
!C- ARRAYS
```

```
allocate (PHI(NP), DIAG(NP), AMAT(2*NP-2), RHS(NP))
allocate (ICELNOD(2*NE))
allocate (INDEX(0:NP), ITEM(2*NP-2), W(NP, 4))
PHI= 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0
```

Program: 1d.f (3/11)

Distributed Local Mesh, Uniform Elements

```
!C
!C-- Local Mesh Size
```

```
Ng= NEg + 1
N = Ng / PETOT
```

Global Number of Nodes
Local Number of Nodes

```
nr = Ng - N*PETOT
if (my_rank. lt. nr) N= N+1
```

mod(Ng, PETOT) . ne. 0

```
NE= N - 1 + 2
NP= N + 2
```

Number of Elements (Local)
Total Number of Nodes (Local) (Internal + External Nodes)

```
if (my_rank. eq. 0) NE= N - 1 + 1
if (my_rank. eq. 0) NP= N + 1
```

```
if (my_rank. eq. PETOT-1) NE= N - 1 + 1
if (my_rank. eq. PETOT-1) NP= N + 1
```

```
if (PETOT. eq. 1) NE= N-1
if (PETOT. eq. 1) NP= N
```



#0:
N+1 nodes
N elements

```
!C
!C- ARRAYS
```

```
allocate (PHI(NP), DIAG(NP), AMAT(2*NP-2), RHS(NP))
allocate (ICELNOD(2*NE))
allocate (INDEX(0:NP), ITEM(2*NP-2), W(NP, 4))
PHI= 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0
```

Program: 1d.f (3/11)

Distributed Local Mesh, Uniform Elem

!C
!C-- Local Mesh Size

$Ng = NEg + 1$
 $N = Ng / PETOT$

Global Number of Nodes
Local Number of Nodes

$nr = Ng - N * PETOT$
if (my_rank. lt. nr) $N = N + 1$

mod(Ng, PETOT) . ne. 0

$NE = N - 1 + 2$
 $NP = N + 2$

Number of Elements (Local)
Total Number of Nodes (Local) (Internal + External Nodes)

if (my_rank. eq. 0) $NE = N - 1 + 1$
if (my_rank. eq. 0) $NP = N + 1$

if (my_rank. eq. PETOT-1) $NE = N - 1 + 1$
if (my_rank. eq. PETOT-1) $NP = N + 1$

if (PETOT. eq. 1) $NE = N - 1$
if (PETOT. eq. 1) $NP = N$



#PETot-1:
N+1 nodes
N elements

!C
!C- ARRAYS

```
allocate (PHI (NP), DIAG (NP), AMAT (2*NP-2), RHS (NP))
allocate (ICELNOD (2*NE))
allocate (INDEX (0:NP), ITEM (2*NP-2), W (NP, 4))
PHI= 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0
```

Program: 1d.f (3/11)

Distributed Local Mesh, Uniform Elements

```

!C
!C-- Local Mesh Size

Ng= NEg + 1                                Global Number of Nodes
N = Ng / PETOT                               Local Number of Nodes

nr = Ng - N*PETOT                            mod(Ng, PETOT) . ne. 0
if (my_rank. lt. nr) N= N+1

NE= N - 1 + 2                                Number of Elements (Local)
NP= N + 2                                    Total Number of Nodes (Local) (Internal + External Nodes)

if (my_rank. eq. 0) NE= N - 1 + 1
if (my_rank. eq. 0) NP= N + 1

if (my_rank. eq. PETOT-1) NE= N - 1 + 1
if (my_rank. eq. PETOT-1) NP= N + 1

if (PETOT. eq. 1) NE= N-1
if (PETOT. eq. 1) NP= N

```

```

!C
!C- ARRAYS

```

```

allocate (PHI(NP), DIAG(NP), AMAT(2*NP-2), RHS(NP))      Size of arrays is "NP" , not "N"
allocate (ICELNOD(2*NE))
allocate (INDEX(0:NP), ITEM(2*NP-2), W(NP, 4))
PHI= 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0

```

Program: 1d.f (4/11)

Initialization of Arrays, Elements-Nodes

```

do icel= 1, NE
  ICELNOD(2*icel-1)= icel
  ICELNOD(2*icel    )= icel + 1
enddo

if (PETOT.gt. 1) then

  if (my_rank.eq. 0) then
    icel= NE
    ICELNOD(2*icel-1)= N
    ICELNOD(2*icel    )= N + 1

  else if (my_rank.eq. PETOT-1) then
    icel= NE
    ICELNOD(2*icel-1)= N + 1
    ICELNOD(2*icel    )= 1

  else
    icel= NE - 1
    ICELNOD(2*icel-1)= N + 1
    ICELNOD(2*icel    )= 1
    icel= NE
    ICELNOD(2*icel-1)= N
    ICELNOD(2*icel    )= N + 2

  endif
endif

```



*Icelnod(2*icel-1)*
 $=icel$

*Icelnod(2*icel)*
 $=icel+1$

Program: 1d.f (4/11)

Initialization of Arrays, Elements-Nodes

```
do icel= 1, NE
  ICELNOD(2*icel-1)= icel
  ICELNOD(2*icel    )= icel + 1
enddo
```

```
if (PETOT.gt. 1) then
```

```
if (my_rank.eq. 0) then
```

```
  icel= NE
```

```
  ICELNOD(2*icel-1)= N
```

```
  ICELNOD(2*icel    )= N + 1
```

e.g. Element-1 includes node-1 and node-2



#0:
N+1 nodes
N elements

```
else if (my_rank.eq. PETOT-1) then
```

```
  icel= NE
```

```
  ICELNOD(2*icel-1)= N + 1
```

```
  ICELNOD(2*icel    )= 1
```



#PETot-1:
N+1 nodes
N elements

```
else
```

```
  icel= NE - 1
```

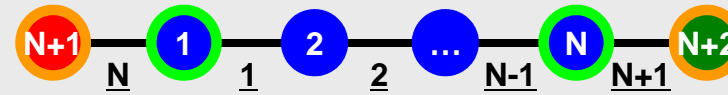
```
  ICELNOD(2*icel-1)= N + 1
```

```
  ICELNOD(2*icel    )= 1
```

```
  icel= NE
```

```
  ICELNOD(2*icel-1)= N
```

```
  ICELNOD(2*icel    )= N + 2
```



Others (General):
N+2 nodes
N+1 elements

```
endif
```

```
endif
```

Program: 1d.f (5/11)

"Index"

```
KMAT(1, 1)= +1. d0
KMAT(1, 2)= -1. d0
KMAT(2, 1)= -1. d0
KMAT(2, 2)= +1. d0
```

!C==

!C
!C +-----+
!C | CONNECTIVITY |
!C +-----+
!C==

INDEX = 2

INDEX(0)= 0

INDEX(N+1)= 1

INDEX(NP)= 1

if (my_rank. eq. 0) INDEX(1)= 1
if (my_rank. eq. PETOT-1) INDEX(N)= 1

do i= 1, NP
INDEX(i)= INDEX(i) + INDEX(i-1)
enddo

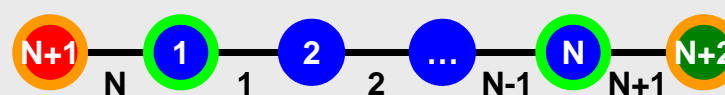
NPLU= INDEX(NP)
ITEM= 0



#0:
N+1 nodes
N elements



#PETot-1:
N+1 nodes
N elements



Others (General):
N+2 nodes
N+1 elements

Program: 1d.f (6/11)

"Item"

```

do i= 1, N
  jS= INDEX(i-1)
  if (my_rank.eq.0.and.i.eq.1) then
    ITEM(jS+1)= i+1
  else if (my_rank.eq.PETOT-1.and.i.eq.N) then
    ITEM(jS+1)= i-1
  else
    ITEM(jS+1)= i-1
    ITEM(jS+2)= i+1
    if (i.eq.1) ITEM(jS+1)= N + 1
    if (i.eq.N) ITEM(jS+2)= N + 2
    if (my_rank.eq.0.and.i.eq.N) ITEM(jS+2)= N + 1
  endif
enddo

```

```

i = N + 1
jS= INDEX(i-1)
if (my_rank.eq.0) then
  ITEM(jS+1)= N
else
  ITEM(jS+1)= 1
endif

i = N + 2
if (my_rank.ne.0.and.my_rank.ne.PETOT-1) then
  jS= INDEX(i-1)
  ITEM(jS+1)= N
endif

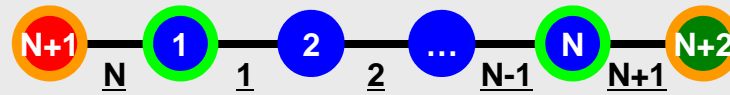
```



#0:
N+1 nodes
N elements



#PETot-1:
N+1 nodes
N elements



Others (General):
N+2 nodes
N+1 elements

Program: 1d.f (7/11)

Communication Tables

```

!C
!C-- COMMUNICATION
NEIBPETOT= 2
if (my_rank.eq.0) NEIBPETOT= 1
if (my_rank.eq.PETOT-1) NEIBPETOT= 1
if (PETOT.eq.1) NEIBPETOT= 0

NEIBPE(1)= my_rank - 1
NEIBPE(2)= my_rank + 1

if (my_rank.eq.0) NEIBPE(1)= my_rank + 1
if (my_rank.eq.PETOT-1) NEIBPE(1)= my_rank - 1

BUFlength= 1

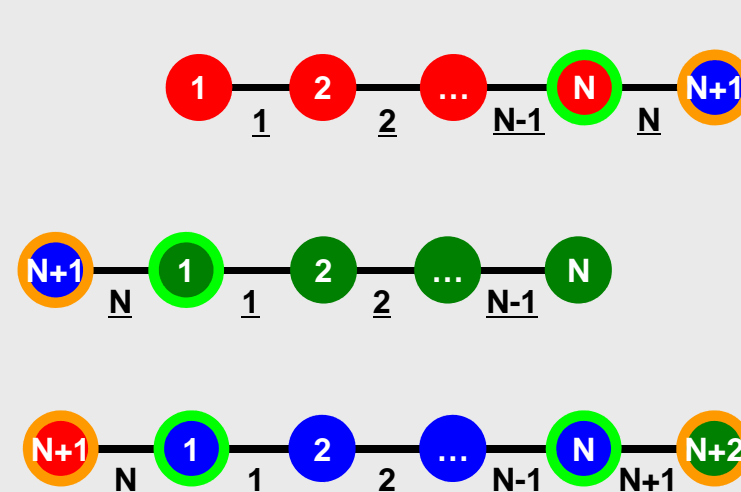
import_index(1)= 1
import_index(2)= 2
import_item (1)= N+1
import_item (2)= N+2

export_index(1)= 1
export_index(2)= 2
export_item (1)= 1
export_item (2)= N

if (my_rank.eq.0) then
  import_item (1)= N+1
  export_item (1)= N
endif

!C
!C-- INIT. arrays for MPI_Waitall
allocate (stat_send(MPI_STATUS_SIZE,NEIBPETOT), stat_recv(MPI_STATUS_SIZE,NEIBPETOT))
allocate (request_send(NEIBPETOT), request_recv(NEIBPETOT))

```



#0:
N+1 nodes
N elements

#PETot-1:
N+1 nodes
N elements

Others (General):
N+2 nodes
N+1 elements

MPI_ISEND

- Begins a non-blocking send
 - Send the contents of sending buffer (starting from **sendbuf**, number of messages: **count**) to **dest** with **tag**.
 - Contents of sending buffer cannot be modified before calling corresponding **MPI_Waitall**.
- **call MPI_ISEND**
(sendbuf, count, datatype, dest, tag, comm, request, ierr)

– sendbuf	choice	I	starting address of sending buffer
– count	I	I	number of elements sent to each process
– datatype	I	I	data type of elements of sending buffer
– dest	I	I	rank of destination
– tag	I	I	message tag This integer can be used by the application to distinguish messages. Communication occurs if tag's of MPI_Isend and MPI_Irecv are matched. Usually tag is set to be "0" (in this class),
– comm	I	I	communicator
– request	I	O	communication request array used in MPI_Waitall
– ierr	I	O	completion code

MPI_IRecv

- Begins a non-blocking receive
 - Receiving the contents of receiving buffer (starting from **recvbuf**, number of messages: **count**) from **source** with **tag** .
 - Contents of receiving buffer cannot be used before calling corresponding **MPI_Waitall**.
- **call MPI_IRecv**
(recvbuf, count, datatype, dest, tag, comm, request, ierr)

– recvbuf	choice	I	starting address of receiving buffer
– count	I	I	number of elements in receiving buffer
– datatype	I	I	data type of elements of receiving buffer
– source	I	I	rank of source
– tag	I	I	message tag This integer can be used by the application to distinguish messages. Communication occurs if tag's of MPI_Isend and MPI_Irecv are matched. Usually tag is set to be "0" (in this class),
– comm	I	I	communicator
– request	I	O	communication request used in MPI_Waitall
– ierr	I	O	completion code

MPI_WAITALL

- **`MPI_Waitall`** blocks until all comm's, associated with request in the array, complete. It is used for synchronizing **`MPI_Isend`** and **`MPI_Irecv`** in this class.
- At sending phase, contents of sending buffer cannot be modified before calling corresponding **`MPI_Waitall`**. At receiving phase, contents of receiving buffer cannot be used before calling corresponding **`MPI_Waitall`**.
- **`MPI_Isend`** and **`MPI_Irecv`** can be synchronized simultaneously with a single **`MPI_Waitall`** if it is consistent.
 - Same request should be used in **`MPI_Isend`** and **`MPI_Irecv`**.
- Its operation is similar to that of **`MPI_Barrier`** but, **`MPI_Waitall`** can not be replaced by **`MPI_Barrier`**.
 - Possible troubles using **`MPI_Barrier`** instead of **`MPI_Waitall`**: Contents of request and status are not updated properly, very slow operations etc.
- **call MPI_WAITALL (count, request, status, ierr)**
 - count I I number of processes to be synchronized
 - request I I/O comm. request used in `MPI_Waitall` (array size: count)
 - status I O array of status objects
MPI_STATUS_SIZE: defined in 'mpif.h', 'mpi.h'
 - ierr I O completion code

Generalized Comm. Table: Send

- Neighbors
 - NEIBPETOT, NEIBPE(neib)
- Message size for each neighbor
 - export_index(neib), neib= 0, NEIBPETOT
- ID of boundary points
 - export_item(k), k= 1, export_index(NEIBPETOT)
- Messages to each neighbor
 - SENDbuf(k), k= 1, export_index(NEIBPETOT)

SEND: MPI_ISEND/IRecv/WAITALL

SENDbuf



```

do neib= 1, NEIBPETOT
  do k= export_index(neib-1)+1, export_index(neib)
    kk= export_item(k)
    SENDbuf (k) = VAL (kk)
  enddo
enddo

do neib= 1, NEIBPETOT
  iS_e= export_index(neib-1) + 1
  iE_e= export_index(neib )
  BUFlength_e= iE_e + 1 - iS_e

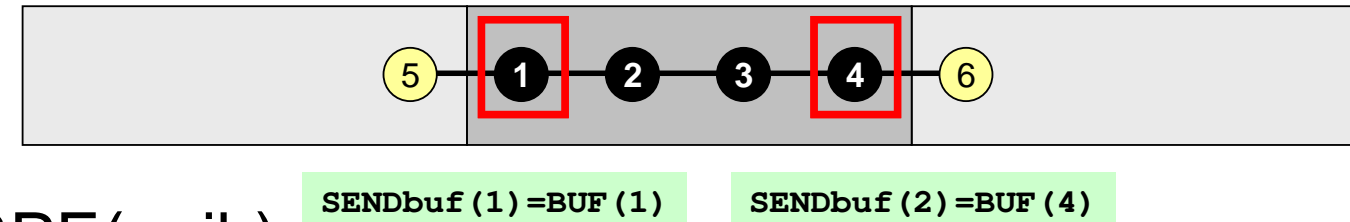
  call MPI_ISEND
  &           (SENDbuf(iS_e), BUFlength_e, MPI_INTEGER, NEIBPE(neib), 0, &
  &           MPI_COMM_WORLD, request_send(neib), ierr)
enddo

call MPI_WAITALL (NEIBPETOT, request_send, stat_recv, ierr)

```

Copied to sending buffers

SEND/Export: 1D Problem



- Neighbors
 - NEIBPETOT, NEIBPE(neib)
 - NEIBPETOT=2, NEIB(1)= my_rank-1, NEIB(2)= my_rank+1
- Message size for each neighbor
 - export_index(neib), neib= 0, NEIBPETOT
 - export_index(0)=0, export_index(1)= 1, export_index(2)= 2
- ID of **boundary** points
 - export_item(k), k= 1, export_index(NEIBPETOT)
 - export_item(1)= 1, export_item(2)= N
- Messages to each neighbor
 - SENDbuf(k), k= 1, export_index(NEIBPETOT)
 - SENDbuf(1)= BUF(1), SENDbuf(2)= BUF(N)

Generalized Comm. Table: Receive

- Neighbors
 - NEIBPETOT, NEIBPE(neib)
- Message size for each neighbor
 - import_index(neib), neib= 0, NEIBPETOT
- ID of external points
 - import_item(k), k= 1, import_index(NEIBPETOT)
- Messages from each neighbor
 - RECVbuf(k), k= 1, import_index(NEIBPETOT)

RECV: MPI_Isend/Irecv/Waitall

```

do neib= 1, NEIBPETOT
    iS_i= import_index(neib-1) + 1
    iE_i= import_index(neib  )
    BUFlength_i= iE_i + 1 - iS_i

    call MPI_IRecv
&          (RECVbuf(iS_i), BUFlength_i, MPI_INTEGER, NEIBPE(neib), 0,&
&           MPI_COMM_WORLD, request_recv(neib), ierr)
    enddo

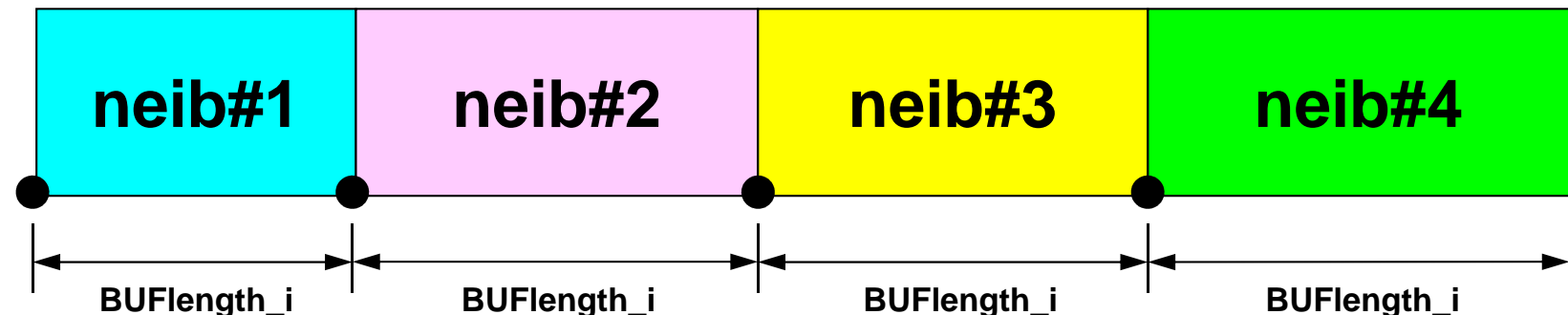
    call MPI_WAITALL (NEIBPETOT, request_recv, stat_recv, ierr)

do neib= 1, NEIBPETOT
    do k= import_index(neib-1)+1, import_index(neib)
        kk= import_item(k)
        VAL(kk)= RECVbuf (k)
    enddo
enddo

```

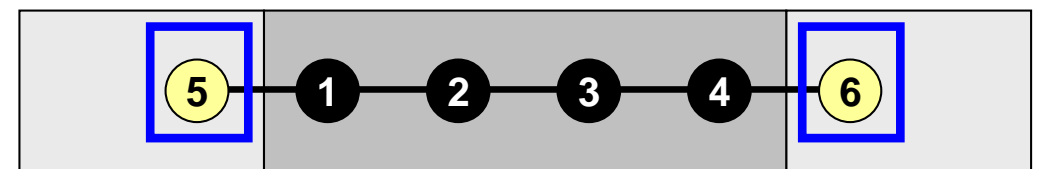
Copied from receiving buffer

RECVbuf



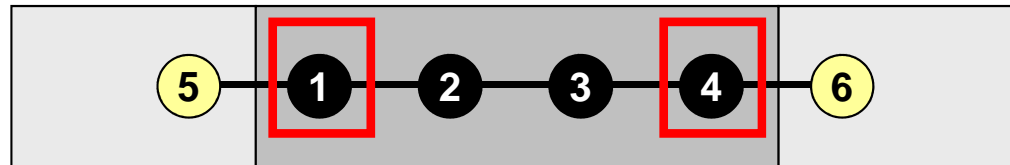
`import_index(0)+1 import_index(1)+1 import_index(2)+1 import_index(3)+1 import_index(4)`

RECV/Import: 1D Proble



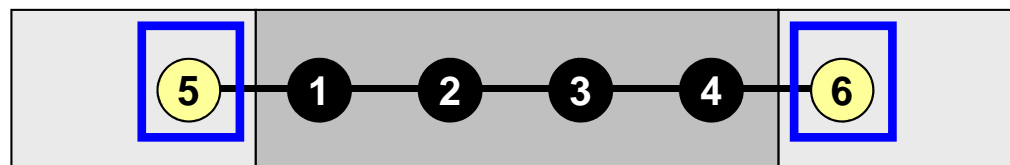
- Neighbors
 - NEIBPETOT, NEIBPE(neib)
 - NEIBPETOT=2, NEIB(1)= my_rank-1, NEIB(2)= my_rank+1
- Message size for each neighbor
 - import_index(neib), neib= 0, NEIBPETOT
 - import_index(0)=0, import_index(1)= 1, import_index(2)= 2
- ID of **external** points
 - import_item(k), k= 1, import_index(NEIBPETOT)
 - import_item(1)= N+1, import_item(2)= N+2
- Messages from each neighbor
 - RECVbuf(k), k= 1, import_index(NEIBPETOT)
 - BUF(N+1)=RECVbuf(1), BUF(N+2)=RECVbuf(2)

Generalized Comm. Table: Fortran



`SENDbuf(1)=BUF(1)`

`SENDbuf(2)=BUF(4)`



`BUF(5)=RECVbuf(1)`

`BUF(6)=RECVbuf(2)`

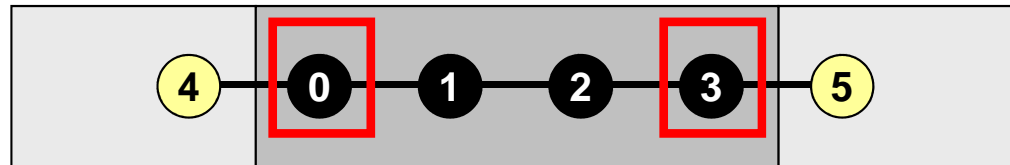
```
NEIBPETOT= 2
NEIBPE(1)= my_rank - 1
NEIBPE(2)= my_rank + 1
```

```
import_index(1)= 1
import_index(2)= 2
import_item (1)= N+1
import_item (2)= N+2
```

```
export_index(1)= 1
export_index(2)= 2
export_item (1)= 1
export_item (2)= N
```

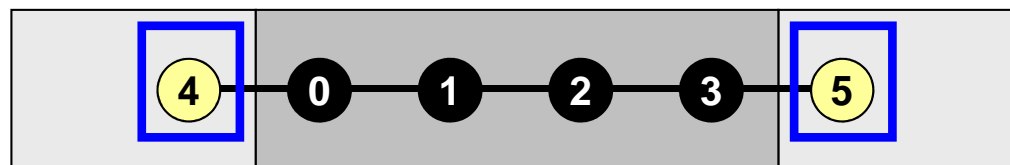
```
if (my_rank.eq.0) then
    import_item (1)= N+1
    export_item (1)= N
    NEIBPE(1)= my_rank+1
endif
```

Generalized Comm. Table: C



`SENDbuf[0]=BUF[0]`

`SENDbuf[1]=BUF[3]`



`BUF[4]=RECVbuf[0]`

`BUF[5]=RECVbuf[1]`

```
NEIBPETOT= 2
NEIBPE[0]= my_rank - 1
NEIBPE[1]= my_rank + 1
```

```
import_index[1]= 0
import_index[2]= 1
import_item [0]= N
import_item [1]= N+1
```

```
export_index[1]= 0
export_index[2]= 1
export_item [0]= 0
export_item [1]= N-1
```

```
if (my_rank.eq.0) then
  import_item [0]= N
  export_item [0]= N-1
  NEIBPE[0]= my_rank+1
endif
```

Program: 1d.f (8/11)

Matrix Assembling, NO changes from 1-CPU co

```
!C
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C==
```

```
do icel= 1, NE
  in1= ICELNOD(2*icel-1)
  in2= ICELNOD(2*icel )
  DL = dx
  cK= AREA*COND/DL
  EMAT (1, 1)= Ck*KMAT (1, 1)
  EMAT (1, 2)= Ck*KMAT (1, 2)
  EMAT (2, 1)= Ck*KMAT (2, 1)
  EMAT (2, 2)= Ck*KMAT (2, 2)

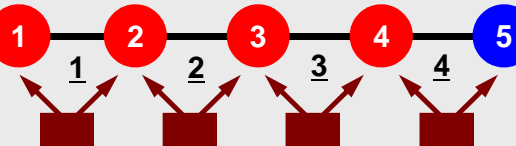
  DIAG(in1)= DIAG(in1) + EMAT (1, 1)
  DIAG(in2)= DIAG(in2) + EMAT (2, 2)

  if (my_rank.eq.0. and. icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1

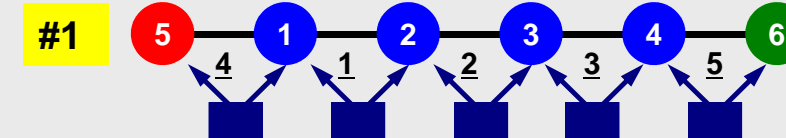
  AMAT (k1)= AMAT (k1) + EMAT (1, 2)
  AMAT (k2)= AMAT (k2) + EMAT (2, 1)

  QN= 0.50d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
enddo
```

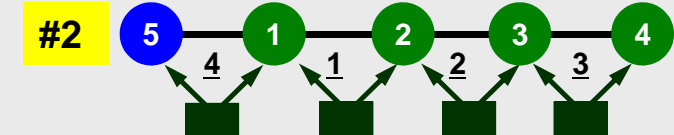
#0



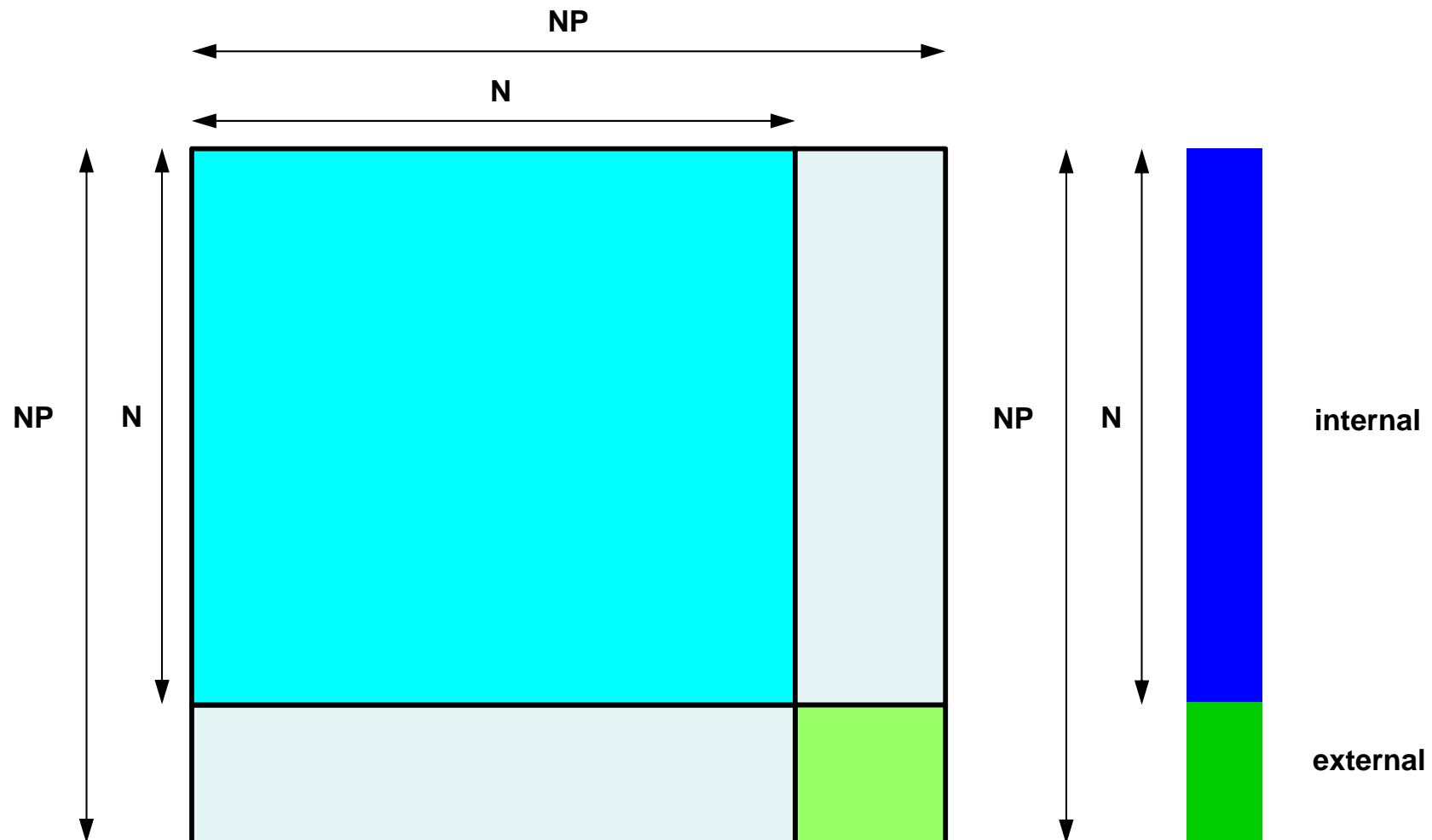
#1



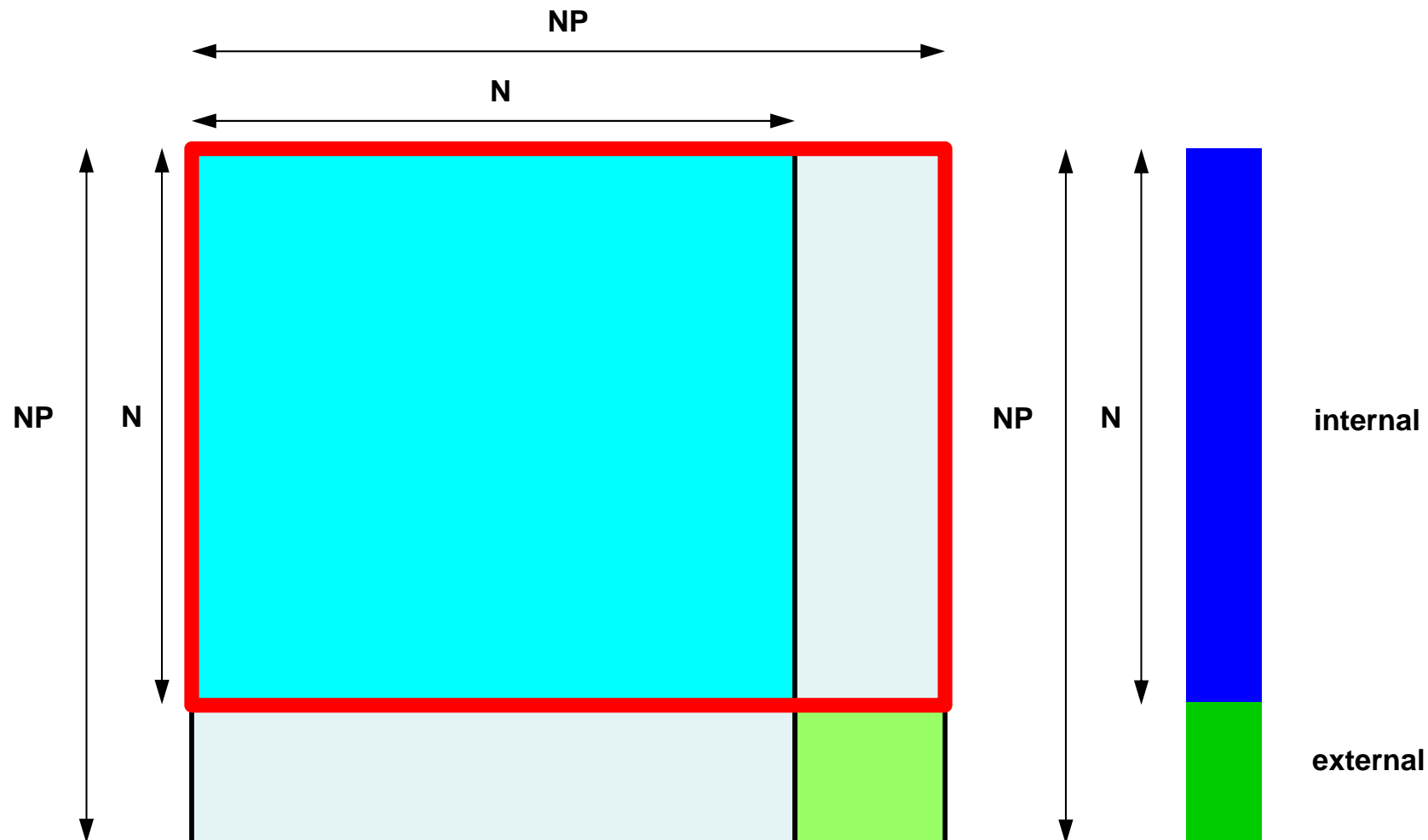
#2



Local Matrix



We really need these parts:



MAT_ASS_MAIN: Overview

```

do kpn= 1, 2      Gaussian Quad. points in  $\zeta$ -direction
  do jpn= 1, 2    Gaussian Quad. points in  $\eta$ -direction
    do ipn= 1, 2   Gaussian Quad. Pointe in  $\xi$ -direction
      Define Shape Function at Gaussian Quad. Points (8-points)
      Its derivative on natural/local coordinate is also defined.
    enddo
  enddo
enddo

```

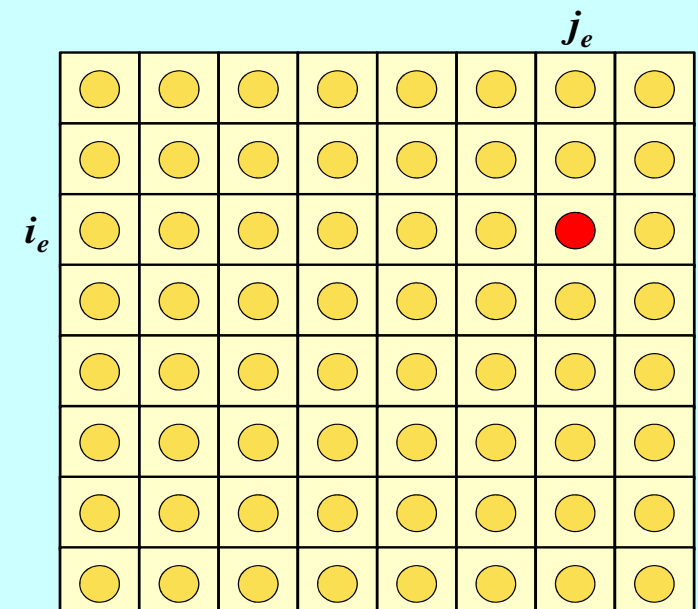
do icel= 1, ICELTOT Loop for Element
 Jacobian and derivative on global coordinate of shape functions at
 Gaussian Quad. Points are defined according to coordinates of 8 nodes. (JACOBI)

```

do ie= 1, 8        Local Node ID
  do je= 1, 8        Local Node ID
    Global Node ID: ip, jp
    Address of  $A_{ip, jp}$  in "item" : kk

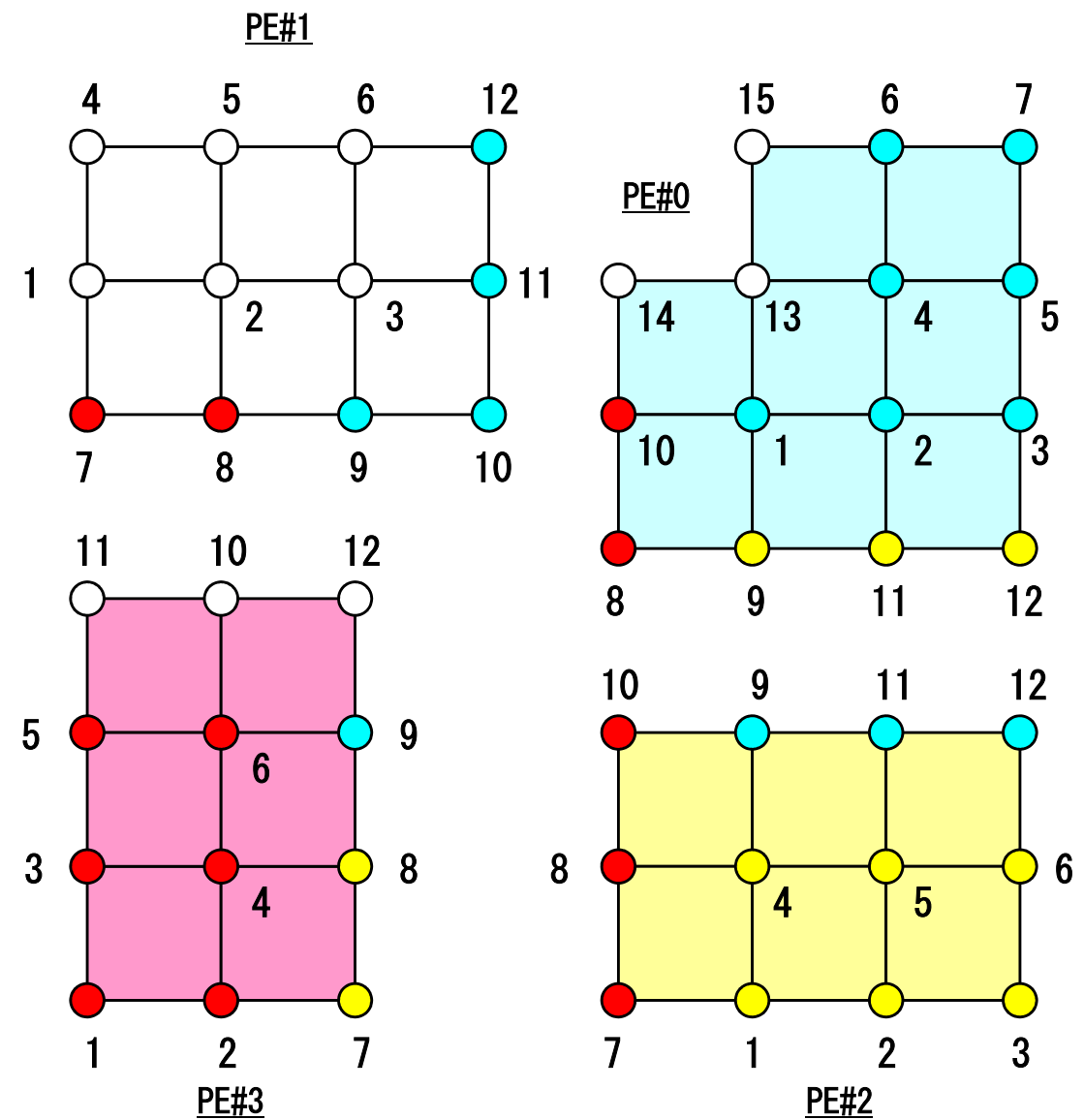
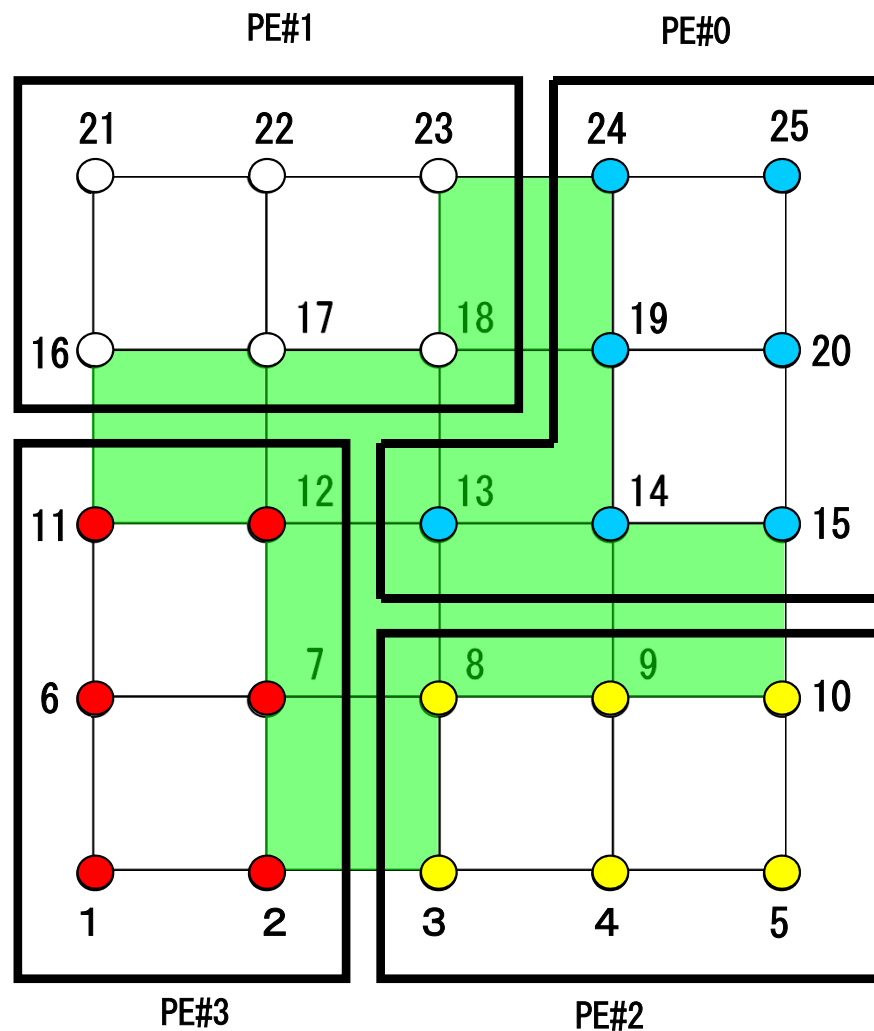
    do kpn= 1, 2      Gaussian Quad. points in  $\zeta$ -direction
      do jpn= 1, 2    Gaussian Quad. points in  $\eta$ -direction
        do ipn= 1, 2   Gaussian Quad. points in  $\xi$ -direction
          integration on each element
          coefficients of element matrices
          accumulation to global matrix
        enddo
      enddo
    enddo
  enddo
enddo
enddo

```

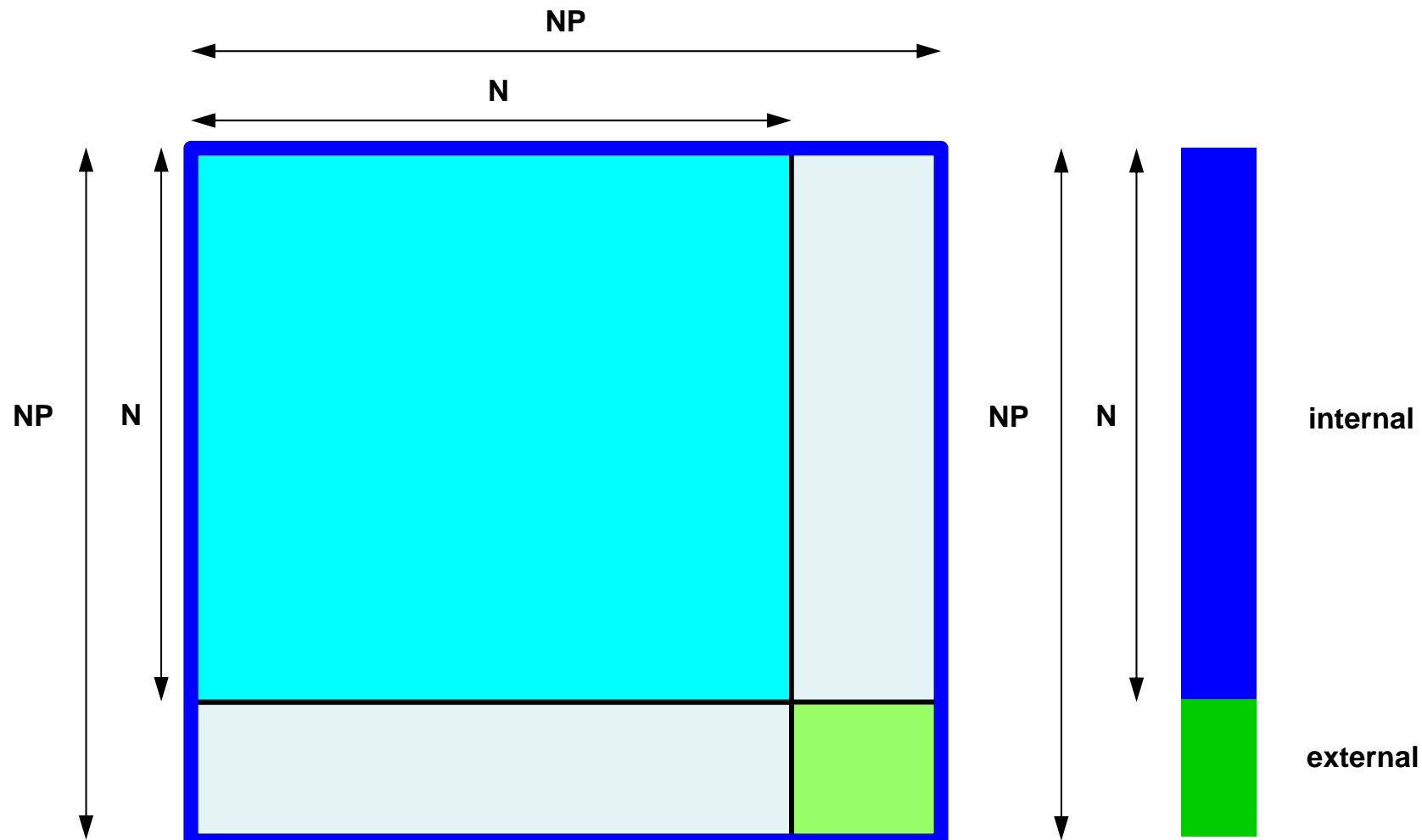


MAT_ASS_MAIN visits all elements

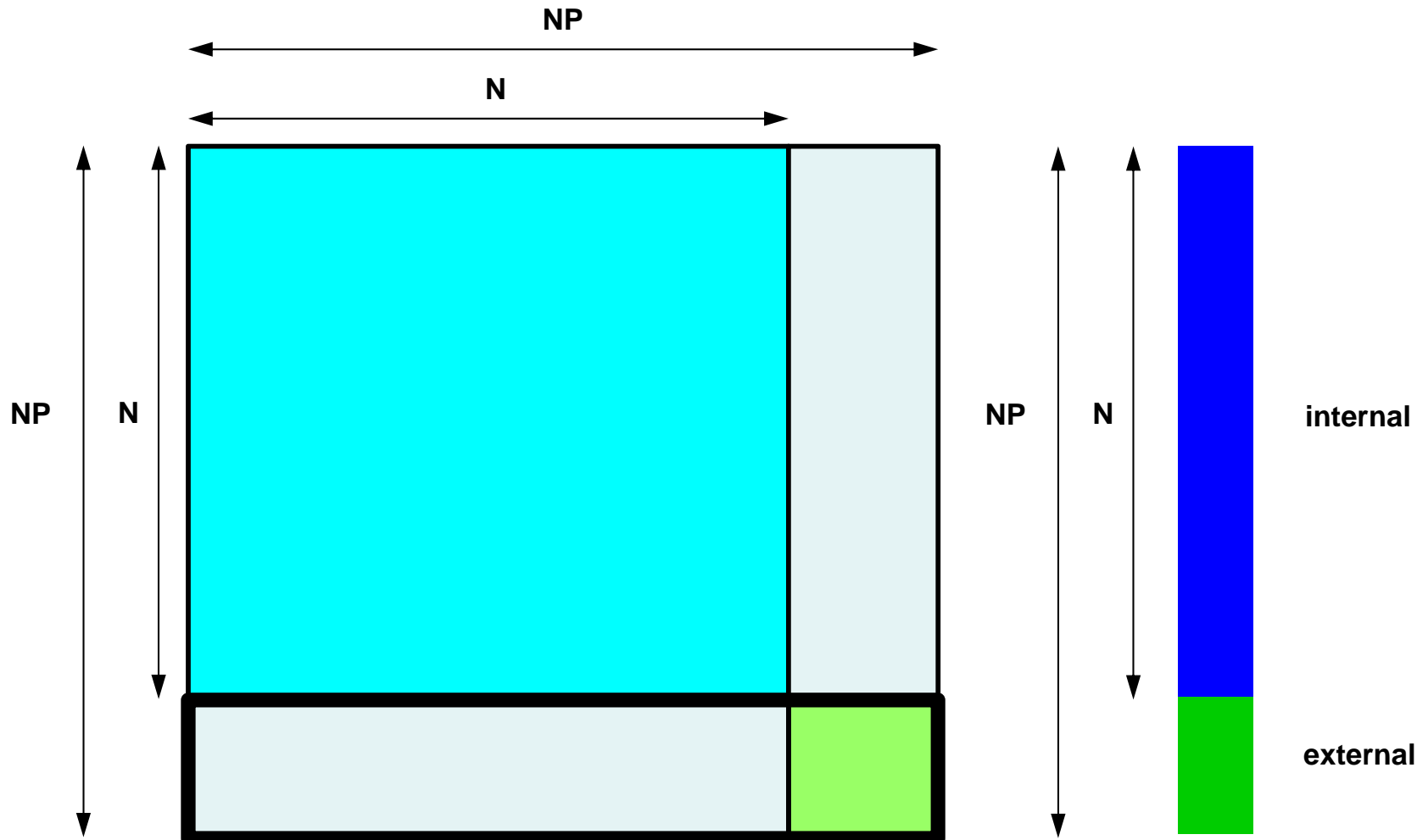
including overlapped elements with external nodes



Therefore, we have this matrix



But components of this part are not complete, and not used in computation



Program: 1d.f (9/11)

Boundary Cond., ALMOST NO changes from 1-CPU code

```
!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C==
```

#0



!C

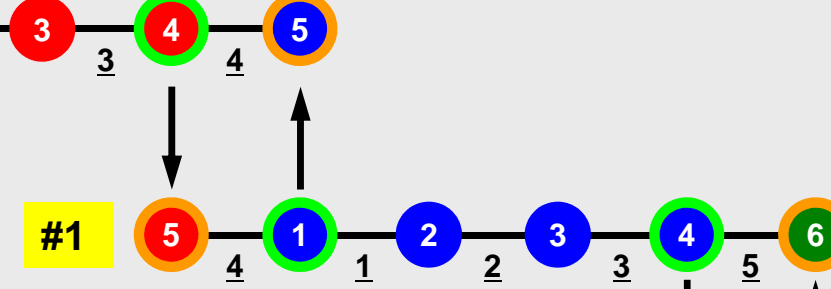
!C-- X=Xmin

```
if (my_rank.eq.0) then
  i = 1
  jS= INDEX(i-1)

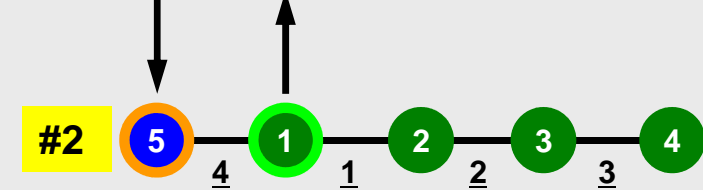
  AMAT(jS+1)= 0. d0
  DIAG(i)= 1. d0
  RHS (i)= 0. d0
  do k= 1, NPLU
    if (ITEM(k).eq. 1) AMAT(k)= 0. d0
  enddo
endif
```

!C==

#1



#2



Program: 1d.c(10/11)

Conjugate Gradient Method

```

!C
!C +-----+
!C | CG iterations |
!C +-----+
!C===
      R = 1
      Z = 2
      Q = 2
      P = 3
      DD= 4

      do i= 1, N
        W(i, DD)= 1.0D0 / DIAG(i)
      enddo

!C-- {r0}= {b} - [A] {xini} |
!C-   init

      do neib= 1, NEIBPETOT
        do k= export_index(neib-1)+1, export_index(neib)
          kk= export_item(k)
          SENDbuf(k)= PHI(kk)
        enddo
      enddo
    enddo
  
```

Compute $r^{(0)} = b - [A]x^{(0)}$

for $i = 1, 2, \dots$

solve $[M]z^{(i-1)} = r^{(i-1)}$

$\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$

if $i = 1$

$p^{(1)} = z^{(0)}$

else

$\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$

$p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$

endif

$q^{(i)} = [A]p^{(i)}$

$\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$

$x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$

$r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$

check convergence $|r|$

end

Conjugate Gradient Method (CG)

- Matrix-Vector Multiply
- Dot Product
- Preconditioning: in the same way as 1CPU code
- DAXPY: in the same way as 1CPU code

Preconditioning, DAXPY

```
!C
!C-- {z} = [M-1] {r}

do i= 1, N
    W(i, Z) = W(i, DD) * W(i, R)
enddo
```

```
!C
!C-- {x} = {x} + ALPHA*{p}
!C {r} = {r} - ALPHA*{q}

do i= 1, N
    PHI(i) = PHI(i) + ALPHA * W(i, P)
    W(i, R) = W(i, R) - ALPHA * W(i, Q)
enddo
```

Matrix-Vector Multiply (1/2)

Using Comm. Table, {p} is updated before computation

```

!C
!C-- {q} = [A] {p}

do neib= 1, NEIBPETOT
  do k= export_index(neib-1)+1, export_index(neib)
    kk= export_item(k)
    SENDbuf(k)= W(kk, P)
  enddo
enddo

do neib= 1, NEIBPETOT
  is  = export_index(neib-1) + 1
  len_s= export_index(neib) - export_index(neib-1)
  call MPI_Isend (SENDbuf(is), len_s, MPI_DOUBLE_PRECISION, &
&                 NEIBPE(neib), 0, MPI_COMM_WORLD, request_send(neib), ierr)
enddo

do neib= 1, NEIBPETOT
  ir  = import_index(neib-1) + 1
  len_r= import_index(neib) - import_index(neib-1)
  call MPI_Irecv (RECVbuf(ir), len_r, MPI_DOUBLE_PRECISION, &
&                 NEIBPE(neib), 0, MPI_COMM_WORLD, request_recv(neib), ierr)
enddo

call MPI_Waitall (NEIBPETOT, request_recv, stat_recv, ierr)

do neib= 1, NEIBPETOT
  do k= import_index(neib-1)+1, import_index(neib)
    kk= import_item(k)
    W(kk, P)= RECVbuf(k)
  enddo
enddo

```

Matrix-Vector Multiply (2/2)

$$\{q\} = [A]\{p\}$$

```
call MPI_Waitall (NEIBPETOT, request_send, stat_send, ierr)

do i= 1, N
    W(i,Q) = DIAG(i)*W(i,P)
    do j= INDEX(i-1)+1, INDEX(i)
        W(i,Q) = W(i,Q) + AMAT(j)*W(ITEM(j), P)
    enddo
enddo
```

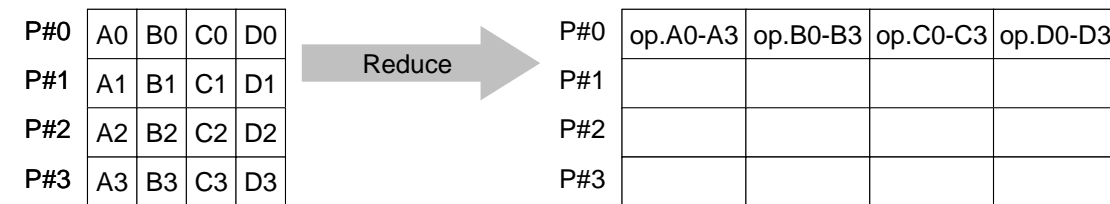
Dot Product

Global Summation by MPI_Allreduce

```
!C
!C-- RH0= {r} {z}

RH00= 0. d0
do i= 1, N
  RH00= RH00 + W(i, R)*W(i, Z)
enddo
call MPI_Allreduce (<b>RH00</b>, <b>RHO</b>, 1, MPI_DOUBLE_PRECISION,
&
                  MPI_SUM, MPI_COMM_WORLD, ierr)
```

MPI_REDUCE



- Reduces values on all processes to a single value
 - Summation, Product, Max, Min etc.

- call MPI_REDUCE**

(sendbuf, recvbuf, count, datatype, op, root, comm, ierr)

- **sendbuf** choice I starting address of send buffer
- **recvbuf** choice O starting address receive buffer
type is defined by "**datatype**"
- **count** I I number of elements in send/receive buffer
- **datatype** I I data type of elements of send/receive buffer
FORTRAN MPI_INTEGER, MPI_REAL, MPI_DOUBLE_PRECISION, MPI_CHARACTER etc.
C MPI_INT, MPI_FLOAT, MPI_DOUBLE, MPI_CHAR etc

- **op** I I reduce operation
MPI_MAX, MPI_MIN, MPI_SUM, MPI_PROD, MPI_LAND, MPI_BAND etc

Users can define operations by **MPI_OP_CREATE**

- **root** I I rank of root process
- **comm** I I communicator
- **ierr** I O completion code

Send/Receive Buffer (Sending/Receiving)

- Arrays of “send (sending) buffer” and “receive (receiving) buffer” often appear in MPI.
- Addresses of “send (sending) buffer” and “receive (receiving) buffer” must be different.

Example of MPI_Reduce (1/2)

```
call MPI_REDUCE  
(sendbuf, recvbuf, count, datatype, op, root, comm, ierr)
```

```
real(kind=8):: x0, x1  
  
call MPI_REDUCE  
(x0, x1, 1, MPI_DOUBLE_PRECISION, MPI_MAX, 0, <comm>, ierr)
```

```
real(kind=8):: x0(4), xmax(4)  
  
call MPI_REDUCE  
(x0, xmax, 4, MPI_DOUBLE_PRECISION, MPI_MAX, 0, <comm>, ierr)
```

Global Max values of X0(i) go to XMAX(i) on #0 process (i=1~4)

Example of MPI_Reduce (2/2)

```
call MPI_REDUCE  
(sendbuf, recvbuf, count, datatype, op, root, comm, ierr)
```

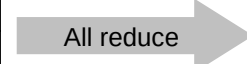
```
real(kind=8) :: X0, XSUM  
  
call MPI_REDUCE  
(X0, XSUM, 1, MPI_DOUBLE_PRECISION, MPI_SUM, 0, <comm>, ierr)
```

Global summation of X0 goes to XSUM on #0 process.

```
real(kind=8) :: X0(4)  
  
call MPI_REDUCE  
(X0(1), X0(3), 2, MPI_DOUBLE_PRECISION, MPI_SUM, 0, <comm>, ierr)
```

- Global summation of X0(1) goes to X0(3) on #0 process.
- Global summation of X0(2) goes to X0(4) on #0 process.

MPI_ALLREDUCE



P#0	A0	B0	C0	D0		P#0	op.A0-A3	op.B0-B3	op.C0-C3	op.D0-D3
P#1	A1	B1	C1	D1	All reduce	P#1	op.A0-A3	op.B0-B3	op.C0-C3	op.D0-D3
P#2	A2	B2	C2	D2		P#2	op.A0-A3	op.B0-B3	op.C0-C3	op.D0-D3
P#3	A3	B3	C3	D3		P#3	op.A0-A3	op.B0-B3	op.C0-C3	op.D0-D3

- MPI_Reduce + MPI_Bcast
- Summation (of dot products) and MAX/MIN values are likely to utilized in each process
- **call MPI_ALLREDUCE**
(sendbuf, recvbuf, count, datatype, op, comm, ierr)
 - **sendbuf** choice I starting address of send buffer
 - **recvbuf** choice O starting address receive buffer
type is defined by "**datatype**"
 - **count** I I number of elements in send/recv buffer
 - **datatype** I I data type of elements in send/recv buffer
 - **op** I I reduce operation
 - **comm** I I communicator
 - **ierr** I O completion code

CG method (1/5)

```

!C
!C-- {r0} = {b} - [A] {xini}
do neib= 1, NEIBPETOT
  do k= export_index(neib-1)+1, export_index(neib)
    kk= export_item(k)
    SENDbuf(k)= PHI(kk)
  enddo
enddo

do neib= 1, NEIBPETOT
  is = export_index(neib-1) + 1
  len_s= export_index(neib) - export_index(neib-1)
  call MPI_Isend (SENDbuf(is), len_s,
                  MPI_DOUBLE_PRECISION,
                  NEIBPE(neib), 0, MPI_COMM_WORLD,
                  request_send(neib), ierr)
&
&
&
enddo

do neib= 1, NEIBPETOT
  ir = import_index(neib-1) + 1
  len_r= import_index(neib) - import_index(neib-1)
  call MPI_Irecv (RECVbuf(ir), len_r,
                  MPI_DOUBLE_PRECISION,
                  NEIBPE(neib), 0, MPI_COMM_WORLD,
                  request_recv(neib), ierr)
&
&
&
enddo
call MPI_Waitall (NEIBPETOT, request_recv, stat_recv, ier end

do neib= 1, NEIBPETOT
  do k= import_index(neib-1)+1, import_index(neib)
    kk= import_item(k)
    PHI(kk)= RECVbuf(k)
  enddo
enddo
call MPI_Waitall (NEIBPETOT, request_send, stat_send, ierr)

```

Compute $r^{(0)} = b - [A] x^{(0)}$

for $i = 1, 2, \dots$

solve $[M] z^{(i-1)} = r^{(i-1)}$

$\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$

if $i = 1$

$p^{(1)} = z^{(0)}$

else

$\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$

$p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$

endif

$q^{(i)} = [A] p^{(i)}$

$\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$

$x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$

$r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$

check convergence $|r|$

CG method (2/5)

```

do i= 1, N
  W(i, R) = DIAG(i)*PHI(i)
  do j= INDEX(i-1)+1, INDEX(i)
    W(i, R) = W(i, R) + AMAT(j)*PHI(ITEM(j))
  enddo
enddo

BNRM20= 0.0D0
do i= 1, N
  BNRM20 = BNRM20 + RHS(i) **2
  W(i, R) = RHS(i) - W(i, R)
enddo
call MPI_Allreduce (BNRM20, BNRM2, 1,
&                                MPI_DOUBLE_PRECISION,
&                                MPI_SUM, MPI_COMM_WORLD, ierr)

!C*****
do iter= 1, ITERmax

!C-- {z}= [M-1] {r}
  do i= 1, N
    W(i, Z)= W(i, DD) * W(i, R)
  enddo
!C-- RHO= {r} {z}

  RH00= 0. d0
  do i= 1, N
    RH00= RH00 + W(i, R)*W(i, Z)
  enddo
  call MPI_Allreduce (RH00, RHO, 1, MPI_DOUBLE_PRECISION,
&                                MPI_SUM, MPI_COMM_WORLD, ierr)

```

Compute $r^{(0)} = b - [A]x^{(0)}$

for $i = 1, 2, \dots$

solve $[M]z^{(i-1)} = r^{(i-1)}$

$\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$

if $i = 1$

$p^{(1)} = z^{(0)}$

else

$\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$

$p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$

endif

$q^{(i)} = [A]p^{(i)}$

$\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$

$x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$

$r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$

check convergence $|r|$

end

CG method (3/5)

```

!C
!C-- {p} = {z} if ITER=1
!C  BETA= RHO / RH01 otherwise

  if ( iter.eq.1 ) then
    do i= 1, N
      W(i,P)= W(i,Z)
    enddo
  else
    BETA= RHO / RH01
    do i= 1, N
      W(i,P)= W(i,Z) + BETA*W(i,P)
    enddo
  endif

!C-- {q}= [A] {p}

  do neib= 1, NEIBPETOT
    do k= export_index(neib-1)+1, export_index(neib)
      kk= export_item(k)
      SENDbuf(k)= W(kk,P)
    enddo
  enddo

  do neib= 1, NEIBPETOT
    is  = export_index(neib-1) + 1
    len_s= export_index(neib) - export_index(neib-1)
    call MPI_Isend (SENDbuf(is), len_s, MPI_DOUBLE_PRECISION,
    &                               NEIBPE(neib), 0, MPI_COMM_WORLD,
    &                               request_send(neib), ierr)
    &
  enddo

```

Compute $r^{(0)} = b - [A]x^{(0)}$

for $i = 1, 2, \dots$

solve $[M]z^{(i-1)} = r^{(i-1)}$

$\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$

if $i=1$

$p^{(1)}= z^{(0)}$

else

$\beta_{i-1} = \rho_{i-1}/\rho_{i-2}$

$p^{(i)}= z^{(i-1)} + \beta_{i-1} p^{(i-1)}$

endif

$q^{(i)}= [A]p^{(i)}$

$\alpha_i = \rho_{i-1}/p^{(i)}q^{(i)}$

$x^{(i)}= x^{(i-1)} + \alpha_i p^{(i)}$

$r^{(i)}= r^{(i-1)} - \alpha_i q^{(i)}$

check convergence $|r|$

end

&

&

CG method (4/5)

```

do neib= 1, NEIBPETOT
    ir = import_index(neib-1) + 1
    len_r= import_index(neib) - import_index(neib-1)
    call MPI_Recv (RECVbuf(ir), len_r,
&           MPI_DOUBLE_PRECISION,
&           NEIBPE(neib), 0, MPI_COMM_WORLD,
&           request_recv(neib), ierr)
enddo
call MPI_Waitall (NEIBPETOT, request_recv, stat_recv, ierr)

do neib= 1, NEIBPETOT
    do k= import_index(neib-1)+1, import_index(neib)
        kk= import_item(k)
        W(kk, P)= RECVbuf(k)
    enddo
enddo
call MPI_Waitall (NEIBPETOT, request_send, stat_send, ierr)

do i= 1, N
    W(i, Q) = DIAG(i)*W(i, P)
    do j= INDEX(i-1)+1, INDEX(i)
        W(i, Q) = W(i, Q) + AMAT(j)*W(ITEM(j), P)
    enddo
enddo

!C
!C-- ALPHA= RHO / {p} {q}

C10= 0. d0
do i= 1, N
    C10= C10 + W(i, P)*W(i, Q)
enddo
call MPI_Allreduce (C10, C1, 1, MPI_DOUBLE_PRECISION, MPI_SUM, MPI_COMM_WORLD, ierr)
ALPHA= RHO / C1

```

Compute $r^{(0)} = b - [A]x^{(0)}$

for $i = 1, 2, \dots$

solve $[M]z^{(i-1)} = r^{(i-1)}$

$\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$

if $i = 1$

$p^{(1)} = z^{(0)}$

else

$\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$

$p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$

endif

$q^{(i)} = [A]p^{(i)}$

$\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$

$x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$

$r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$

check convergence $|r|$

end

CG method (5/5)

```

!C
!C-- {x} = {x} + ALPHA*{p}
!C {r} = {r} - ALPHA*{q}

do i= 1, N
  PHI(i)= PHI(i) + ALPHA * W(i, P)
  W(i, R)= W(i, R) - ALPHA * W(i, Q)
enddo

DNRM20 = 0.0
do i= 1, N
  DNRM20= DNRM20 + W(i, R)**2
enddo

call MPI_Allreduce (DNRM20, DNRM2, 1,
&                               MPI_DOUBLE_PRECISION,
&                               MPI_SUM, MPI_COMM_WORLD, ierr)

RESID= dsqrt(DNRM2/BNRM2)

if (my_rank.eq.0.and.mod(iter,1000).eq.0) then
  write (*, '(i8,1pe16.6)') iter, RESID
endif

if ( RESID.le.EPS) goto 900
RH01 = RHO

enddo

```

Compute $r^{(0)} = b - [A]x^{(0)}$

for $i = 1, 2, \dots$

solve $[M]z^{(i-1)} = r^{(i-1)}$

$\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$

if $i=1$

$p^{(1)} = z^{(0)}$

else

$\beta_{i-1} = \rho_{i-1}/\rho_{i-2}$

$p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$

endif

$q^{(i)} = [A]p^{(i)}$

$\alpha_i = \rho_{i-1}/p^{(i)} q^{(i)}$

$x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$

$r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$

check convergence $|r|$

end

Program: 1d.f (11/11)

Output by Each Process

```
!C
!C-- OUTPUT
  if (my_rank, eq. 0) then
    write (*, '(2(1pe16. 6))') E1Time-S1Time, E2Time-E1Time
  endif

  write (*, '(/a)' ) '### TEMPERATURE'
  do i= 1, N
    write (*, '(2i8, 2(1pe16. 6))') my_rank, i, PHI(i)
  enddo

call MPI_FINALIZE (ierr)
end program heat1Dp
```

- Overview
- Distributed Local Data
- Program
- **Results**

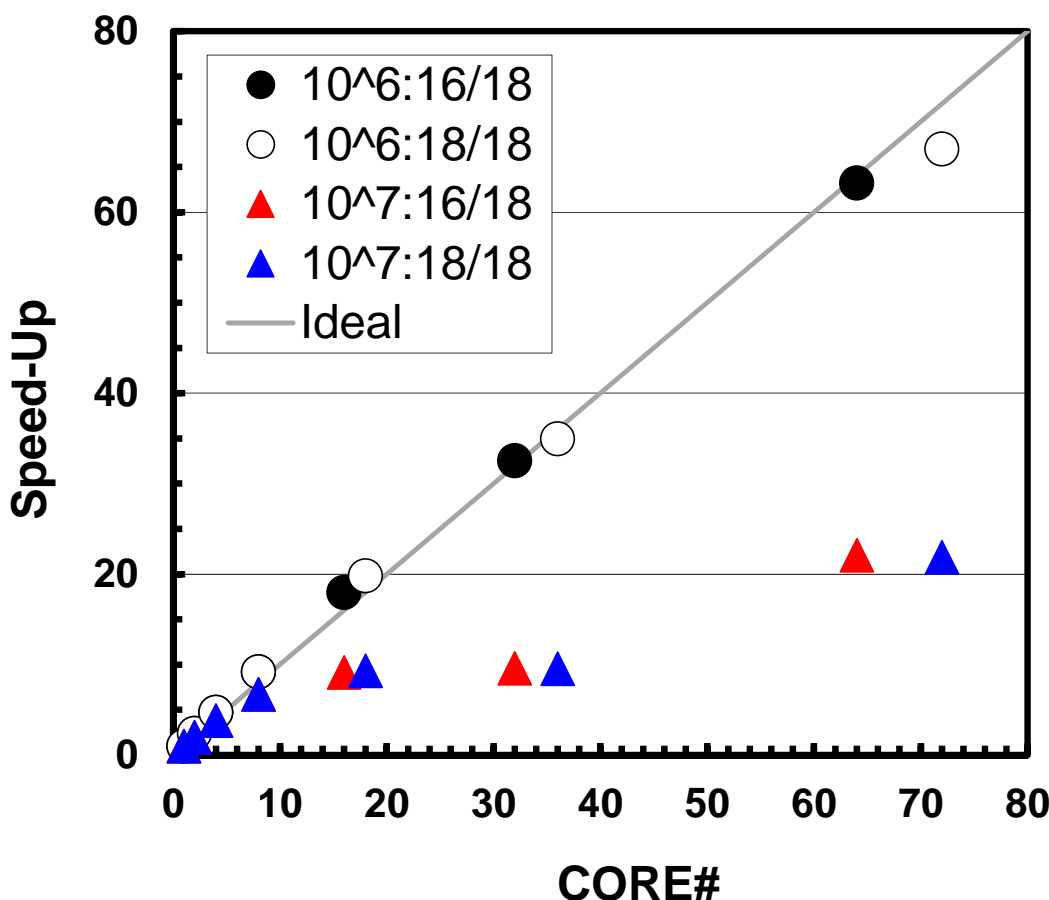
Results: Time for CG Solver

Time for 1,000 iterations, Strong Scaling

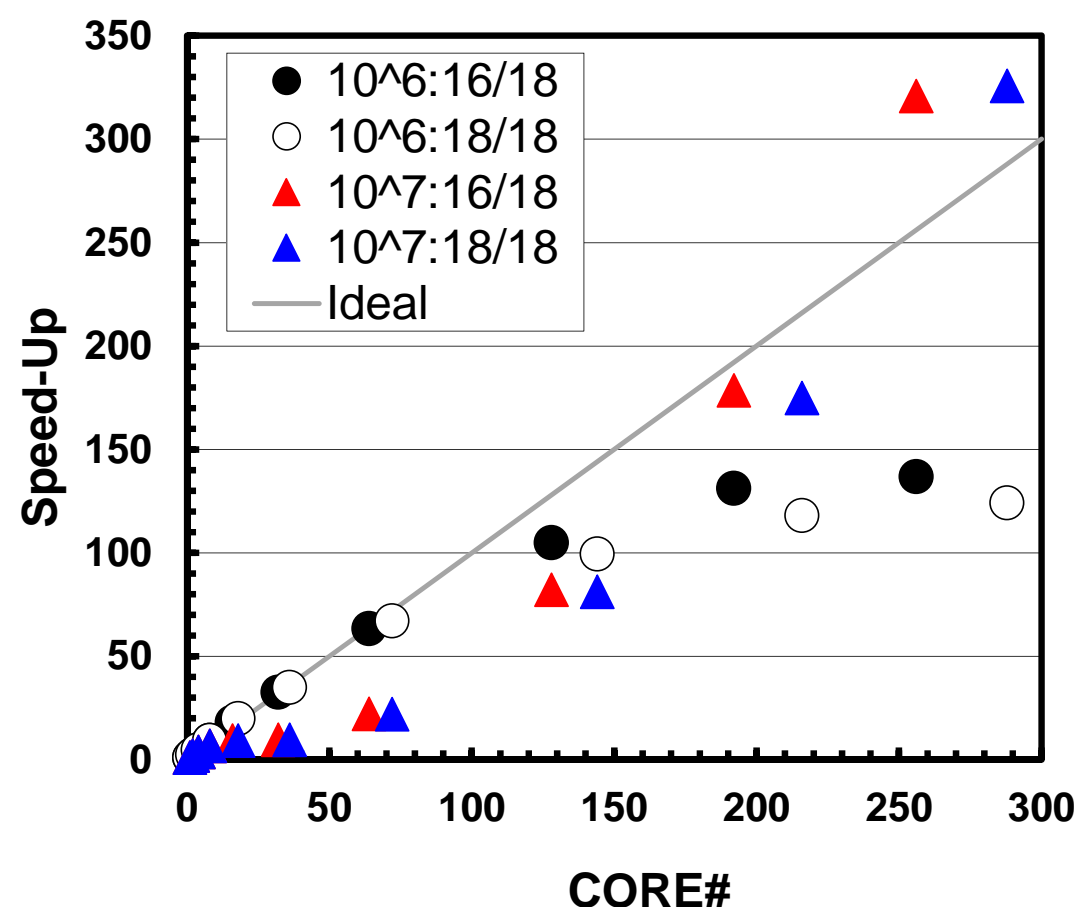
16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large

up to 2 nodes



up to 8 nodes



Performance is lower than ideal one

- Time for MPI communication
 - Time for sending data
 - Communication bandwidth between nodes
 - Time is proportional to size of sending/receiving buffers
- Time for starting MPI
 - latency
 - does not depend on size of buffers
 - depends on number of calling, increases according to process #
 - $O(10^0)$ - $O(10^1)$ μ sec.
- Synchronization of MPI
 - Increases according to number of processes

Performance is lower than ideal one (cont.)

- If computation time is relatively small (N is small in S1-3), these effects are not negligible.
 - If the size of messages is small, effect of “latency” is significant.

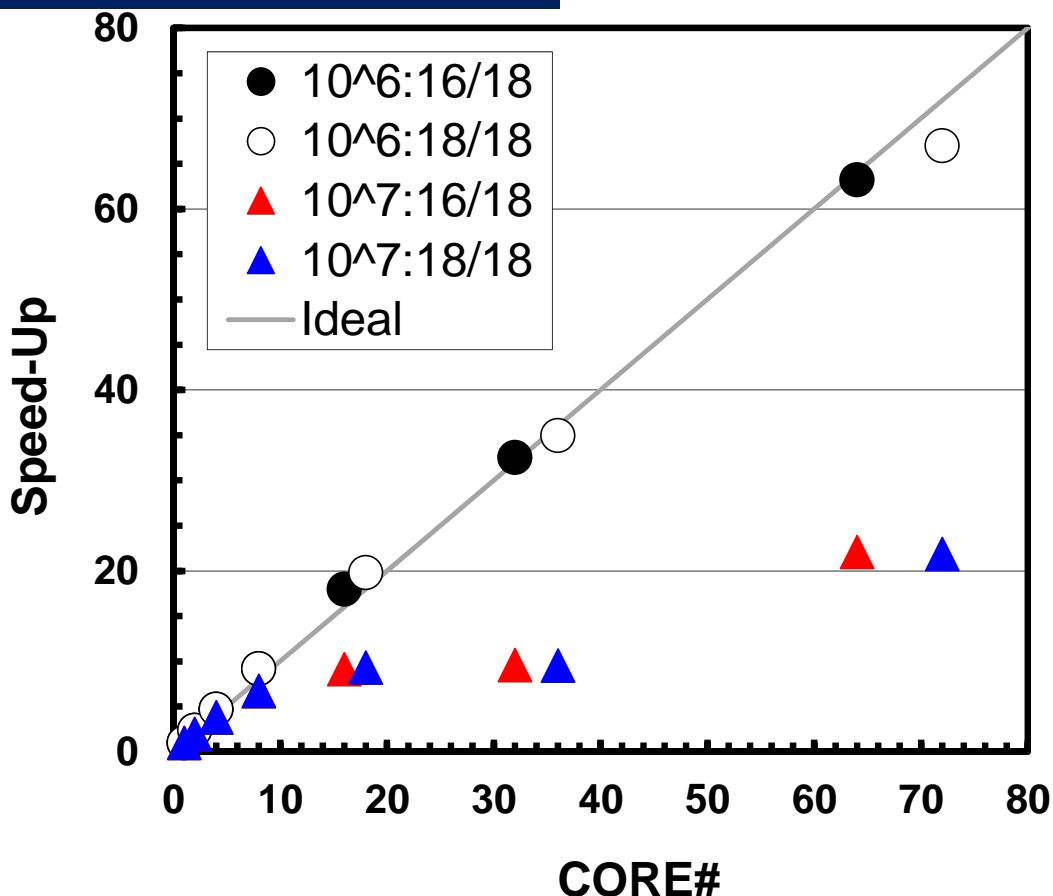
Results: Time for CG Solver

Time for 1,000 iterations, Strong Scaling

16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large

up to 2 nodes



- in a single node (< 36 cores), performance of small cases ($N=10^6$) are rather better.
- Effect of memory contention/saturation (not communication)
- Memory throughput on each node is constant for 8+ cores
- If problem size is small, cache can be well-utilized. Therefore, effect of memory bandwidth is small.

Shell Scripts (1/2)

```

#!/bin/sh
#PBS -q u-lecture4
#PBS -N 1D
#PBS -l select=1:mpiprocs=16
#PBS -Wgroup_list=gt14
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o test.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=16

mpirun ./impimap.sh ./a.out

```

go.sh: 16 cores may be randomly selected from 36 cores

```

#!/bin/sh
#PBS -q u-lecture4
#PBS -N 1D
#PBS -l select=1:mpiprocs=16
#PBS -Wgroup_list=gt14
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o test.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_PROCESSOR_LIST=0-15

mpirun ./impimap.sh ./a.out

```

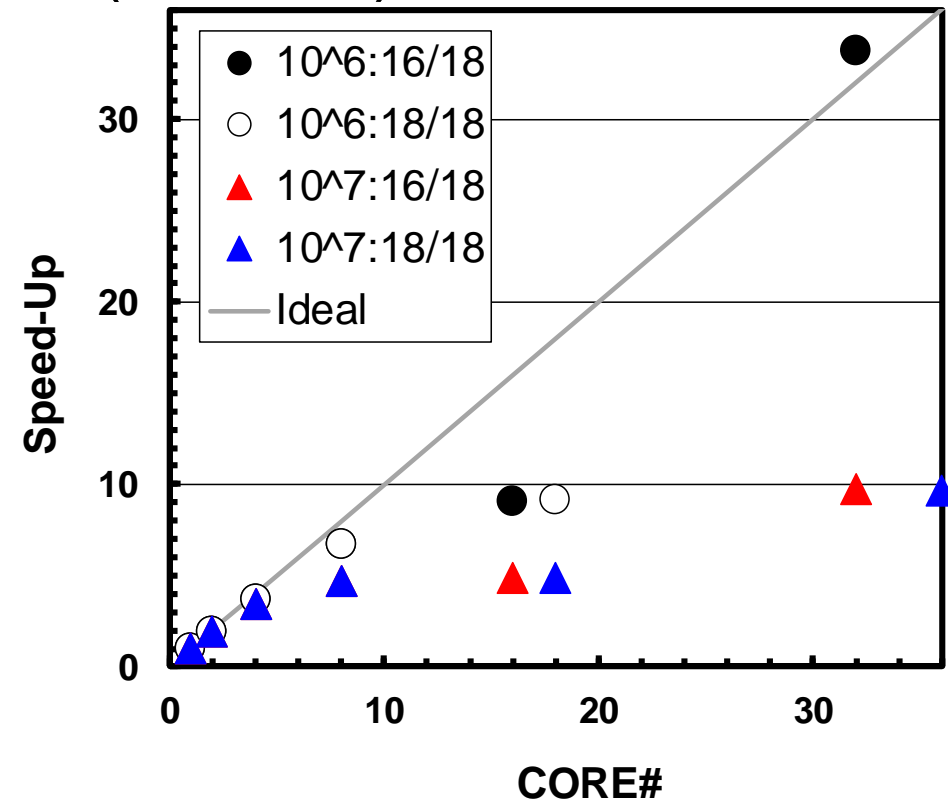
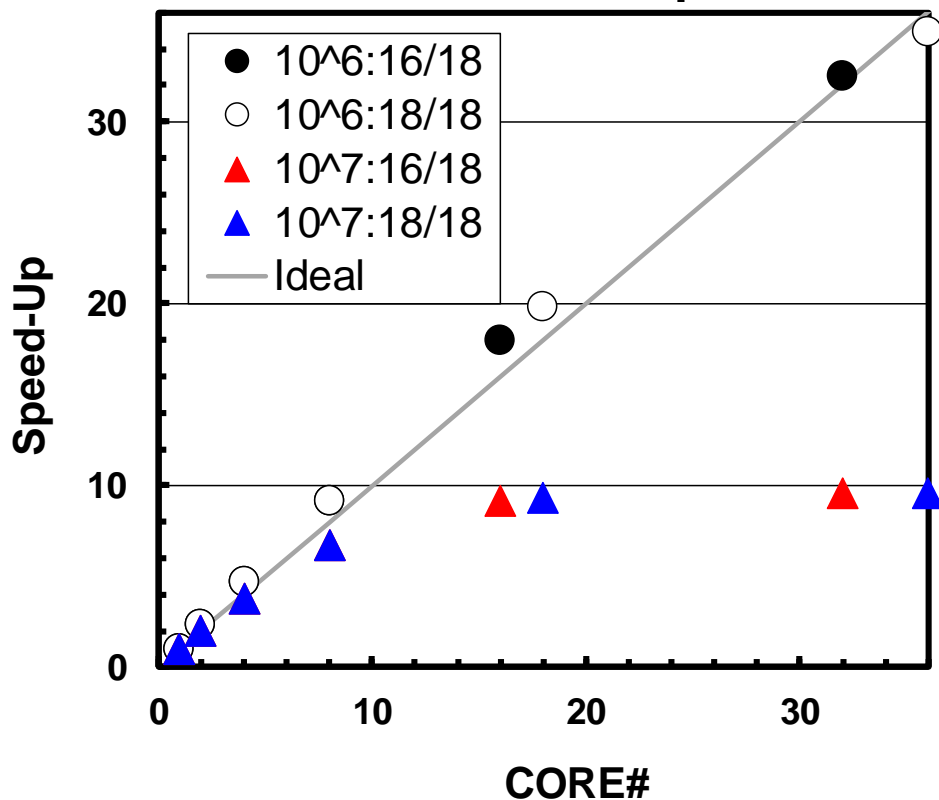
a16.sh: Performance is mostly same, but this one is more stable (small fluctuation)

Results: Time for CG Solver

Time for 1,000 iterations, Strong Scaling

16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large
up to 36 cores (1-node)



```
export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=16
```

```
export
I_MPI_PIN_PROCESSOR_LIST=
0-15
```

Shell Scripts (2/2)

```

#!/bin/sh
#PBS -q u-lecture4
#PBS -N 1D
#PBS -l select=8:mpiprocs=32
#PBS -Wgroup_list=gt14
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o test.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=32

mpirun ./impimap.sh ./a.out

```

go.sh:

```

#!/bin/sh
#PBS -q u-lecture4
#PBS -N 1D
#PBS -l select=8:mpiprocs=32
#PBS -Wgroup_list=gt14
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o test.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_PROCESSOR_LIST=0-15,18-33

mpirun ./impimap.sh ./a.out

```

a32.sh: Performance is mostly same, but this one is more stable (small fluctuation)

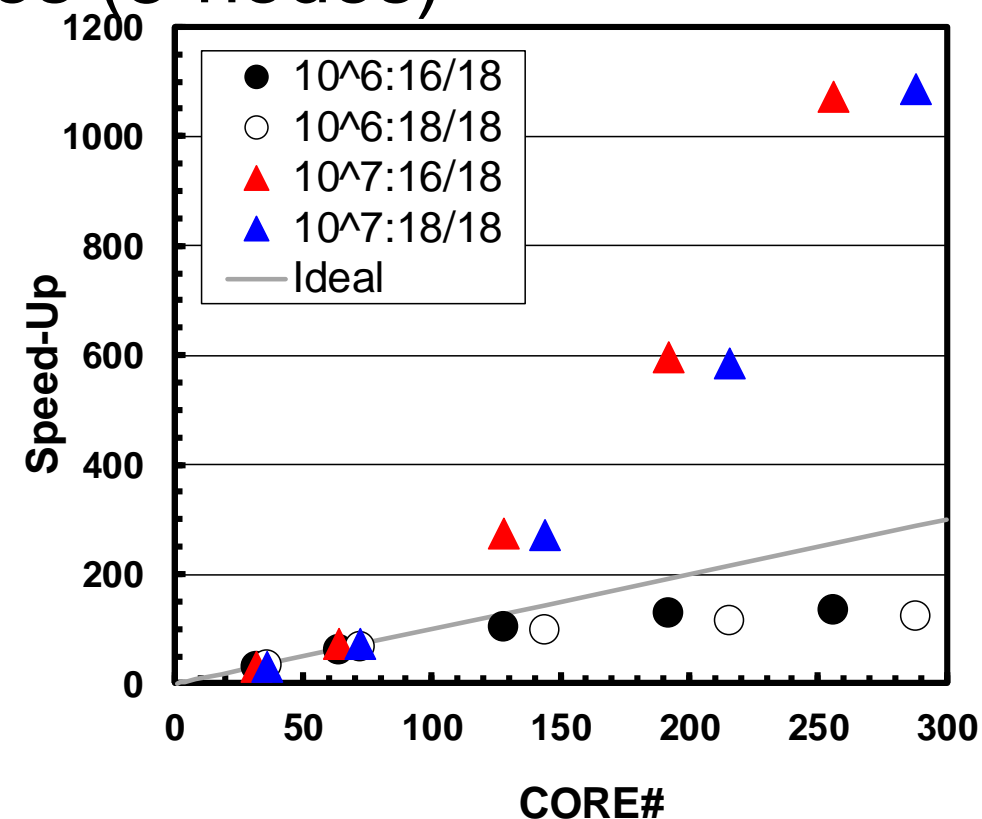
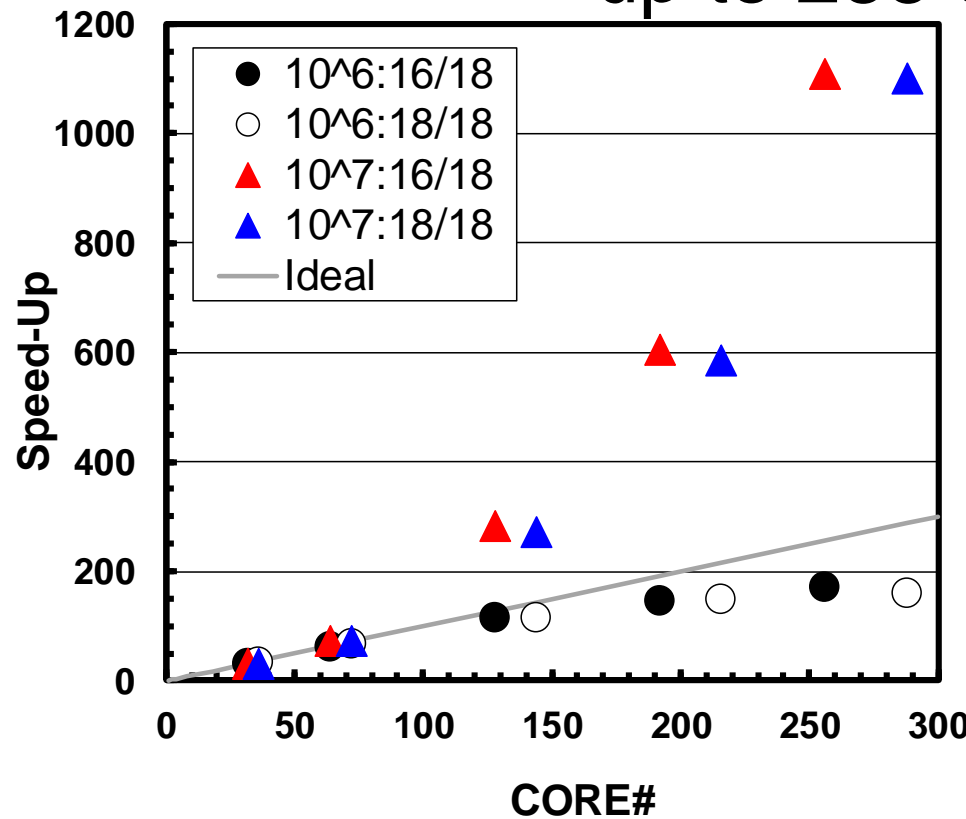
Results: Time for CG Solver

Time for 1,000 iterations, Strong Scaling

16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large

up to 288 cores (8-nodes)



```
export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=32
```

```
export
I_MPI_PIN_PROCESSOR_LIST=
0-15,18-33
```

STREAM benchmark

<http://www.cs.virginia.edu/stream/>

- Benchmarks for Memory Bandwidth
 - Copy: $c(i) = a(i)$
 - Scale: $c(i) = s \cdot b(i)$
 - Add: $c(i) = a(i) + b(i)$
 - Triad: $c(i) = a(i) + s \cdot b(i)$

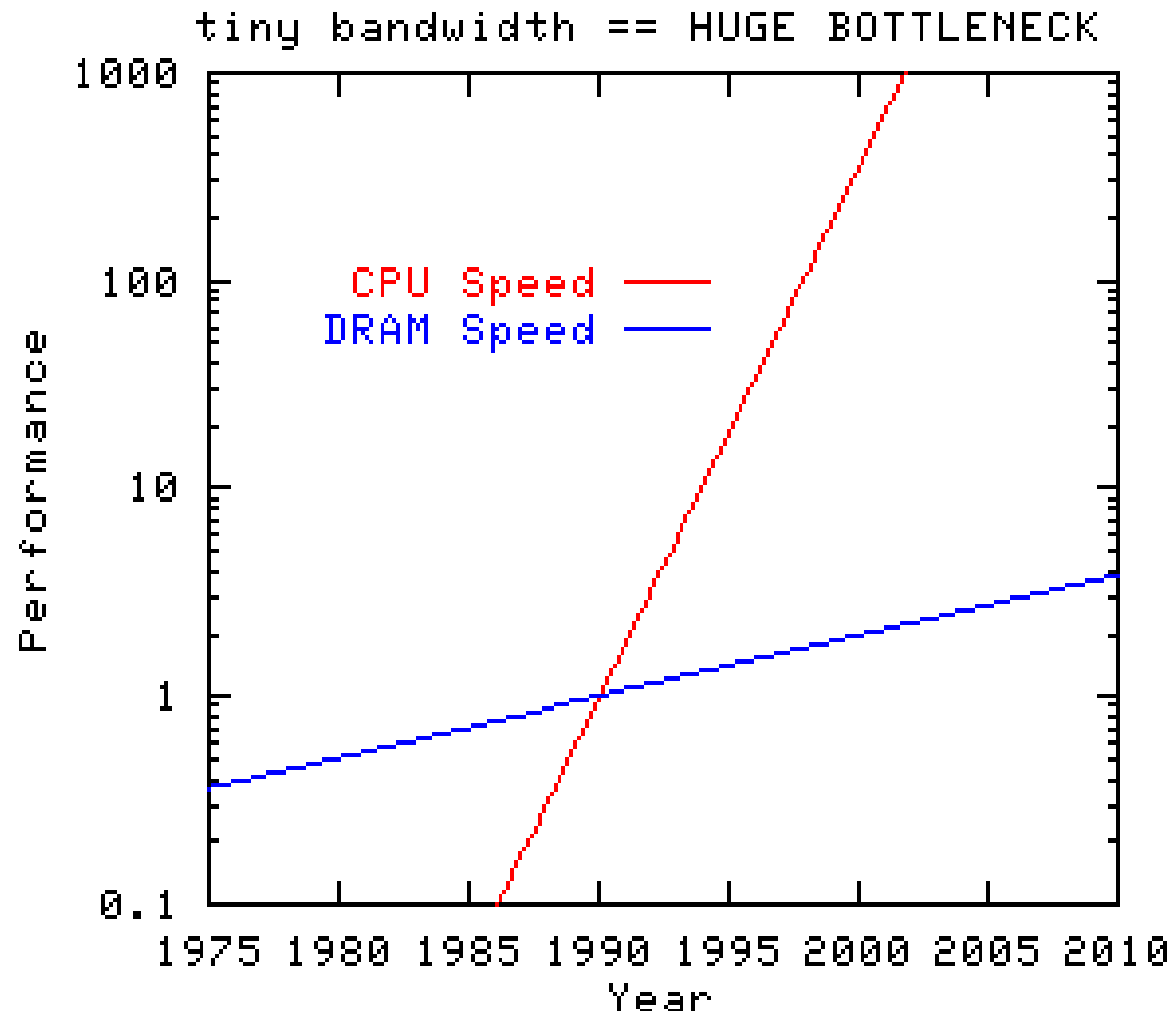
Double precision appears to have 16 digits of accuracy
Assuming 8 bytes per DOUBLE PRECISION word

Number of processors = 16
 Array size = 2000000
 Offset = 0
 The total memory requirement is 732.4 MB (45.8MB/task)
 You are running each test 10 times

The *best* time for each test is used
 EXCLUDING the first and last iterations

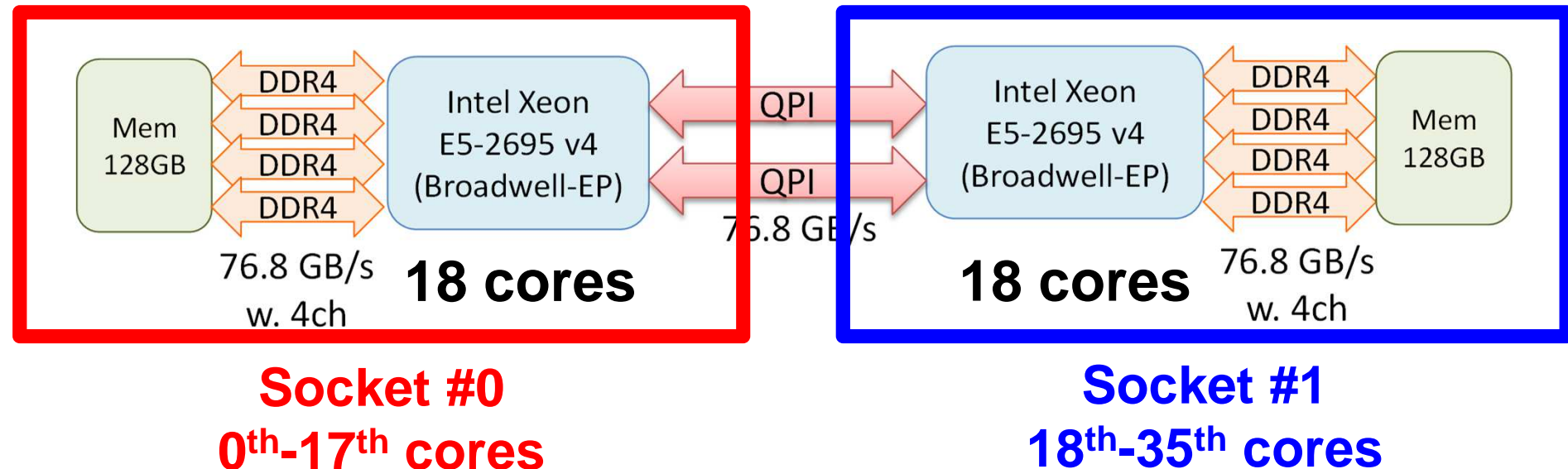
Function	Rate (MB/s)	Avg time	Min time	Max time
Copy:	18334.1898	0.0280	0.0279	0.0280
Scale:	18035.1690	0.0284	0.0284	0.0285
Add:	18649.4455	0.0412	0.0412	0.0413
Triad:	19603.8455	0.0394	0.0392	0.0398

Gap between performance of CPU and Memory



Copy, Compile and Run

```
>$ cd /lustre/gt14/t14xxx/pFEM  
>$ cp /lustre/gt00/z30088/class_eps/F/stream.tar .  
>$ tar xvf stream.tar  
>$ cd mpi/stream  
  
>$ mpiifort -O3 -xCORE-AVX2 -align array32byte stream.f -o stream  
>$ qsub XXX.sh
```



s01.sh: Use 1 core (0th)

```

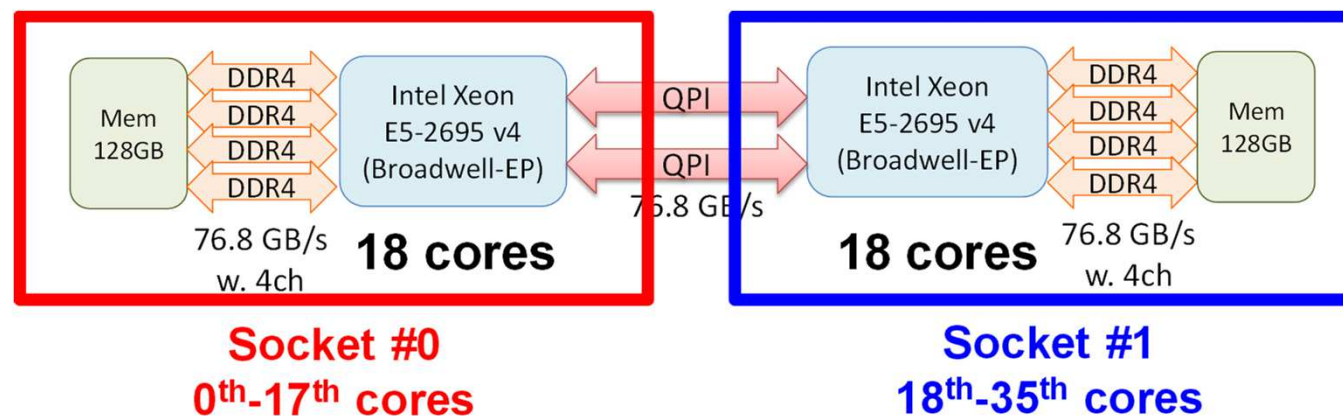
#!/bin/sh

#PBS -q u-lecture7
#PBS -N stream
#PBS -l select=1:mpiprocs=1      MPI Process # (1-36)
#PBS -Wgroup_list=gt07
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o t01.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_PROCESSOR_LIST=0      use 0th core
mpirun ./impimap.sh ./stream

```



s16.sh: Use 16 cores (0-15th)

```

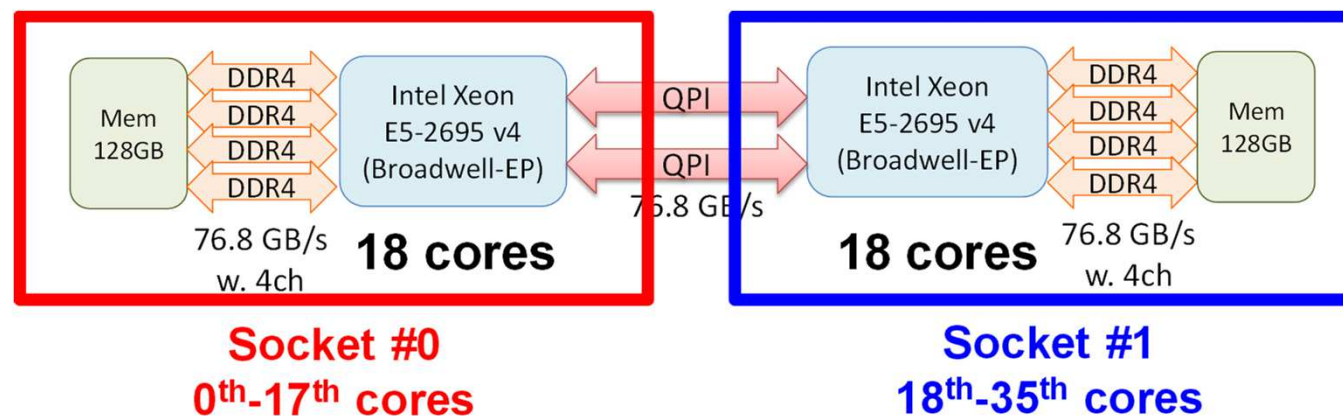
#!/bin/sh

#PBS -q u-lecture7
#PBS -N stream
#PBS -l select=1:mpiprocs=16      MPI Process # (1-36)
#PBS -Wgroup_list=gt07
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o t16.1st

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_PROCESSOR_LIST=0-15      use 0-15th core
mpirun ./impimap.sh ./stream

```



s32.sh: Use 32 cores (16 ea)

```

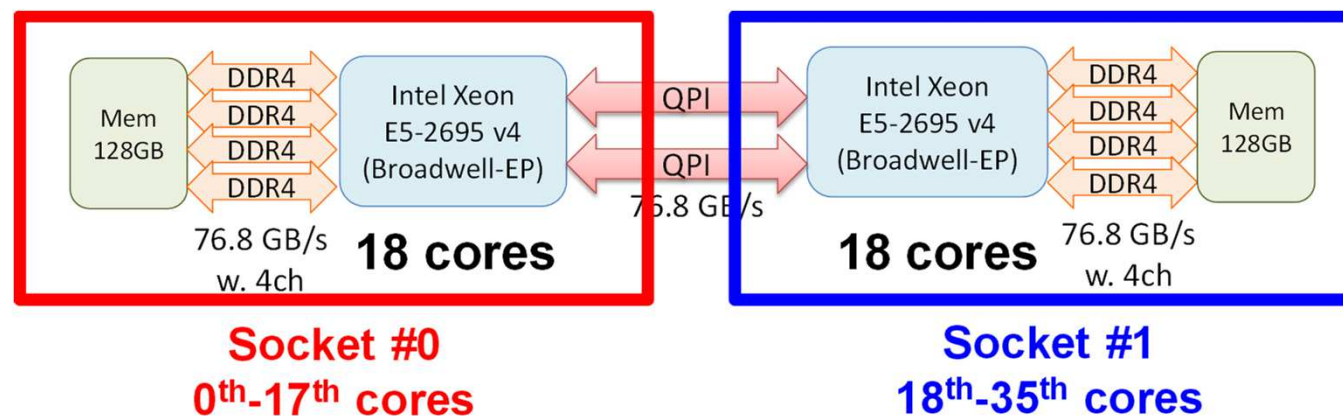
#!/bin/sh

#PBS -q u-lecture7
#PBS -N stream
#PBS -l select=1:mpiprocs=32      MPI Process # (1-36)
#PBS -Wgroup_list=gt07
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o t32.1st

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_PROCESSOR_LIST=0-15,18-33
mpirun ./impimap.sh ./stream

```



s36.sh: Use 36 cores (ALL)

```

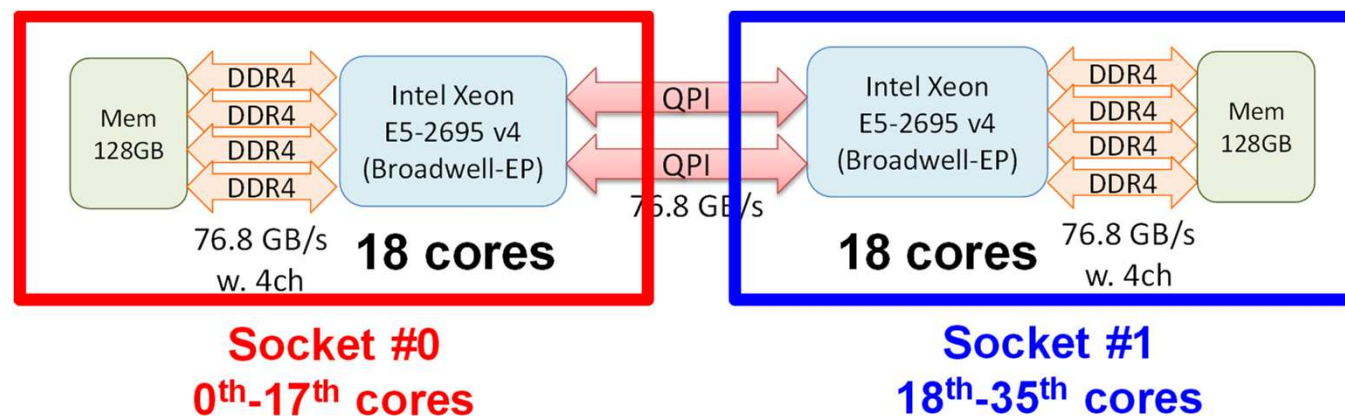
#!/bin/sh

#PBS -q u-lecture7
#PBS -N stream
#PBS -l select=1:mpiprocs=36      MPI Process # (1-36)
#PBS -Wgroup_list=gt07
#PBS -l walltime=00:05:00
#PBS -e err
#PBS -o t36.1st

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_PROCESSOR_LIST=0-35
mpirun ./impimap.sh ./stream

```



Results of Triad on a Single Node of Reedbush-U

Peak is 153.6 GB/sec.

Thread #	GB/sec	Speed-up
1	16.11	1.00
2	30.95	1.92
4	54.72	3.40
8	64.59	4.01
16	65.18	4.04
18	64.97	4.03
32	130.0	8.07
36	128.2	7.96

Exercises

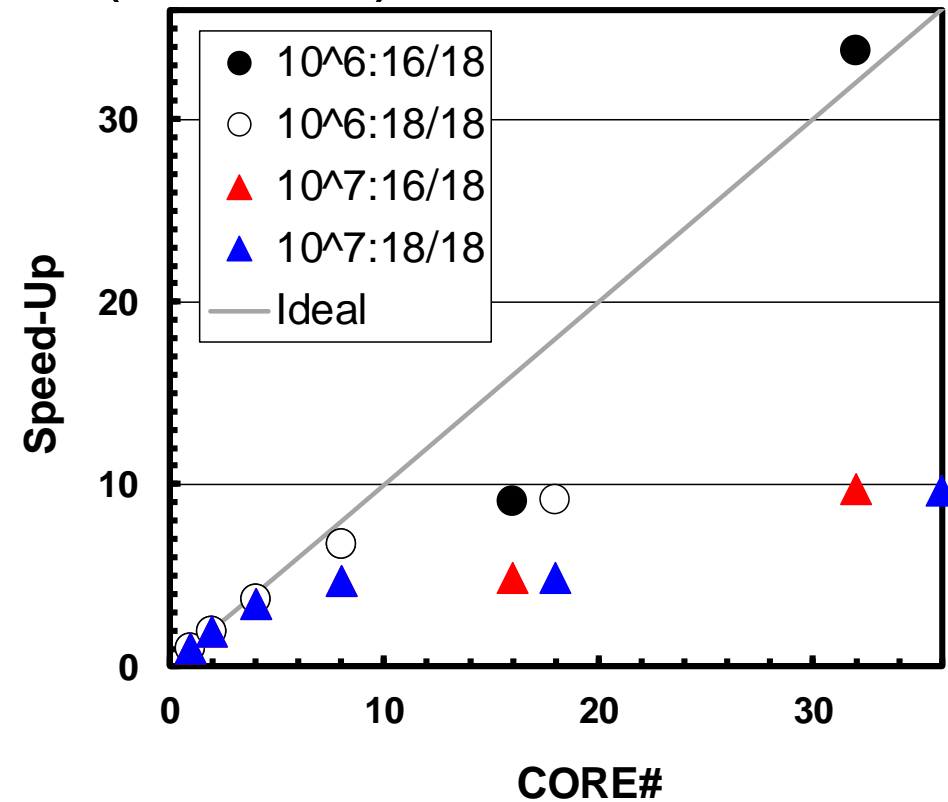
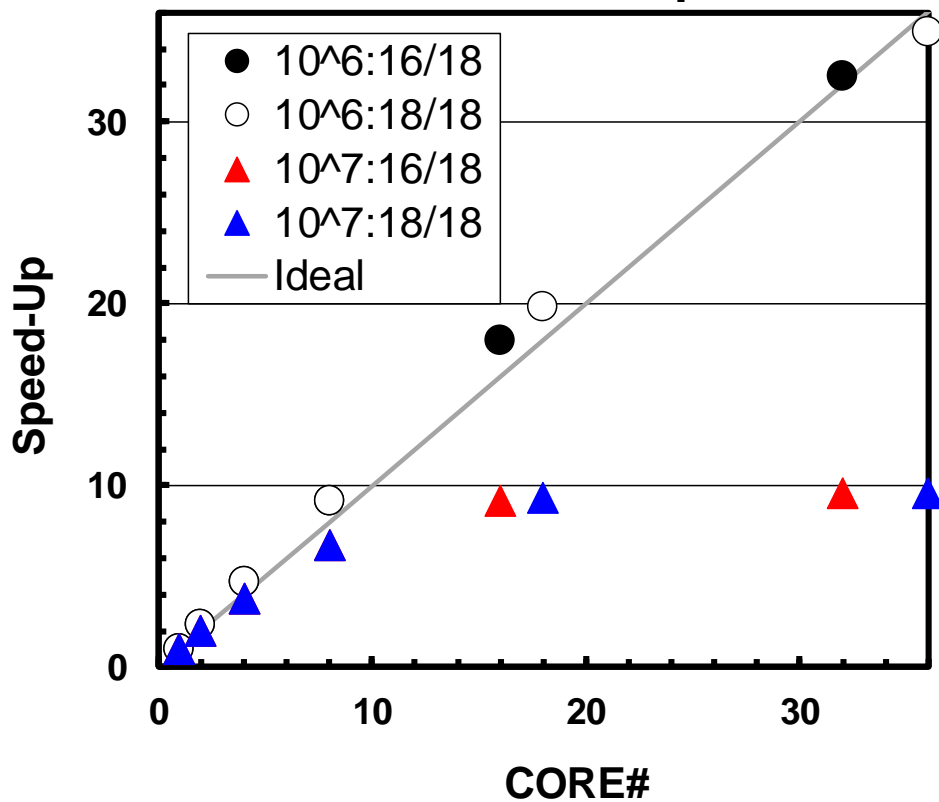
- Running the code
- Try various number of processes (1-36)
- OpenMP-version and Single PE version are available
 - Fortran, C
 - Web-site of STREAM
 - <http://www.cs.virginia.edu/stream/>

Results: Time for CG Solver

Time for 1,000 iterations, Strong Scaling

16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large
up to 36 cores (1-node)



```
export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=16
```

```
export
I_MPI_PIN_PROCESSOR_LIST=
0-15
```

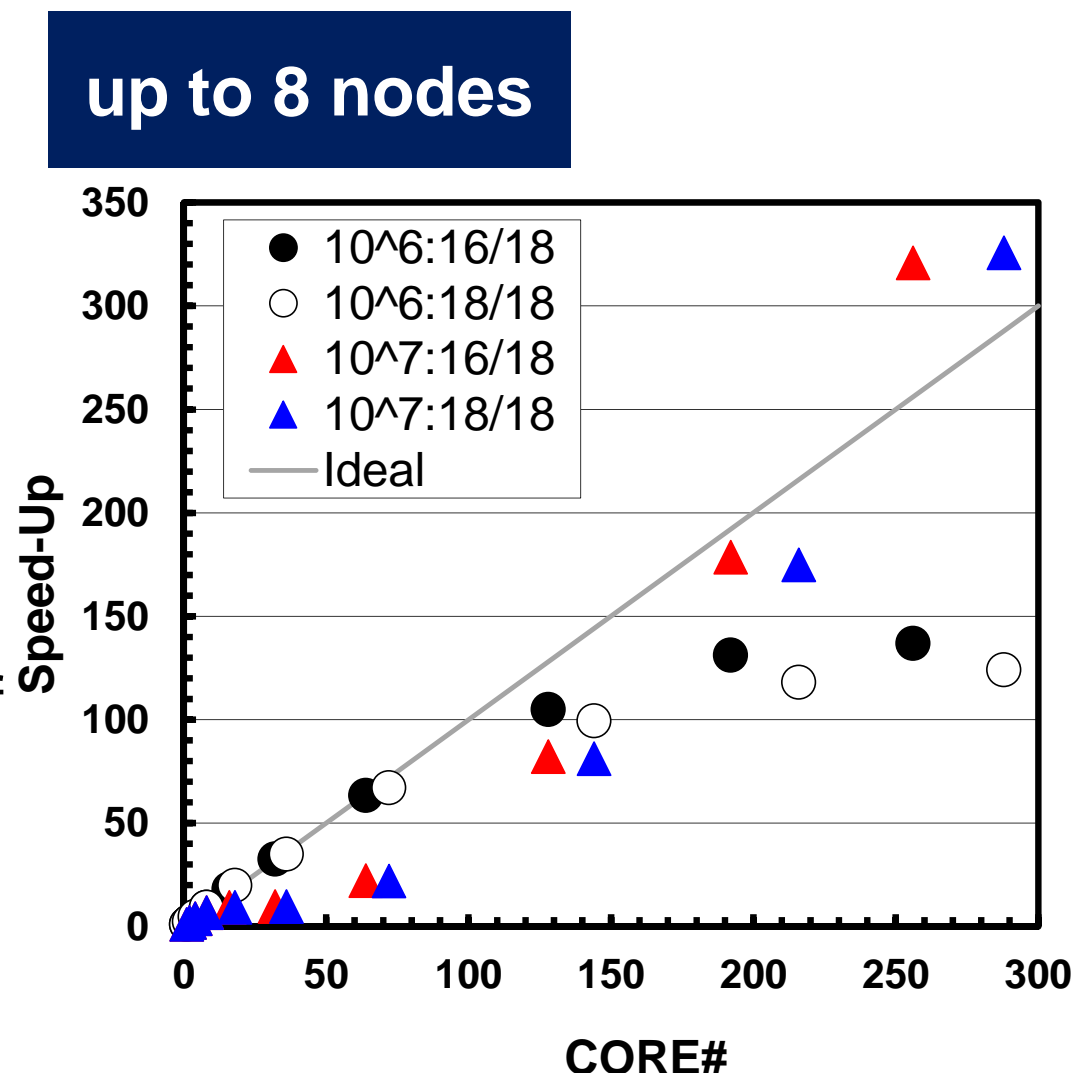
Results: Time for CG Solver

Time for 1,000 iterations, Strong Scaling

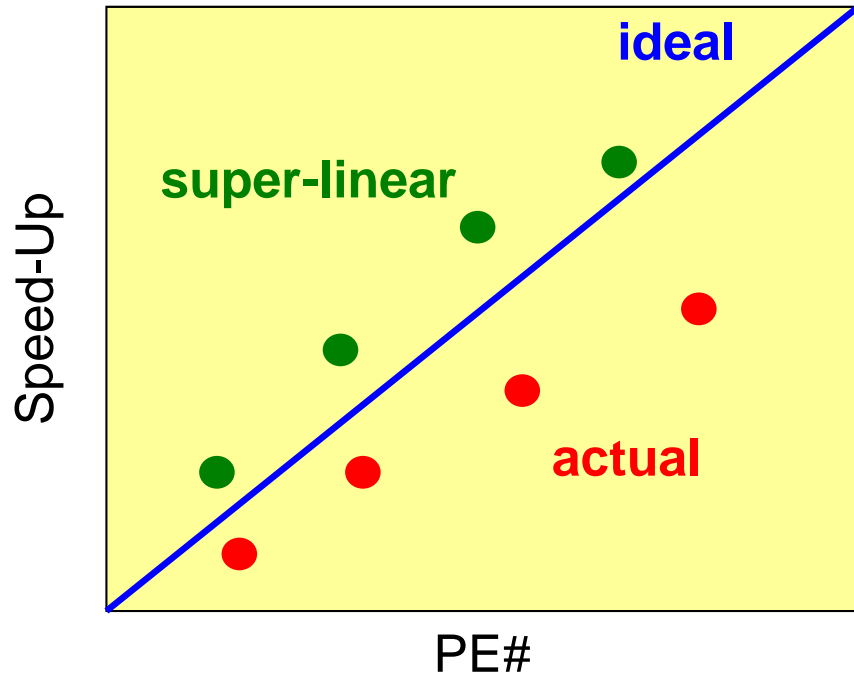
16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large

- Performance of $N=10^6$ case decreases, as node# increases.
- Performance of $N=10^7$ becomes close to ideal one gradually, and superlinear situation occurs at 256 cores (8 nodes).
- 16/18 is rather better if number of node is large.
 - Memory is more well-utilized



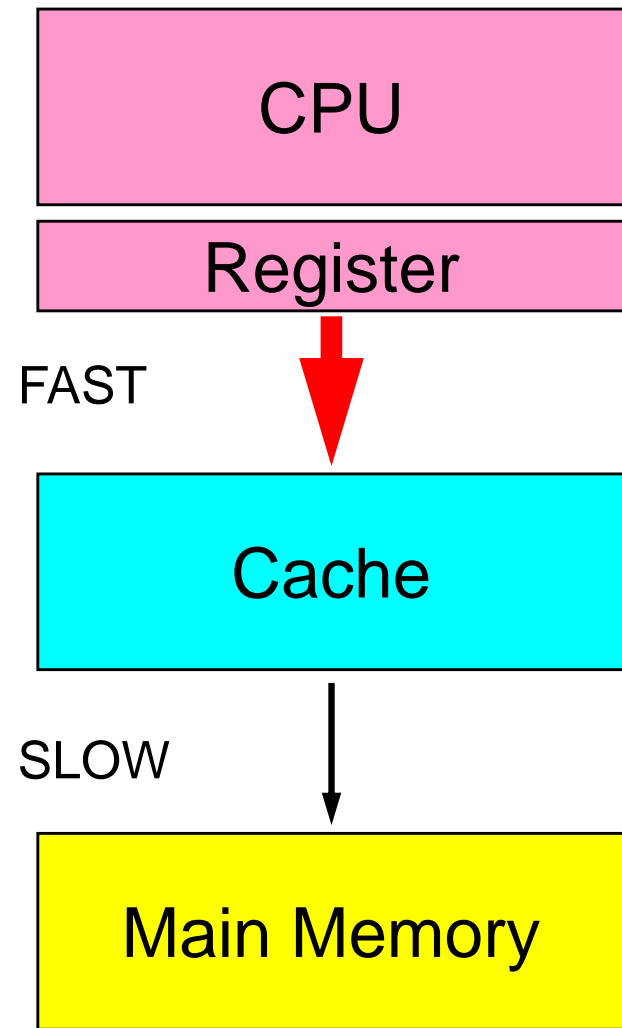
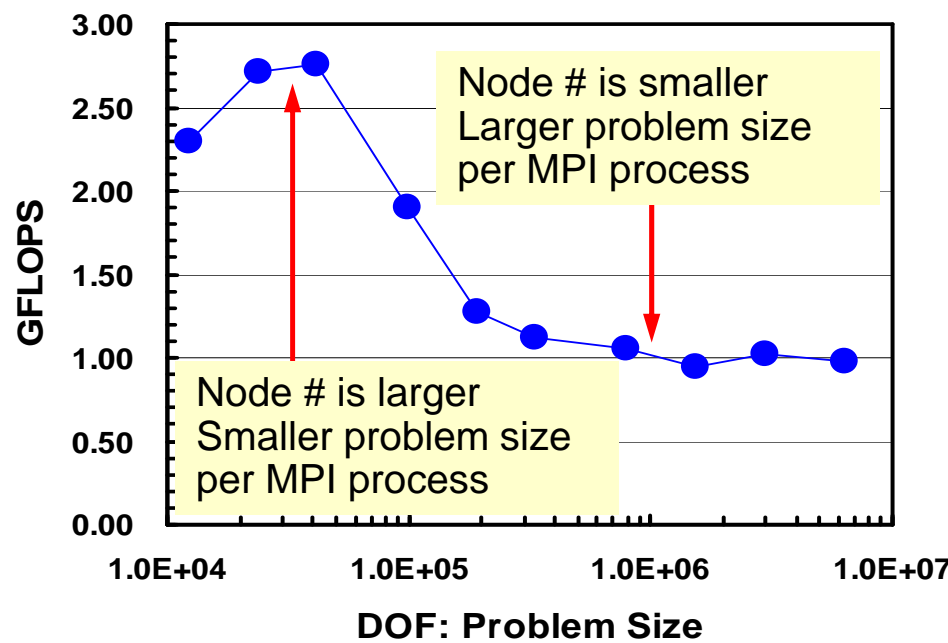
Super-Linear in Strong Scaling



- In strong scaling case where entire problem size is fixed, performance is generally lower than the ideal one due to communication overhead.
- But sometime, actual performance may be better than the ideal one. This is called “super-linear”
 - only for scalar processors
 - does not happen in vector processors

Why does “Super-Linear” happen ?

- Effect of Cache
- In scalar processors, performance for smaller problem is generally better.
 - Cache is well-utilized.



Memory Copy is expensive (1/2)

SENDbuf



```

do neib= 1, NEIBPETOT
  do k= export_index(neib-1)+1, export_index(neib)
    kk= export_item(k)
    SENDbuf (k) = VAL (kk)
  enddo
enddo

do neib= 1, NEIBPETOT
  iS_e= export_index(neib-1) + 1
  iE_e= export_index(neib   )
  BUFlength_e= iE_e + 1 - iS_e

  call MPI_ISEND
  &           (SENDbuf(iS_e), BUFlength_e, MPI_INTEGER, NEIBPE(neib), 0, &
  &           MPI_COMM_WORLD, request_send(neib), ierr)
enddo

call MPI_WAITALL (NEIBPETOT, request_send, stat_recv, ierr)

```

Copied to sending buffers

Memory Copy is expensive (2/2)

```

do neib= 1, NEIBPETOT
    iS_i= import_index(neib-1) + 1
    iE_i= import_index(neib )
    BUFlength_i= iE_i + 1 - iS_i

    call MPI_IRecv
&          (RECVbuf(iS_i), BUFlength_i, MPI_INTEGER, NEIBPE(neib), 0, &
& MPI_COMM_WORLD, request_recv(neib), ierr)
enddo

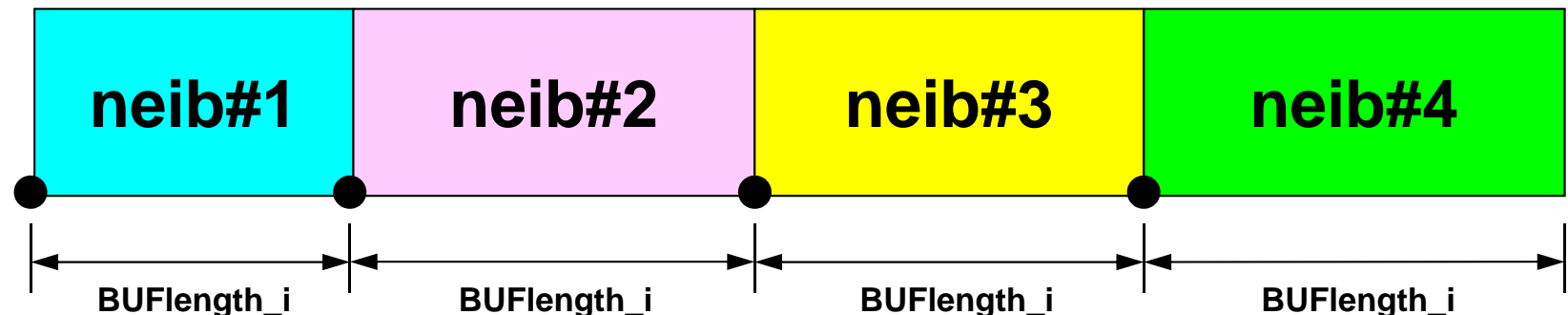
call MPI_WAITALL (NEIBPETOT, request_recv, stat_recv, ierr)

do neib= 1, NEIBPETOT
    do k= import_index(neib-1)+1, import_index(neib)
        kk= import_item(k)
        VAL(kk)= RECVbuf (k)
    enddo
enddo

```

Copied from receiving buffer

RECVbuf



`import_index(0)+1 import_index(1)+1 import_index(2)+1 import_index(3)+1 import_index(4)`

Results: Time for CG Solver

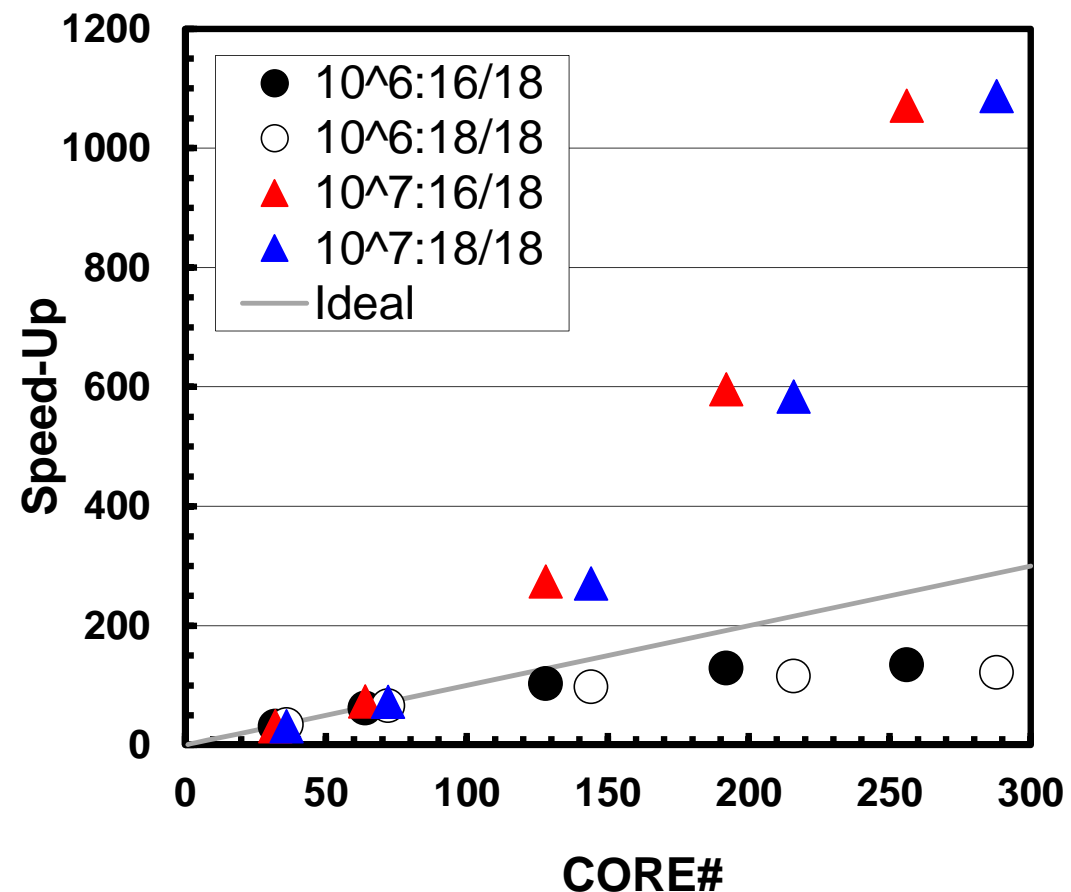
Time for 1,000 iterations, Strong Scaling

16/18: 16 of 18 cores on socket, 18/18: 18 of 18 cores

16/18 is rather better, if number of nodes is large

- It is reasonable to evaluate strong scalability for multiple nodes based on the performance by a node with 32/36 cores
- In this figure, performance of the case with 32 cores is set to 32 (results of 16/18 and 18/18 are directly compared).
- L2 cache: 256kB/core
- L3: 45MB/socket (shared)

up to 8 nodes



Summary: Parallel FEM

- Proper design of data structure of distributed local meshes.
- Open Technical Issues
 - Parallel Mesh Generation, Parallel Visualization
 - Parallel Preconditioner for Ill-Conditioned Problems
 - Large-Scale I/O

Distributed Local Data Structure for Parallel Computation

- Distributed local data structure for domain-to-domain communications has been introduced, which is appropriate for such applications with sparse coefficient matrices (e.g. FDM, FEM, FVM etc.).
 - SPMD
 - Local Numbering: Internal pts to External pts
 - Generalized communication table
- Everything is easy, if proper data structure is defined:
 - Values at boundary pts are copied into sending buffers
 - Send/Recv
 - Values at external pts are updated through receiving buffers

If numbering of external nodes is continuous in each neighboring process ...

	84	81	85	82	83	86	88	87	
96	57	58	59	60	61	62	63	64	73
95	49	50	51	52	53	54	55	56	74
94	41	42	43	44	45	46	47	48	80
93	33	34	35	36	37	38	39	40	79
92	25	26	27	28	29	30	31	32	78
91	17	18	19	20	21	22	23	24	77
90	9	10	11	12	13	14	15	16	76
89	1	2	3	4	5	6	7	8	75
	65	66	67	68	69	70	71	72	

[A]{p} = {q} (Original)

```

allocate (stat_send(MPI_STATUS_SIZE, NEIBPETOT))
allocate (stat_recv(MPI_STATUS_SIZE, NEIBPETOT))
allocate (request_send(NEIBPETOT)); allocate (request_recv(NEIBPETOT))

do neib= 1, NEIBPETOT
  do k= export_index(neib-1)+1, export_index(neib)
    kk= export_item(k)
    SENDbuf(k)= W(kk, P)
  enddo
enddo

do neib= 1, NEIBPETOT
  is   = export_index(neib-1) + 1
  len_s= export_index(neib) - export_index(neib-1)
  call MPI_Isend (SENDbuf(is), len_s, MPI_DOUBLE_PRECISION,
&                      NEIBPE(neib), 0, MPI_COMM_WORLD, &
&                      request_send(neib), ierr)
  &&
  enddo

do neib= 1, NEIBPETOT
  ir   = import_index(neib-1) + 1
  len_r= import_index(neib) - import_index(neib-1)
  call MPI_Irecv (RECVbuf(ir), len_r, MPI_DOUBLE_PRECISION,
&                      NEIBPE(neib), 0, MPI_COMM_WORLD, &
&                      request_recv(neib), ierr)
  &&
  enddo
  call MPI_Waitall (NEIBPETOT, request_recv, stat_recv, ierr)

do neib= 1, NEIBPETOT
  do k= import_index(neib-1)+1, import_index(neib)
    kk= import_item(k)
    W(kk, P)= RECVbuf(k)
  enddo
enddo
call MPI_Waitall (NEIBPETOT, request_send, stat_send, ierr)

```

[A]{p}={q} (Mod.): No Copy for RECV

```

allocate (stat_send(MPI_STATUS_SIZE, 2*NEIBPETOT))
allocate (request_send(2*NEIBPETOT))

do neib= 1, NEIBPETOT
    do k= export_index(neib-1)+1, export_index(neib)
        kk= export_item(k)
        SENDbuf(k)= W(kk, P)
    enddo
enddo

do neib= 1, NEIBPETOT
    is = export_index(neib-1) + 1
    len_s= export_index(neib) - export_index(neib-1)
    call MPI_Isend (SENDbuf(is), len_s, MPI_DOUBLE_PRECISION,
&                               NEIBPE(neib), 0, MPI_COMM_WORLD,
&                               request_send(neib), ierr)
    &&
enddo

do neib= 1, NEIBPETOT
    ir = import_index(neib-1) + 1
    len_r= import_index(neib) - import_index(neib-1)
    call MPI_Irecv (W(ir+N, P), len_r, MPI_DOUBLE_PRECISION,
&                               NEIBPE(neib), 0, MPI_COMM_WORLD,
&                               request_send(neib+NEIBPETOT), ierr)
    &&
enddo

call MPI_Waitall (2*NEIBPETOT, request_send, stat_send, ierr)

```

Comp. Time for 1,000 Iterations

32x8 MPI Processes

```
#PBS -q u-lecture4
#PBS -N 1d
#PBS -l select=8:mpiprocs=32
#PBS -Wgroup_list=gt14
#PBS -l walltime=00:10:00
#PBS -e err
#PBS -o test.lst

cd $PBS_O_WORKDIR
. /etc/profile.d/modules.sh

export I_MPI_PIN_DOMAIN=socket
export I_MPI_PERHOST=32
mpirun ./impimap.sh ./a.out
```

