

# **Introduction to Parallel FEM in Fortran Parallel Data Structure**

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Technical & Scientific Computing II (4820-1028)  
Seminar on Computer Science II (4810-1205)  
Hybrid Distributed Parallel Computing (3747-111)

# Parallel Computing

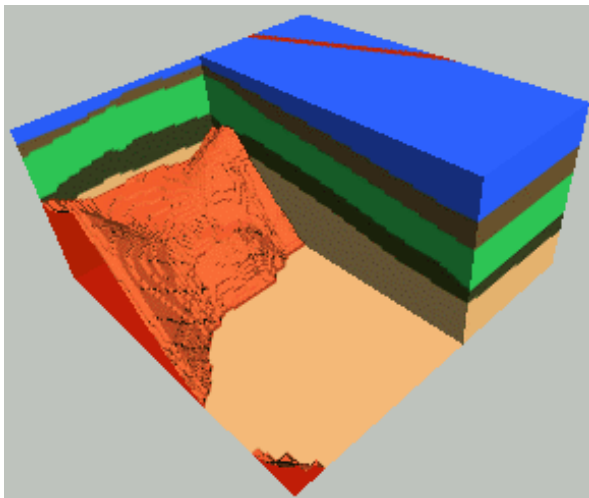
- Faster, Larger & More Complicated
- Scalability
  - Solving  $N^x$  scale problem using  $N^x$  computational resources during same computation time
    - for large-scale problems: Weak Scaling
    - e.g. CG solver: more iterations needed for larger problems
  - Solving a problem using  $N^x$  computational resources during  $1/N$  computation time
    - for faster computation: Strong Scaling

# What is Parallel Computing ? (1/2)

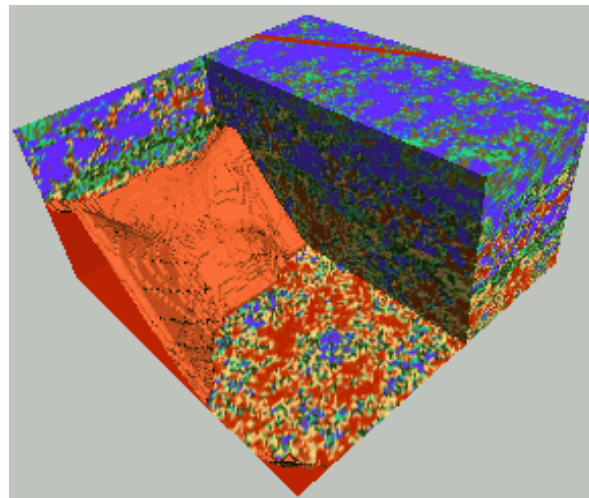
- to solve larger problems faster

## Homogeneous/Heterogeneous Porous Media

Lawrence Livermore National Laboratory



Homogeneous

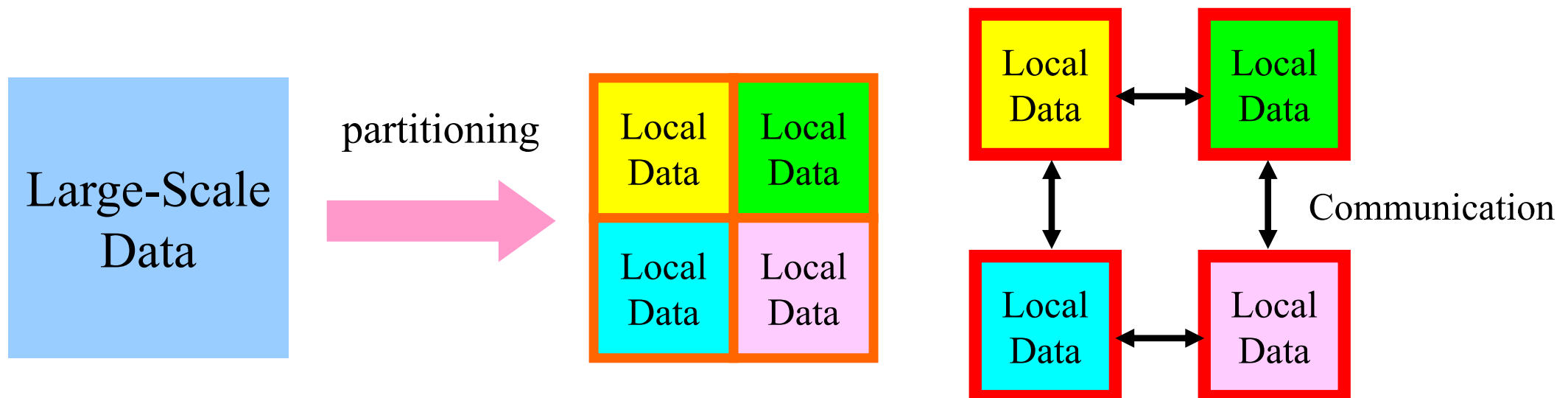


Heterogeneous

**very fine meshes are required for simulations of heterogeneous field.**

# What is Parallel Computing ? (2/2)

- PC with 1GB memory : 1M meshes are the limit for FEM
  - Southwest Japan with 1,000km x 1,000km x 100km in 1km mesh  
->  $10^8$  meshes
- Large Data -> Domain Decomposition -> Local Operation
- Inter-Domain Communication for Global Operation



# What is Communication ?

- Parallel Computing -> Local Operations
- Communications are required in Global Operations for Consistency.

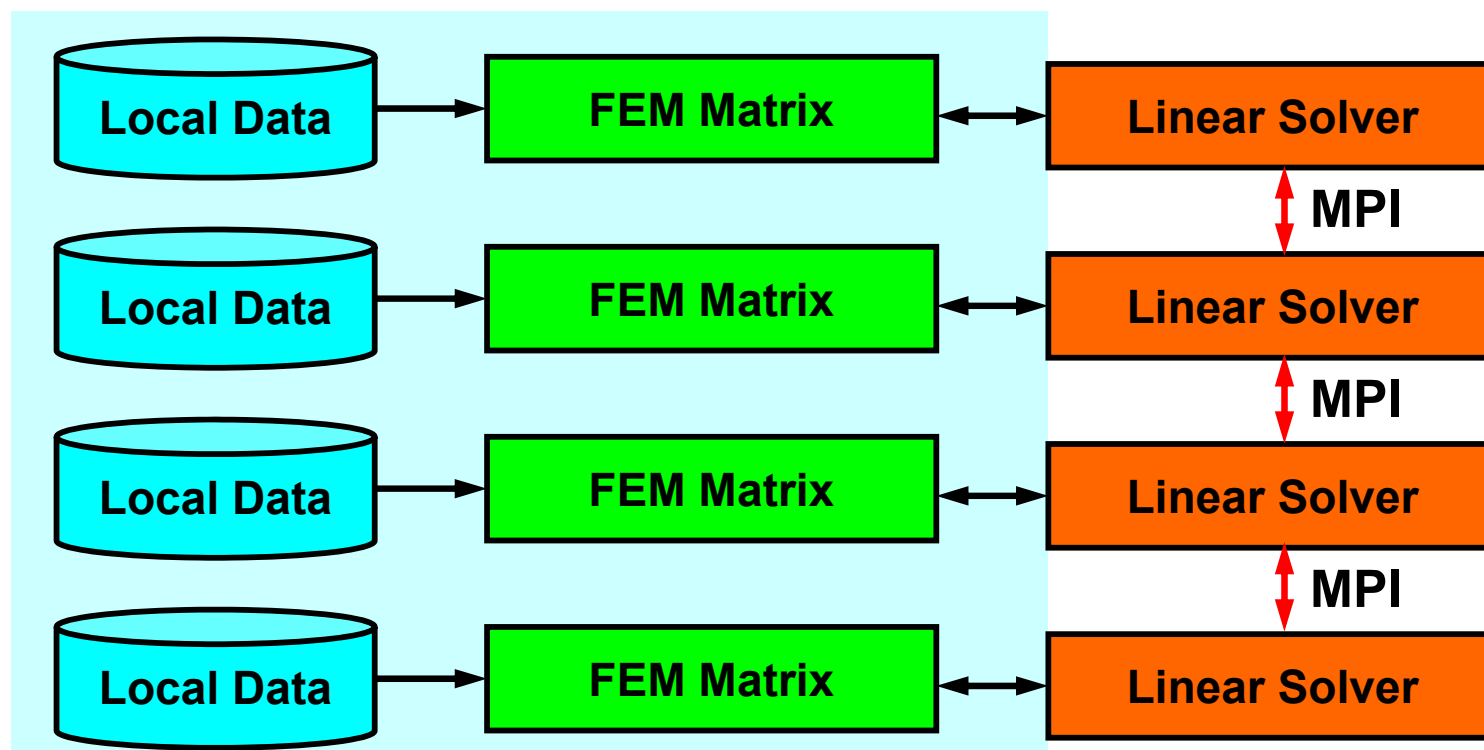
# Operations in Parallel FEM

## SPMD: Single-Program Multiple-Data

Large Scale Data -> partitioned into Distributed Local Data Sets.

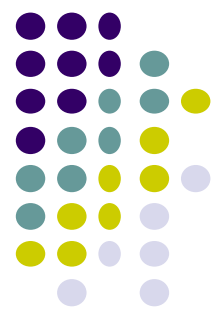
FEM code can assemble coefficient matrix for each local data set :  
this part could be completely local, same as serial operations

Global Operations & Communications happen only in Linear Solvers  
dot products, matrix-vector multiply, preconditioning



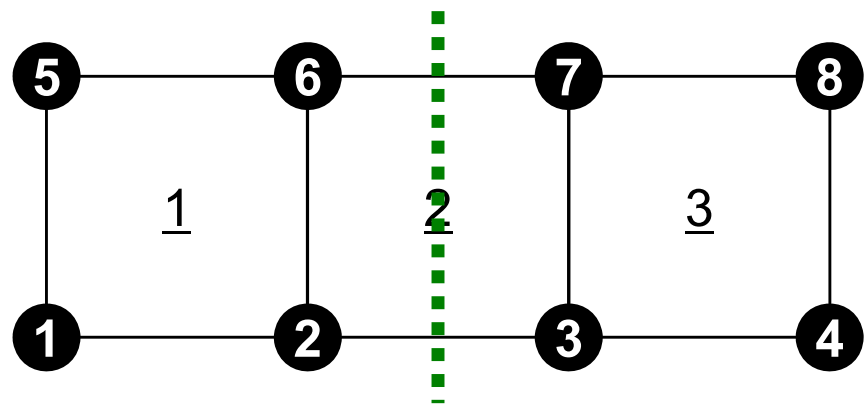
# Parallel FEM Procedures

- Design on “Local Data Structure” is important
  - for SPMD-type operations in the previous page
- Matrix Generation
- Preconditioned Iterative Solvers for Linear Equations



# Bi-Linear Square Elements

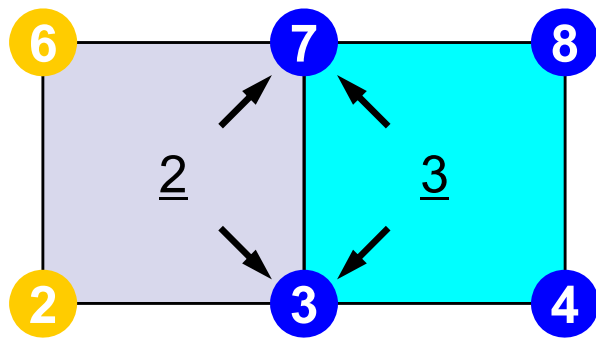
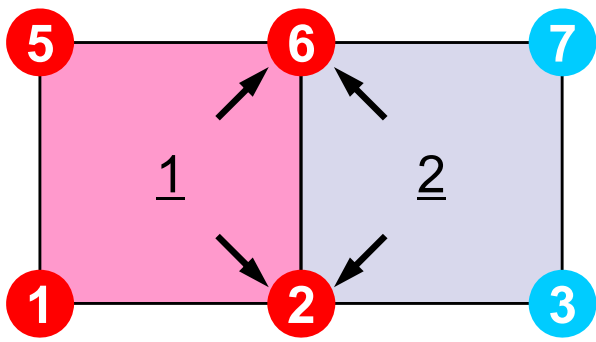
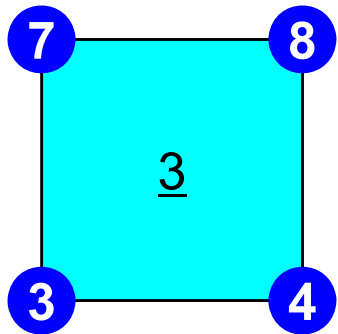
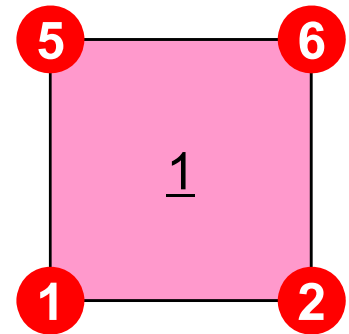
Values are defined on each node



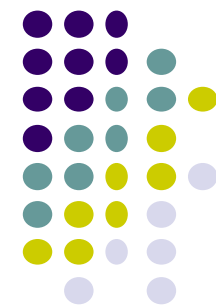
divide into two domains by “node-based” manner, where number of “nodes (vertices)” are balanced.

Local information is not enough for matrix assembling.

Information of overlapped elements and connected nodes are required for matrix assembling on boundary nodes.





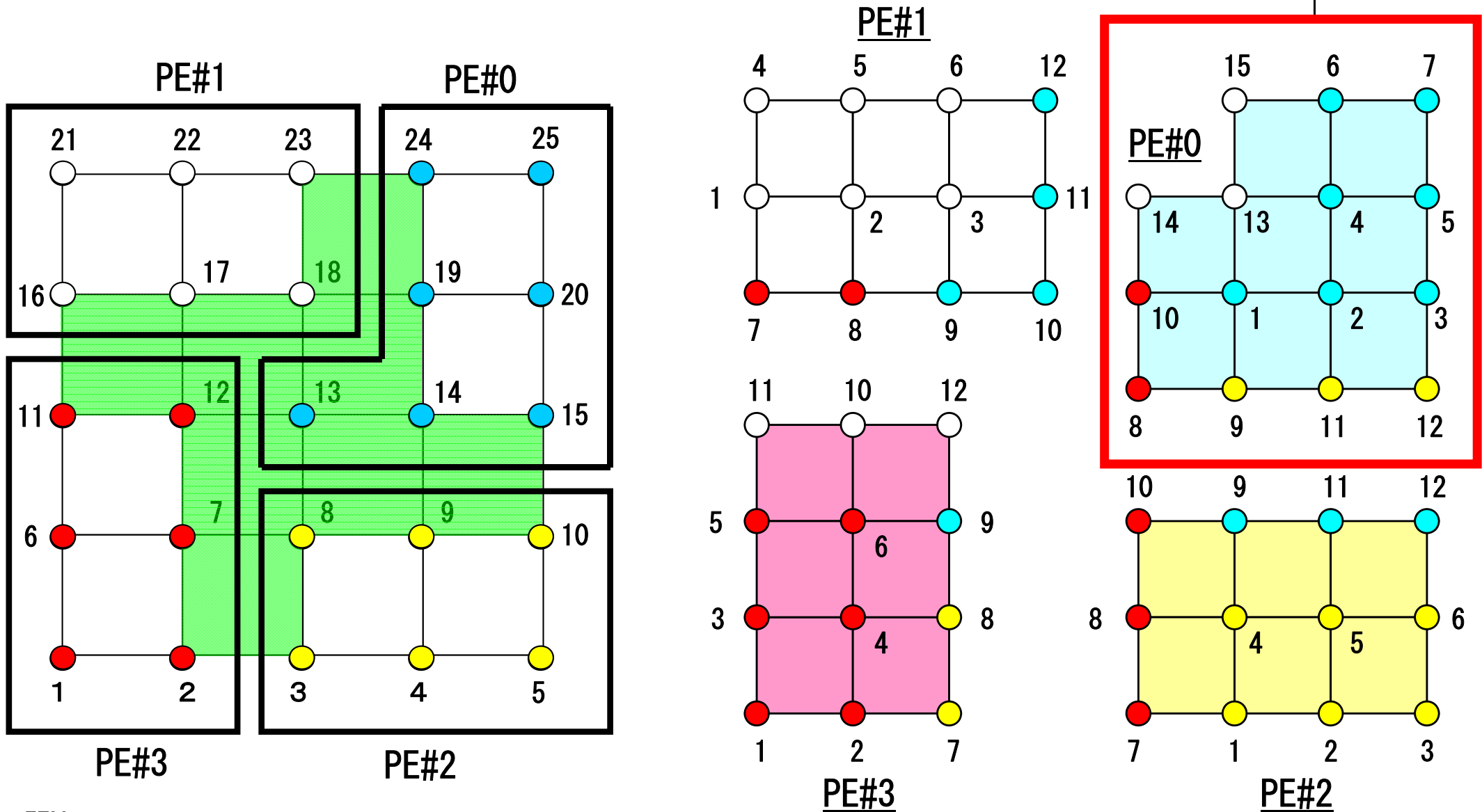


# Local Data of Parallel FEM

- **Node-based partitioning for IC/ILU type preconditioning methods**
- Local data includes information for :
  - Nodes originally assigned to the partition/PE
  - Elements which include the nodes : Element-based operations (Matrix Assemble) are allowed for fluid/structure subsystems.
  - All nodes which form the elements but out of the partition
- Nodes are classified into the following 3 categories from the viewpoint of the message passing
  - **Internal nodes** originally assigned nodes
  - **External nodes** in the overlapped elements but out of the partition
  - **Boundary nodes** *external nodes* of other partition
- Communication table between partitions
- NO global information required except partition-to-partition connectivity

# Node-based Partitioning

internal nodes - elements - external nodes

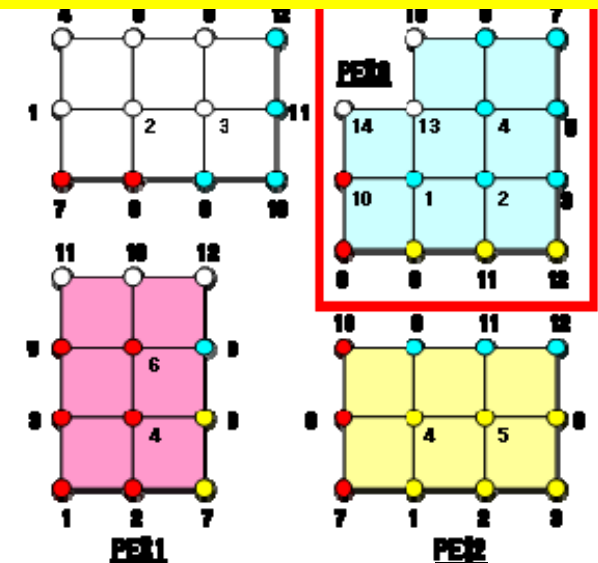
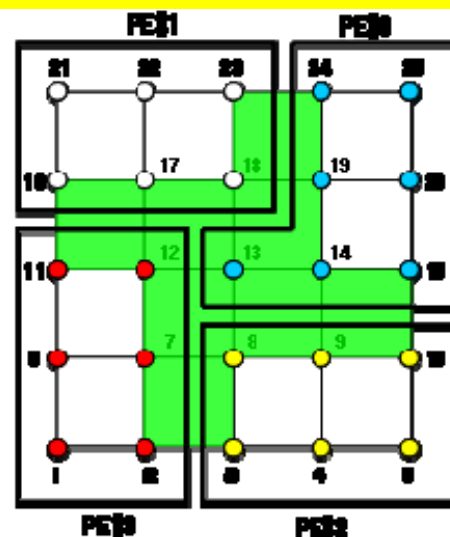
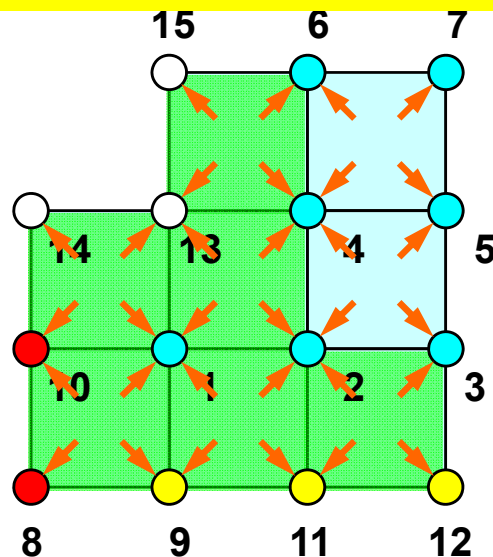


# Node-based Partitioning

internal nodes - elements - external nodes

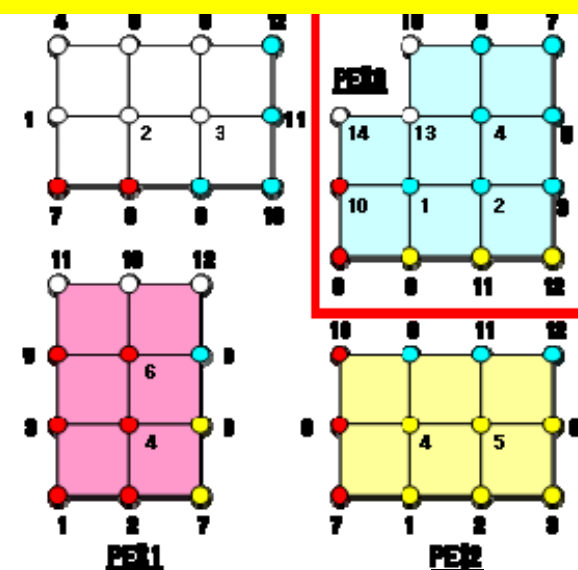
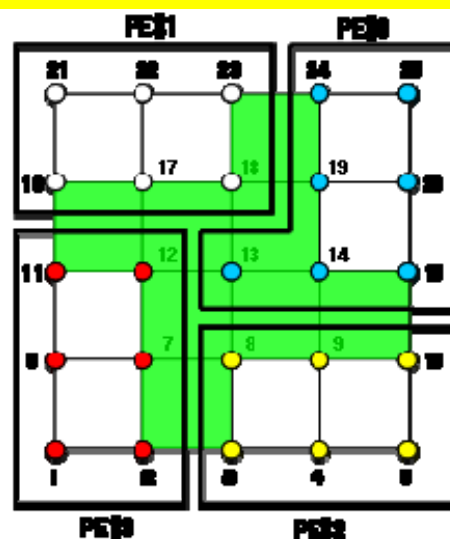
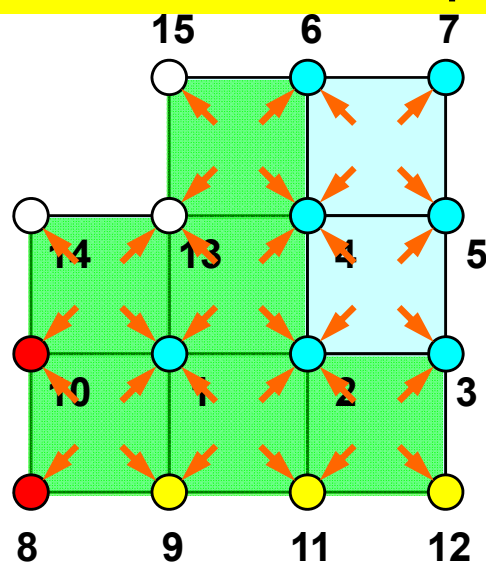


- Partitioned nodes themselves (Internal Nodes) 内点
- Elements which include Internal Nodes 内点を含む要素
- External Nodes included in the Elements 外点  
in overlapped region among partitions.
- Info of External Nodes are required for completely local element-based operations on each processor.



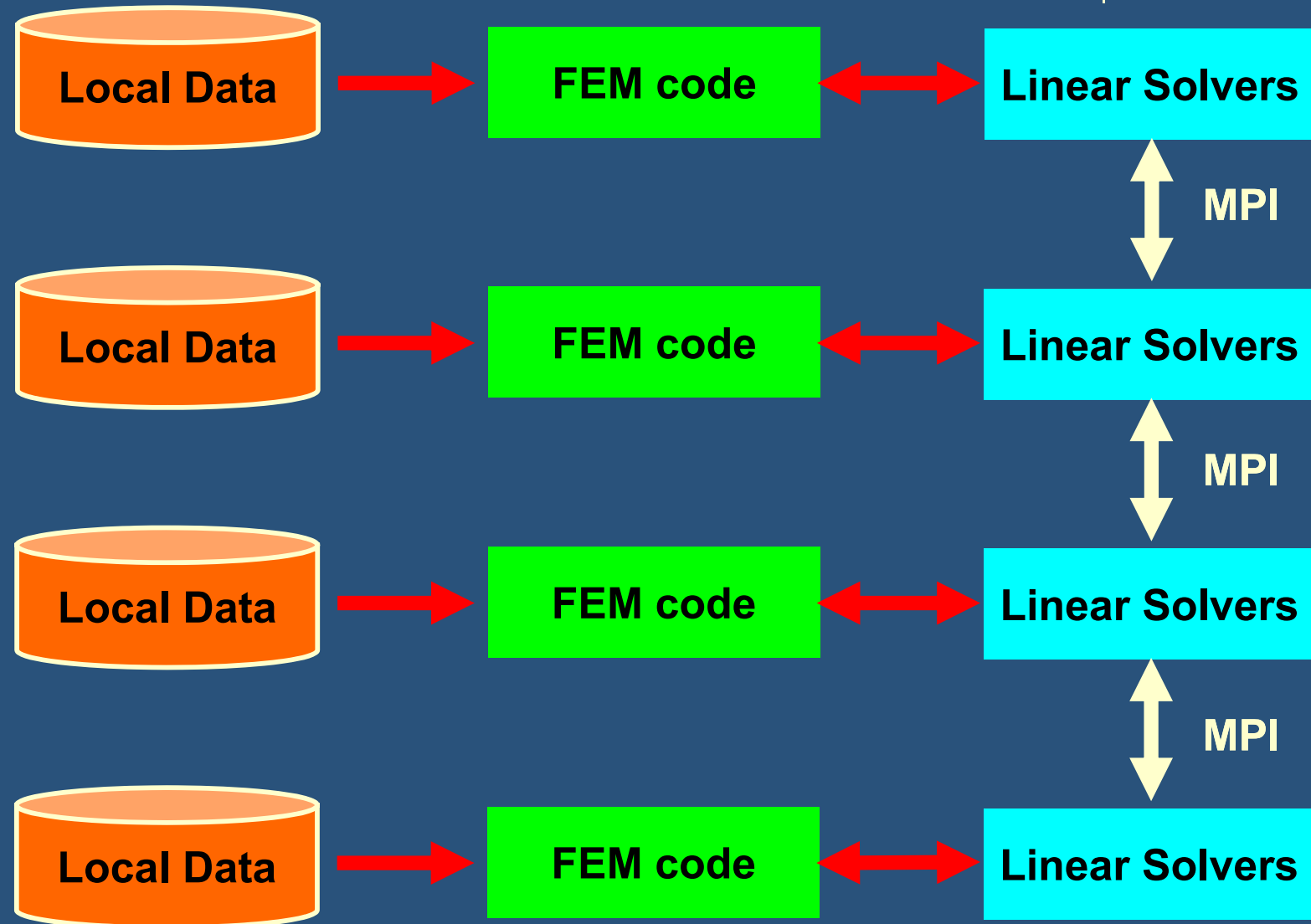
# We do not need communication during matrix assemble !!

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



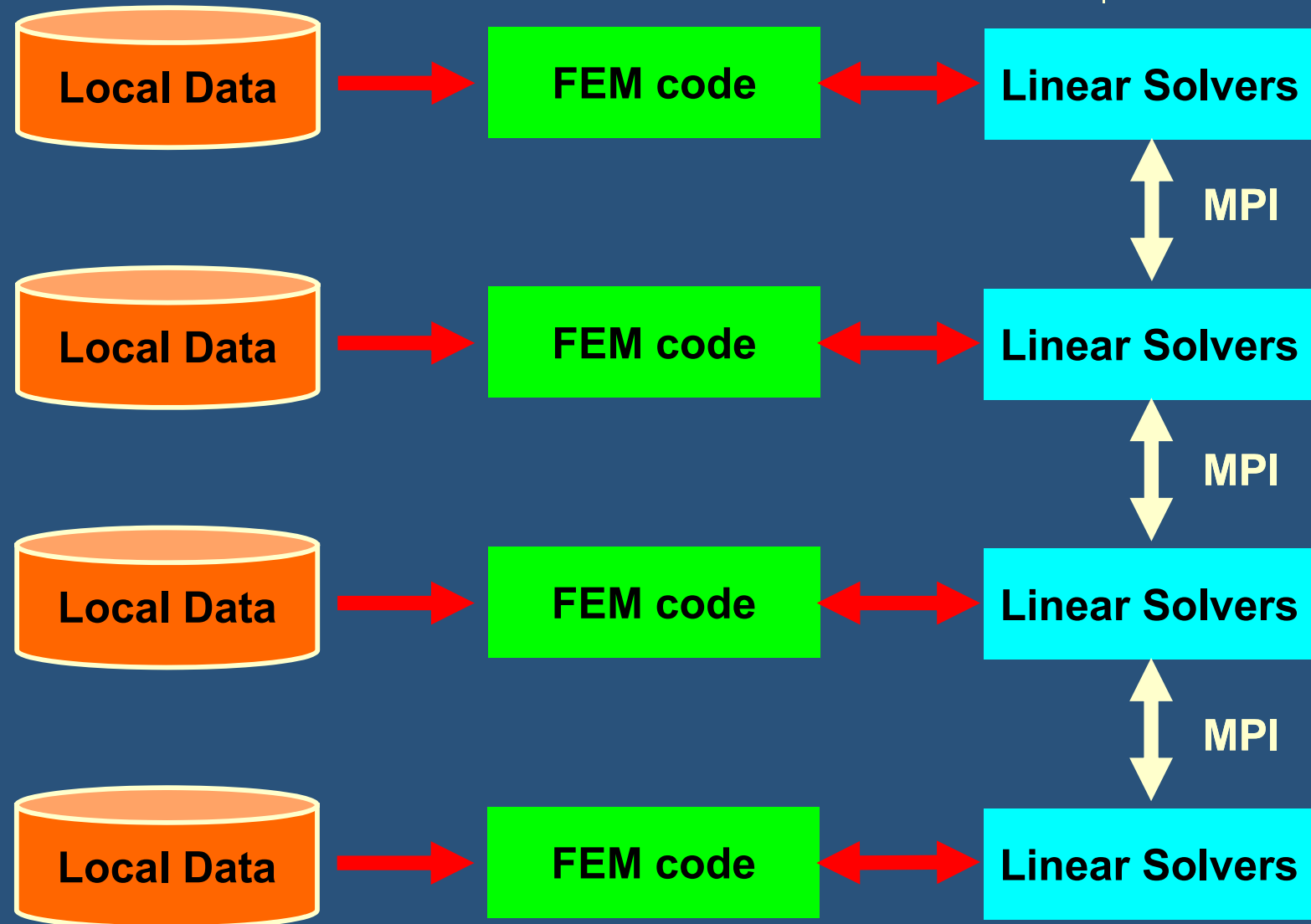
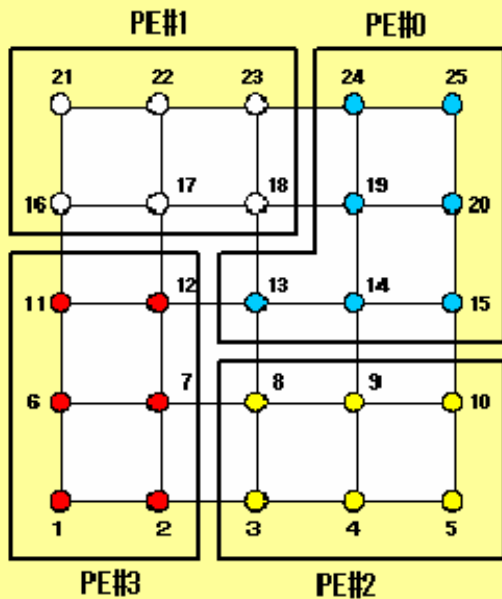
# Parallel Computing in FEM

## SPMD: Single-Program Multiple-Data



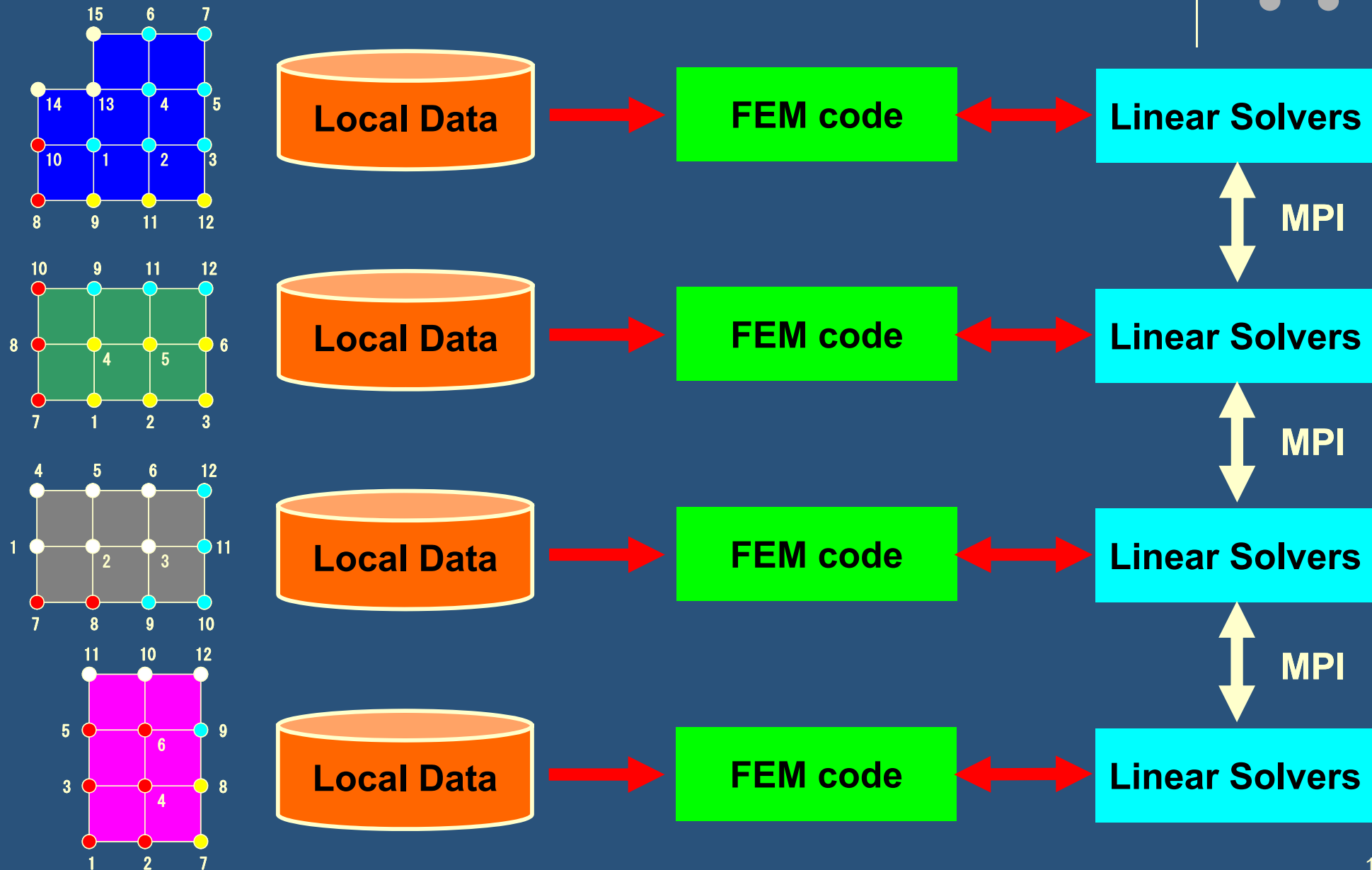
# Parallel Computing in FEM

## SPMD: Single-Program Multiple-Data



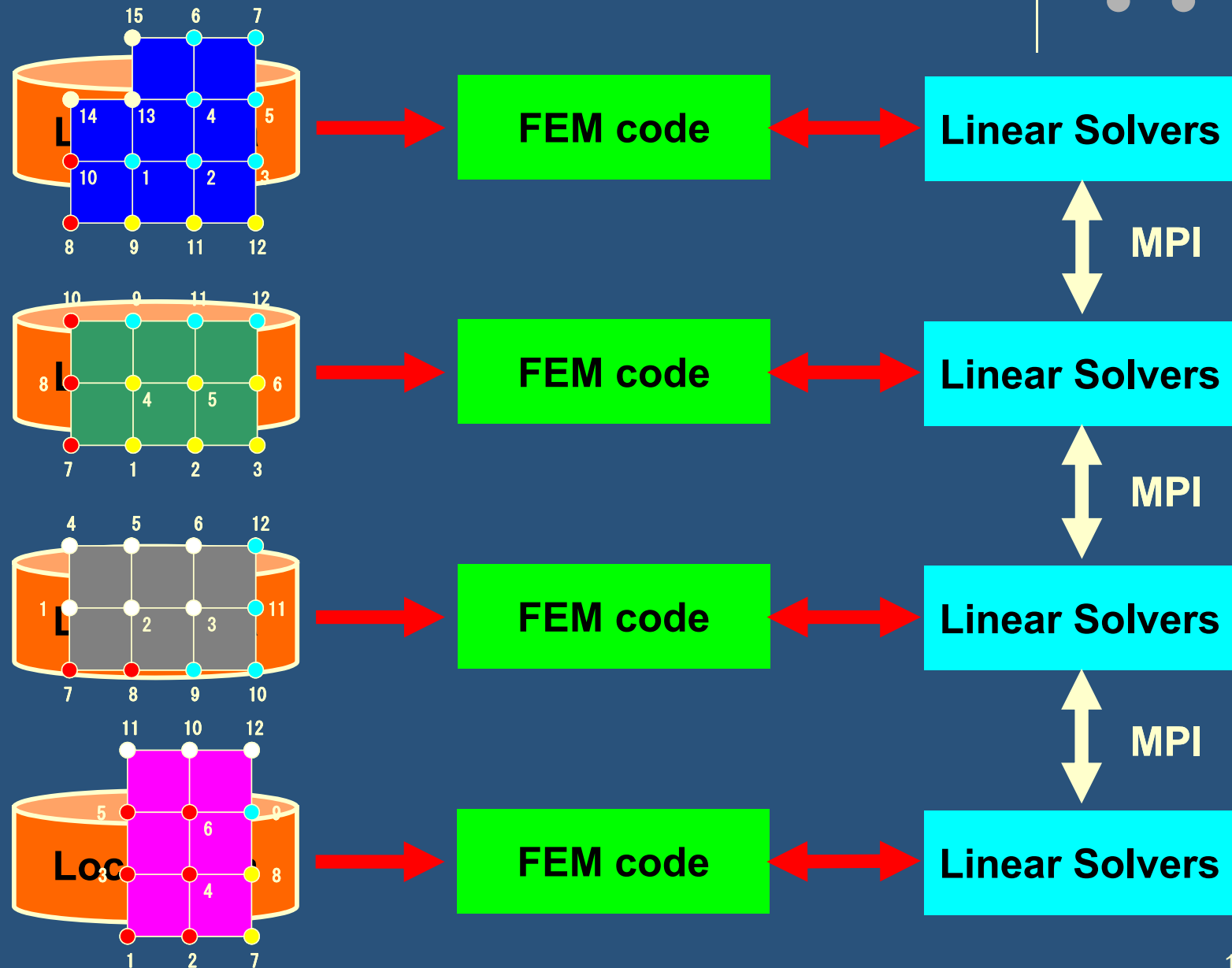
# Parallel Computing in FEM

## SPMD: Single-Program Multiple-Data



# Parallel Computing in FEM

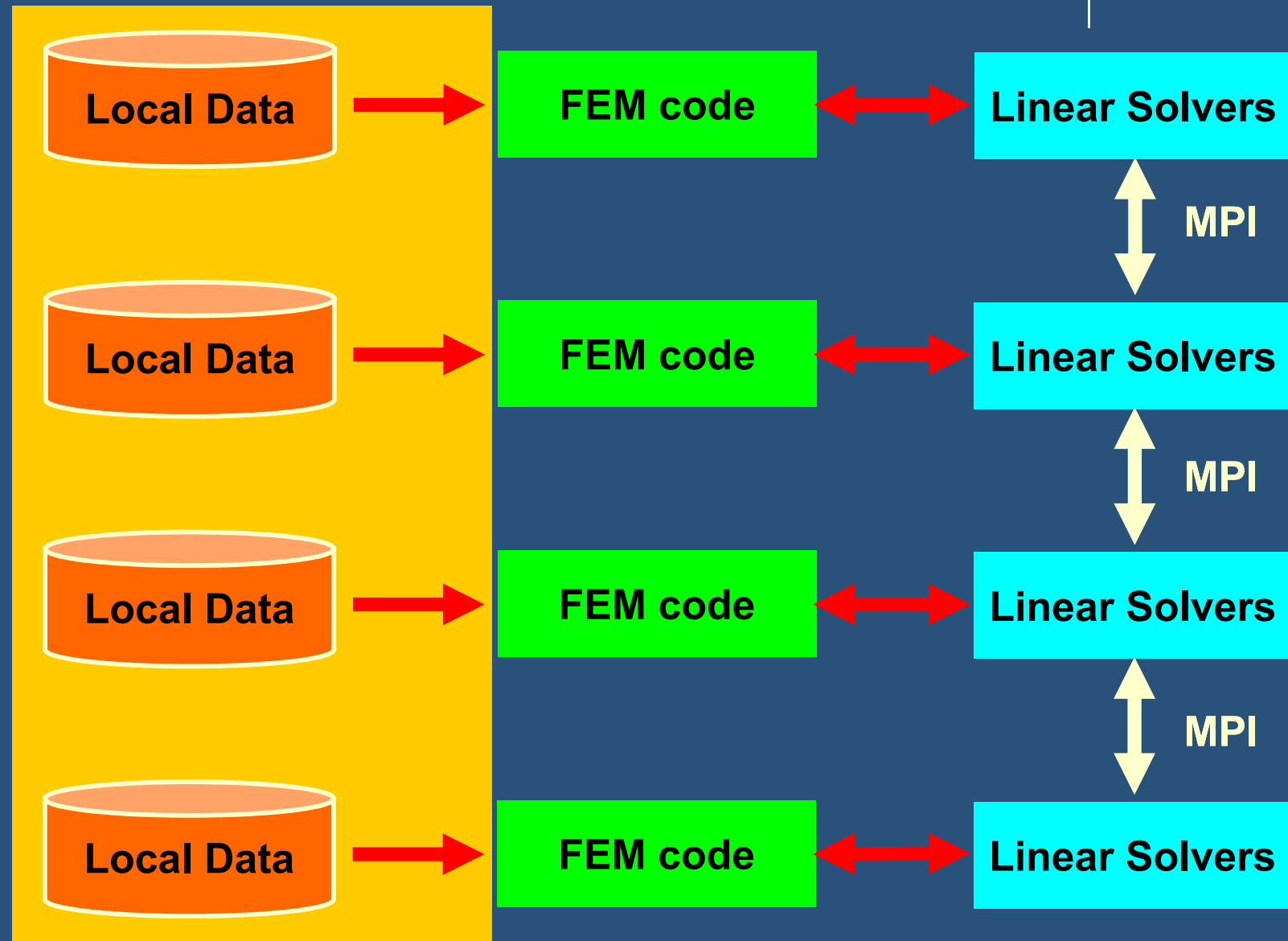
## SPMD: Single-Program Multiple-Data



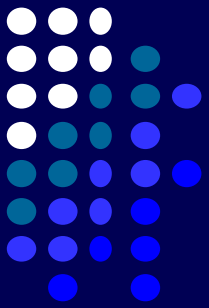


# Parallel Computing in FEM

## SPMD: Single-Program Multiple-Data

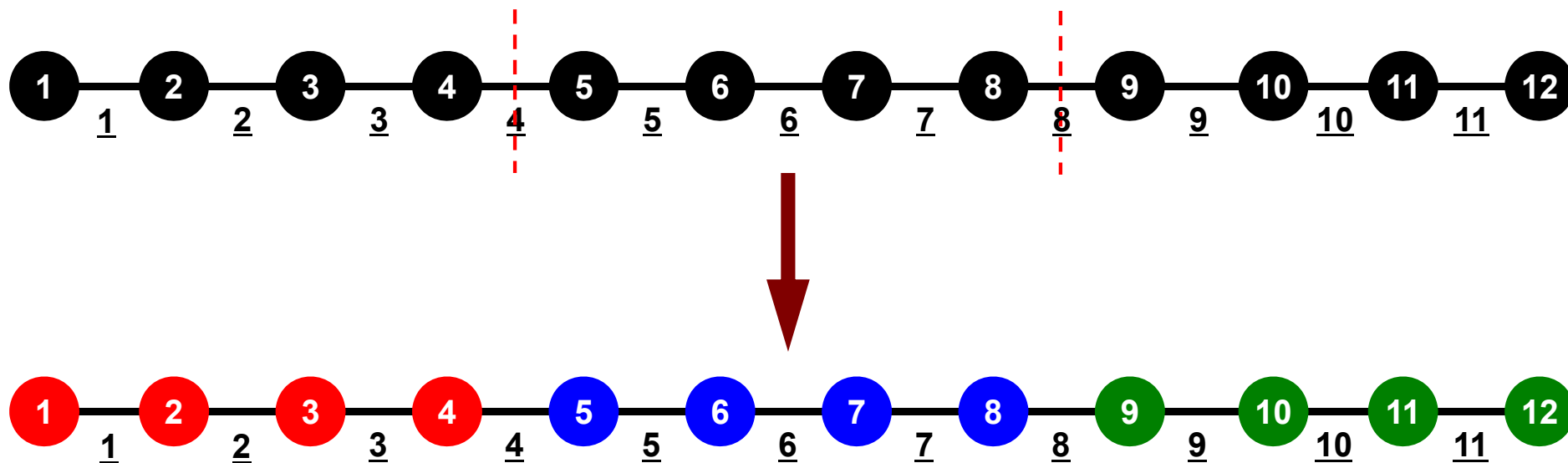


# What is Communications ?

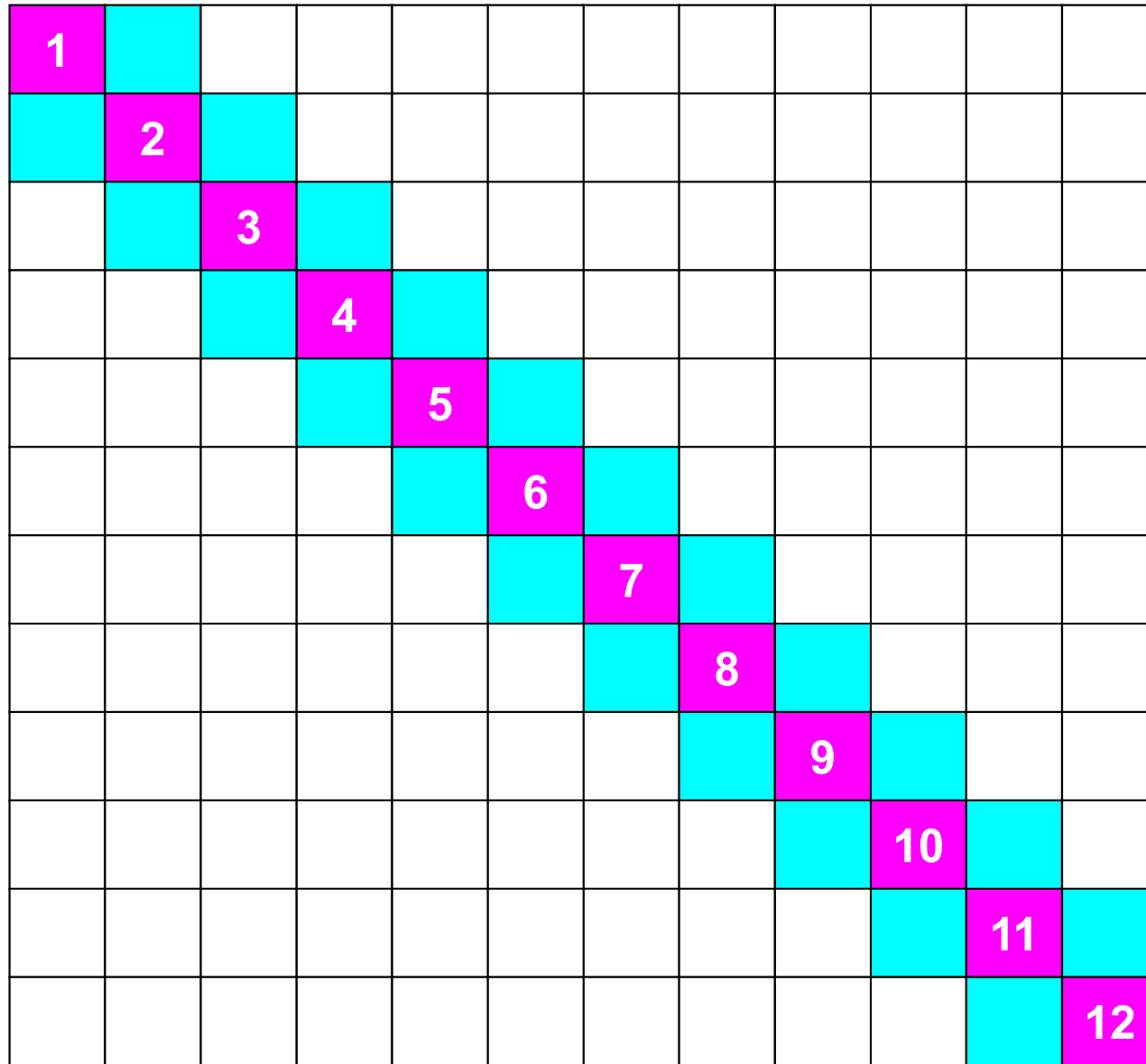


- to get information of “external nodes” from external partitions (local data)
- “Communication tables” contain the information

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

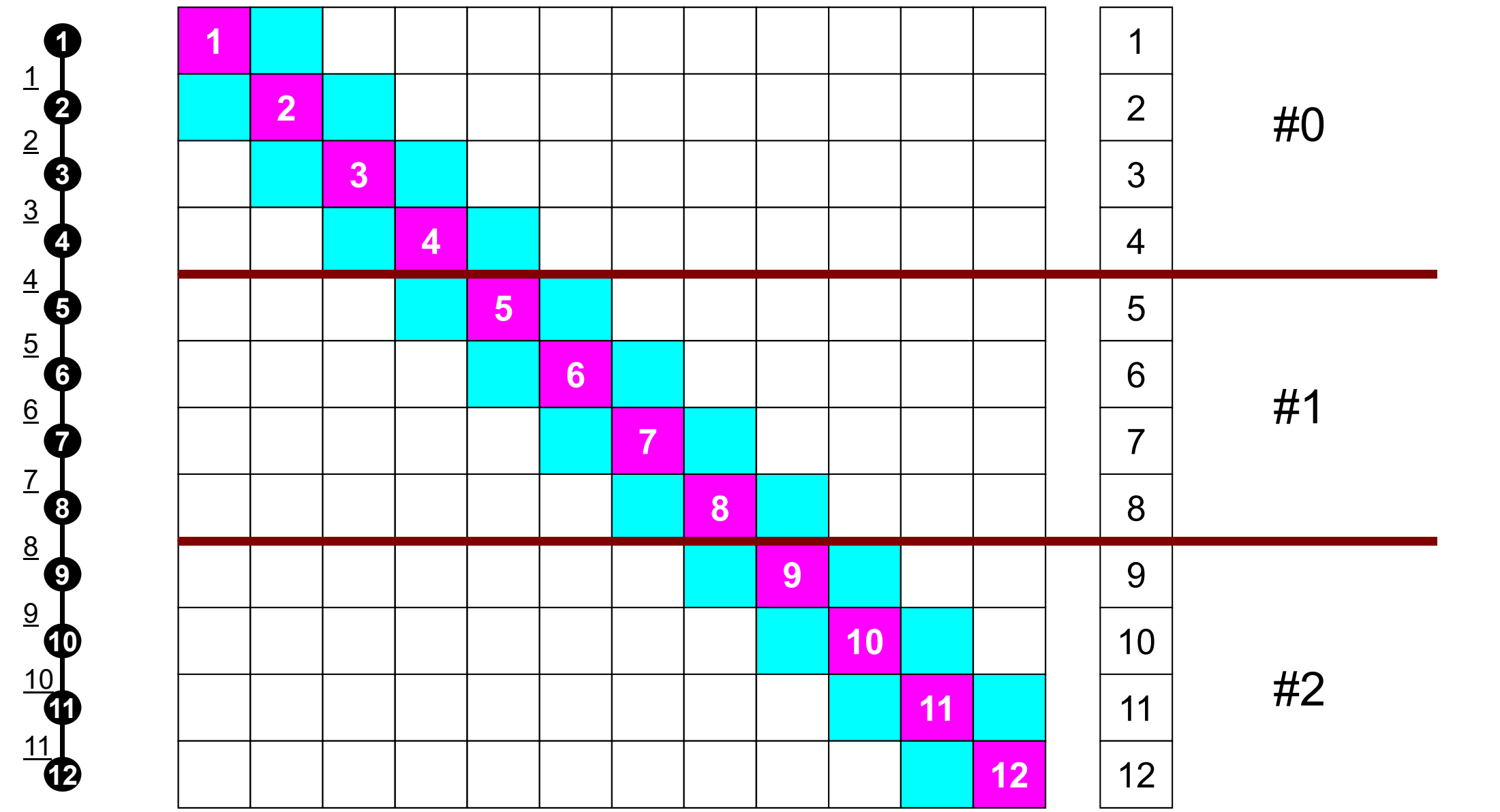


A vertical number line with numbers 1 through 12. The numbers 1 through 10 are placed inside black circles, while 11 and 12 are placed below the line. The numbers 1 through 10 are also underlined.



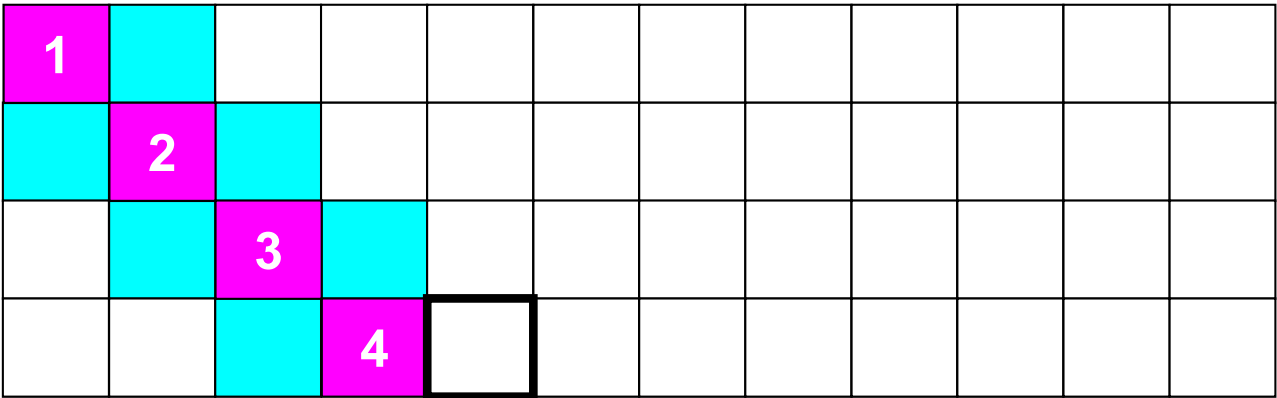
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| 3  |
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| 6  |
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| 9  |
| 10 |
| 11 |
| 12 |

# # “Internal Nodes” should be balanced



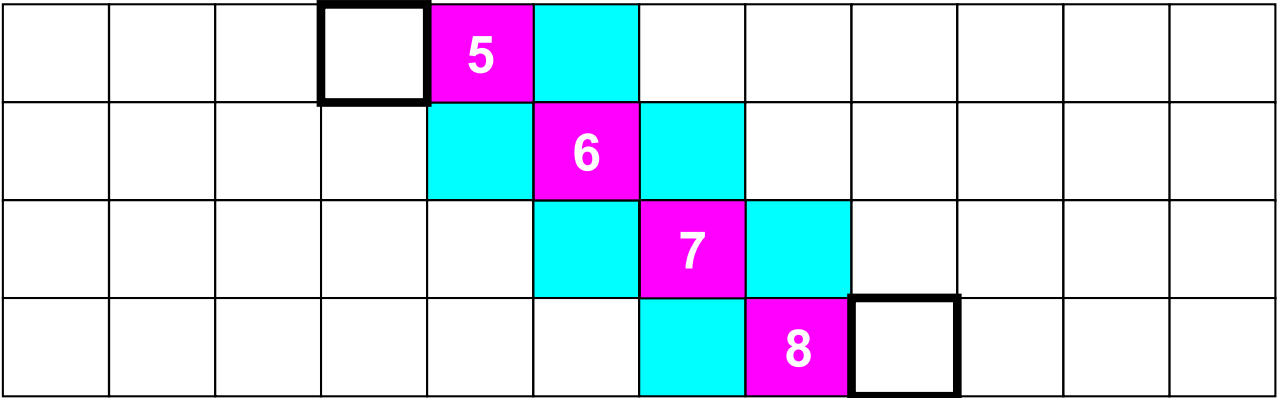
# Matrices are incomplete !

- 1
- 2
- 3
- 4



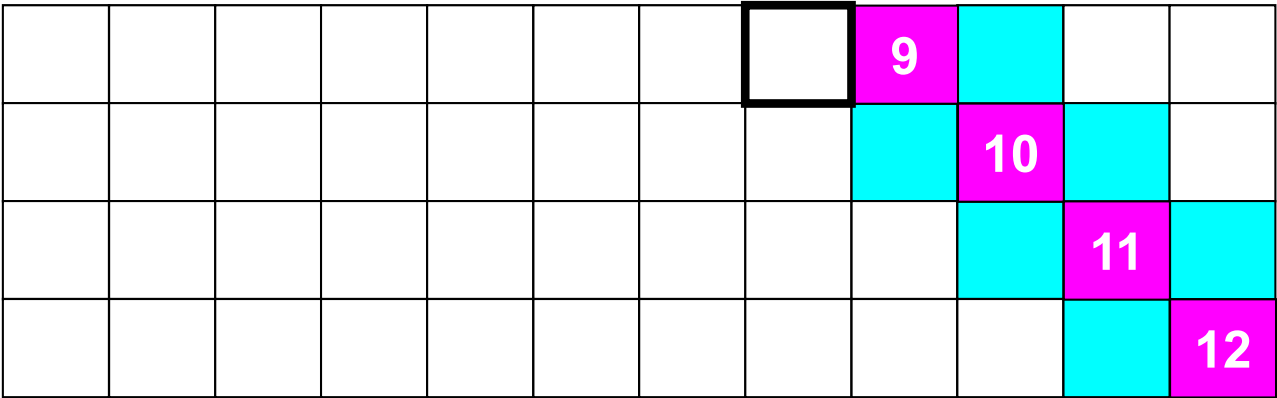
#0

- 5
- 6
- 7
- 8



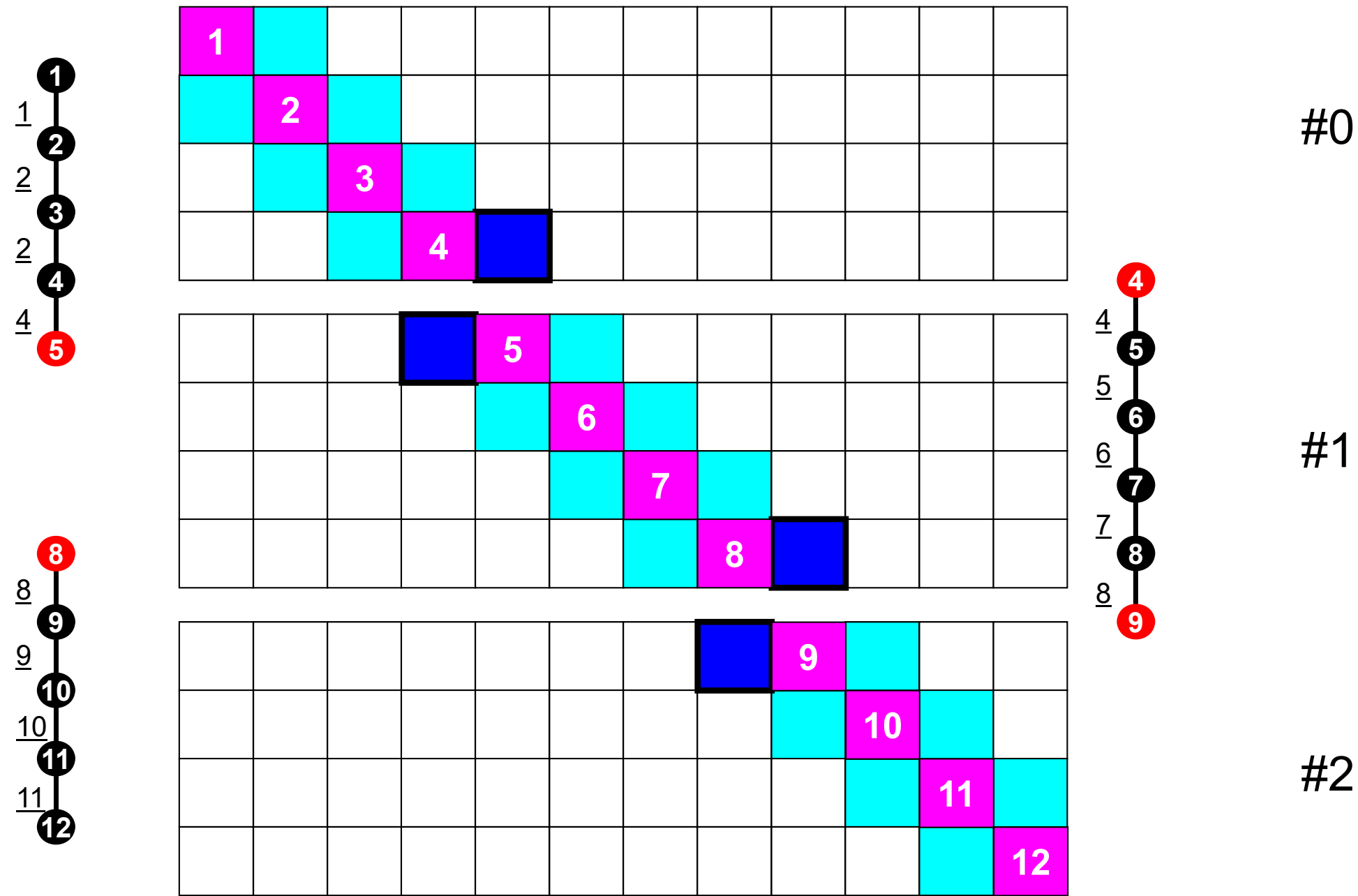
#1

- 9
- 10
- 11
- 12

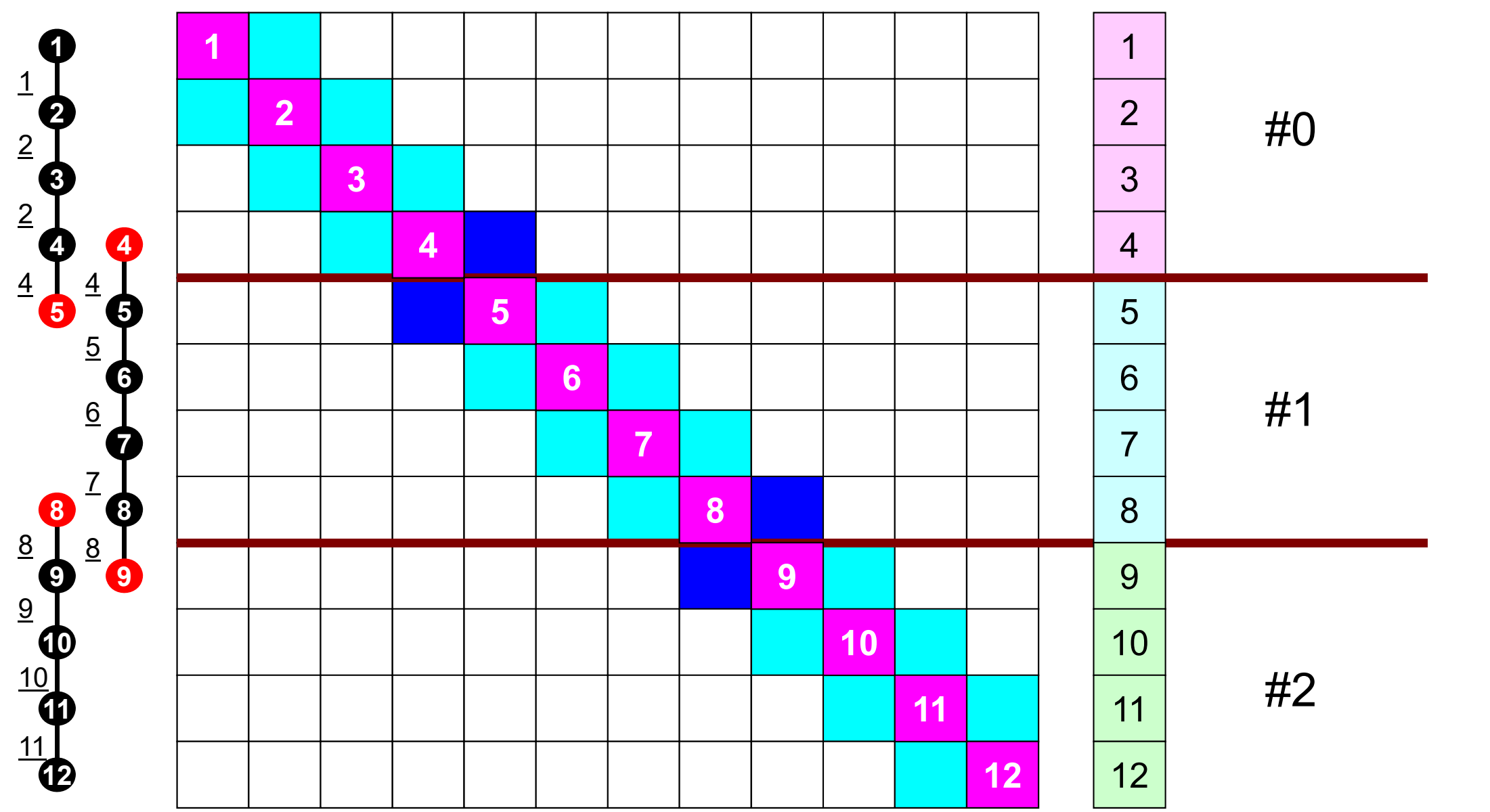


#2

# Connected Elements + External Nodes

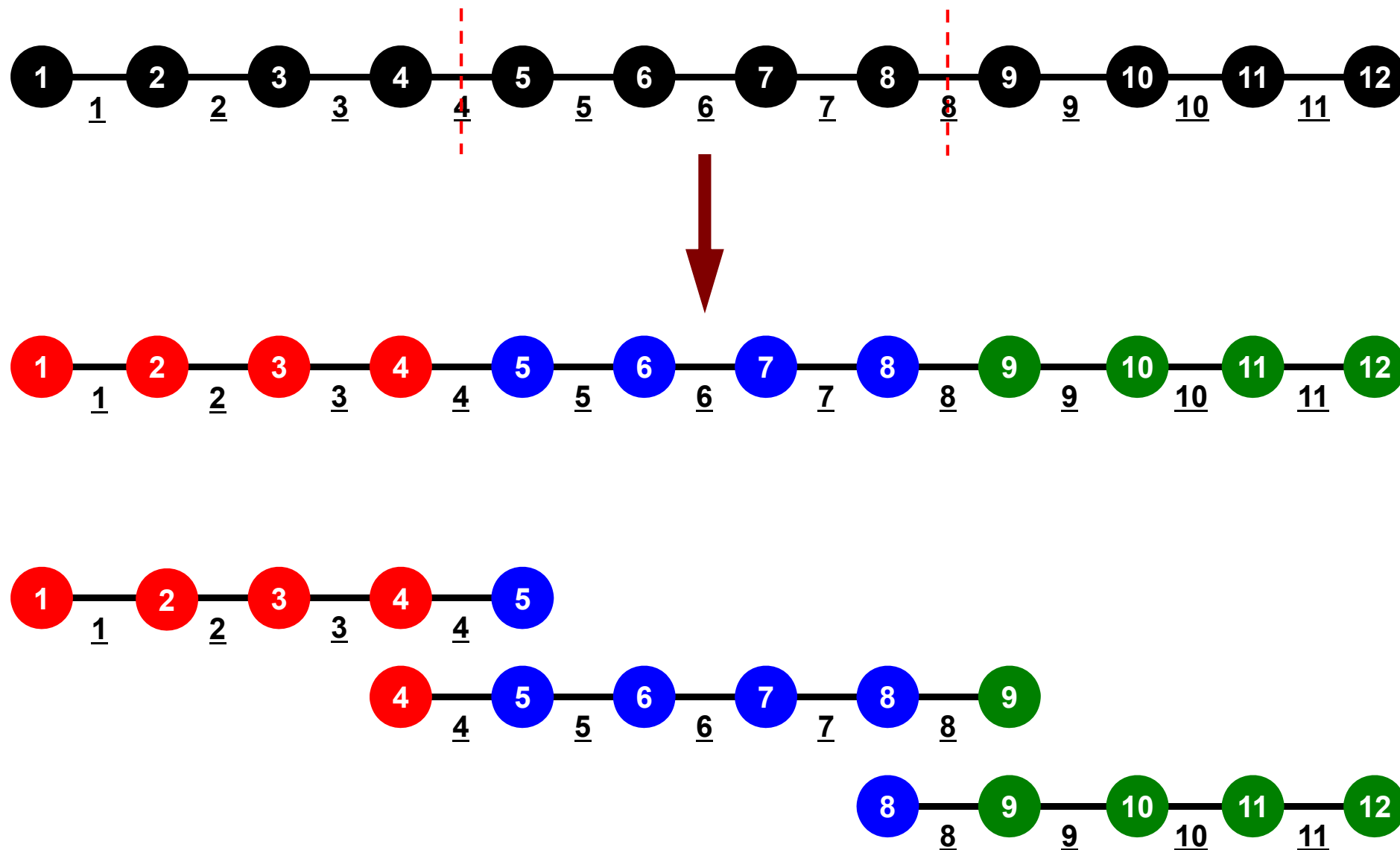


# 1D FEM: 12 nodes/11 elem's/3 domains



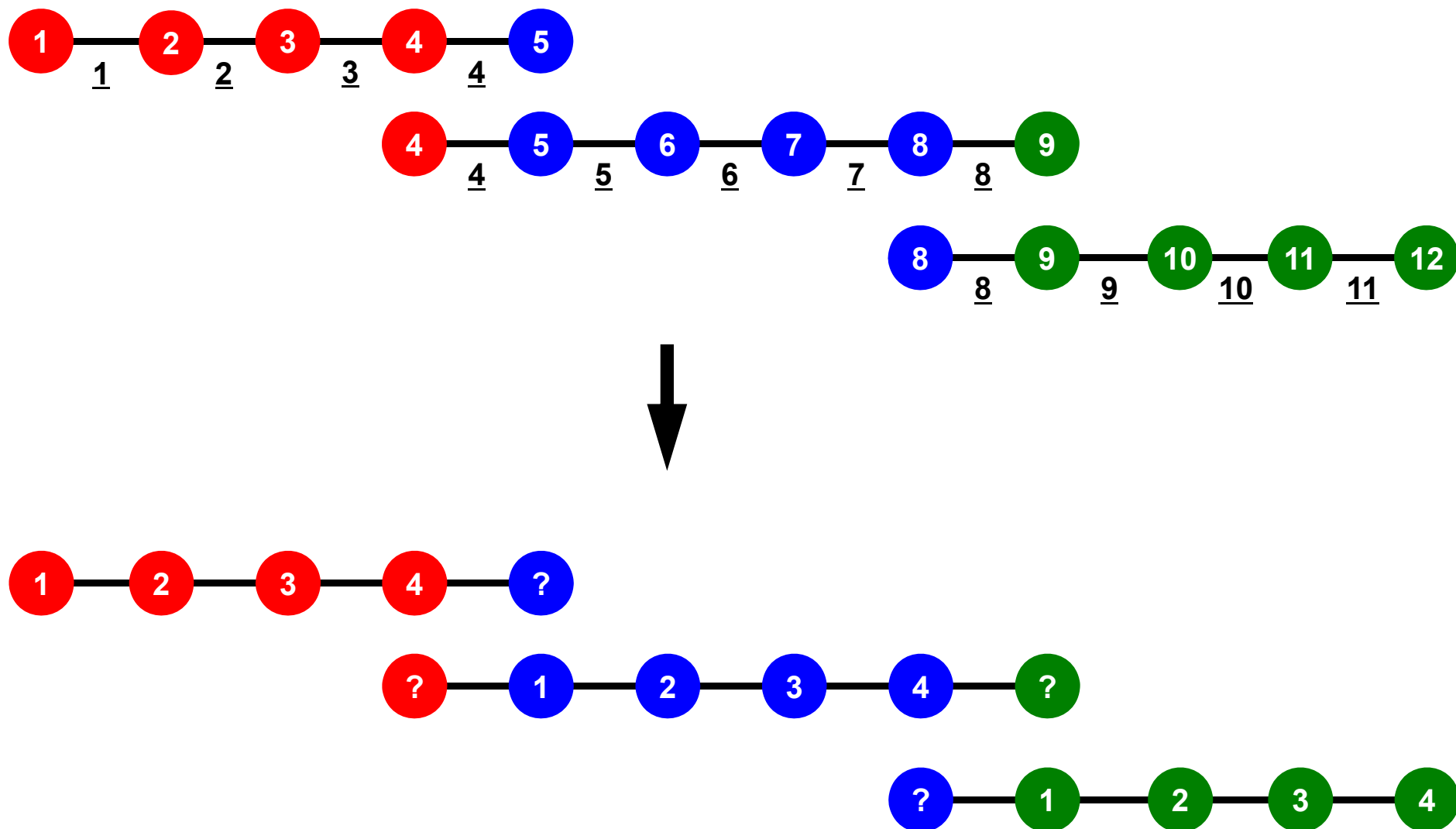


# 1D FEM: 12 nodes/11 elem's/3 domains



# Local Numbering for SPMD

Numbering of internal nodes is 1-N (0-N-1), same operations in serial program can be applied. How about numbering of external nodes ?

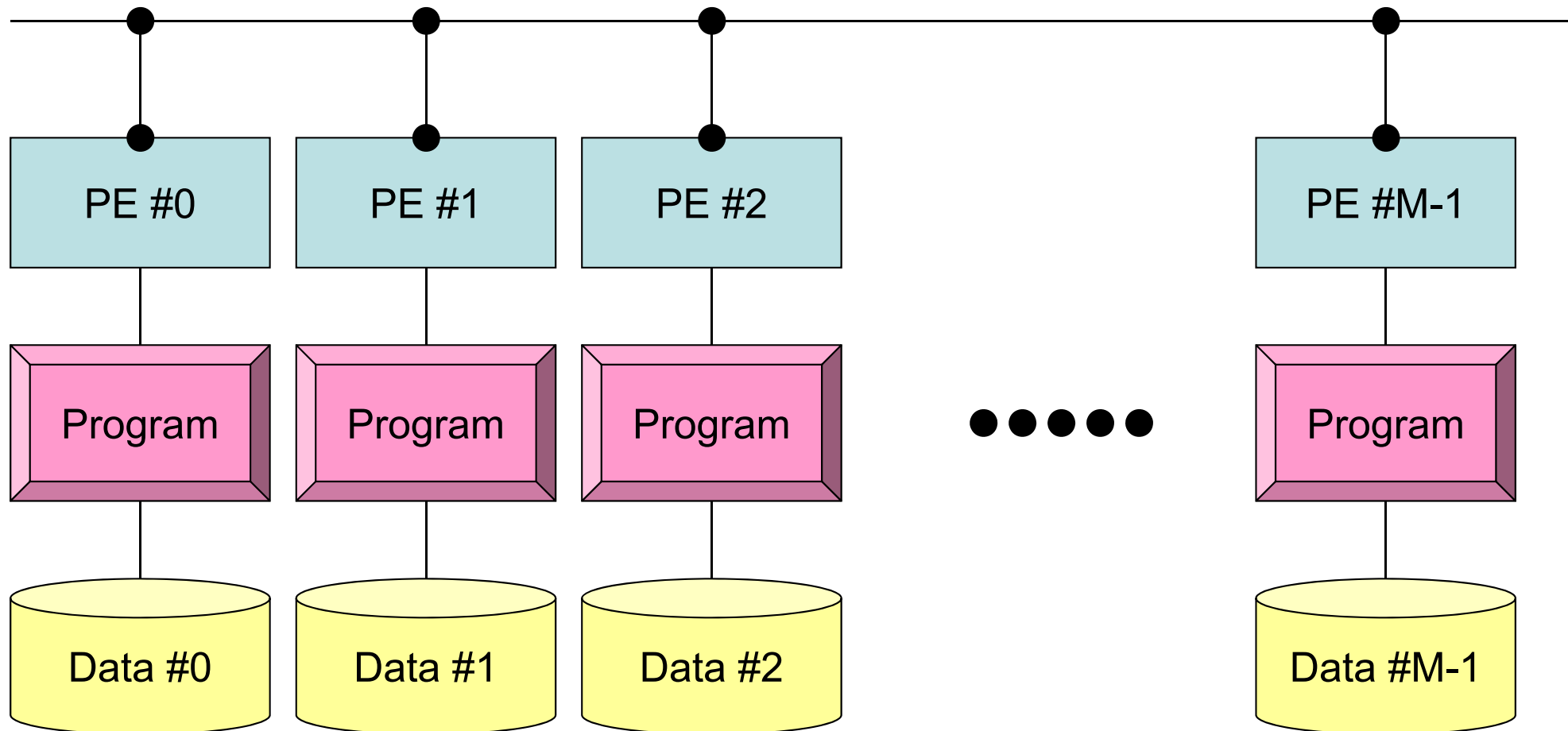


PE: Processing Element  
Processor, Domain, Process

# SPMD:

## Single Program Multiple Data

```
mpirun -np M <Program>
```



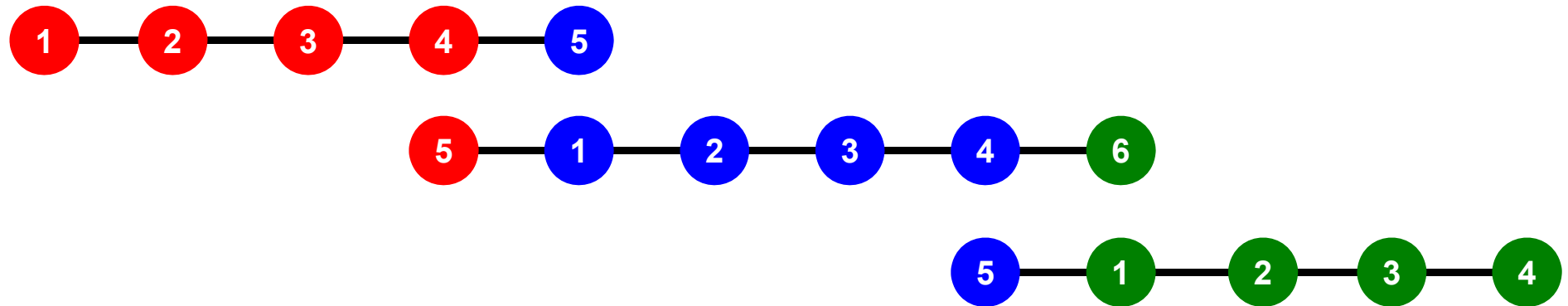
Each process does same operation for different data

Large-scale data is decomposed, and each part is computed by each process

It is ideal that parallel program is not different from serial one except communication.

# Local Numbering for SPMD

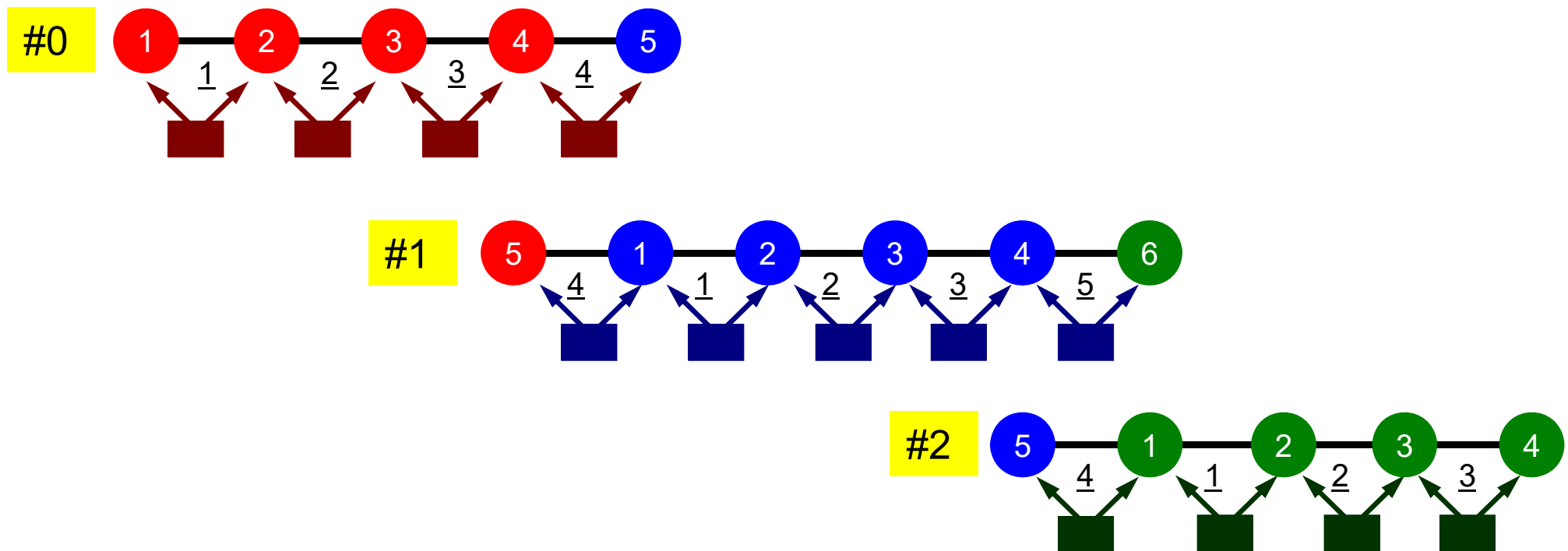
Numbering of external nodes:  $N+1$ ,  $N+2$  ( $N, N+1$ )



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

Integration on each element, element matrix  $\rightarrow$  global matrix

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



# 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

# Preconditioned CG Solver

```

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/Point Jacobi
- Parallel operations are required in
  - Dot Products
  - Mat-Vec. Multiplication
    - SpMV: Sparse Mat-Vec. Mult.

$$[\mathbf{M}] = \begin{bmatrix} D_1 & 0 & \dots & 0 & 0 \\ 0 & D_2 & & 0 & 0 \\ \dots & & \dots & & \dots \\ 0 & 0 & & D_{N-1} & 0 \\ 0 & 0 & \dots & 0 & D_N \end{bmatrix}$$

# 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}      DAXPY: double a{x} plus {y}
!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
```

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| 12 |



# 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
```

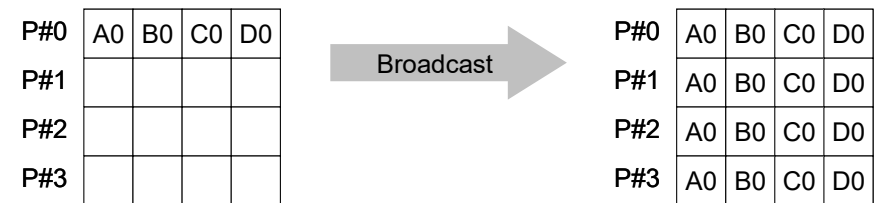
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| 12 |

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

➔  
Reduce

|     |          |          |          |          |
|-----|----------|----------|----------|----------|
| P#0 | op.A0-A3 | op.B0-B3 | op.C0-C3 | op.D0-D3 |
| P#1 |          |          |          |          |
| P#2 |          |          |          |          |
| P#3 |          |          |          |          |

- # Fortran



|     |    |    |    |    |            |     |          |          |          |          |
|-----|----|----|----|----|------------|-----|----------|----------|----------|----------|
| P#0 | A0 | B0 | C0 | D0 | All reduce | P#0 | op.A0-A3 | op.B0-B3 | op.C0-C3 | op.D0-D3 |
| P#1 | A1 | B1 | C1 | D1 |            | 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 |

- **call MPI\_ALLREDUCE**

|   |                 |        |   |                                           |
|---|-----------------|--------|---|-------------------------------------------|
| - | <u>sendbuf</u>  | choice | I | starting address of send buffer           |
| - | <u>recvbuf</u>  | choice | 0 | starting address receive buffer           |
|   |                 |        |   | type is defined by " <u>datatype</u> "    |
| - | <u>count</u>    | I      | I | number of elements in send/recv buffer    |
| - | <u>datatype</u> | I      | I | data type of elements in send/recv buffer |
| - | <u>op</u>       | I      | I | reduce operation                          |
| - | <u>comm</u>     | I      | I | commuinicator                             |
| - | <u>ierr</u>     | I      | 0 | completion code                           |

# “op” of MPI\_Reduce/Allreduce

```
call MPI_REDUCE
```

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

- **MPI\_MAX, MPI\_MIN** Max, Min
- **MPI\_SUM, MPI\_PROD** Summation, Product
- **MPI\_LAND** Logical AND

# Preconditioned CG Solver

```

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  check convergence  $|\mathbf{r}|$ 
end

```

- Preconditioning
  - Diagonal Scaling/Point Jacobi
- Parallel operations are required in
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  - Mat-Vec. Multiplication
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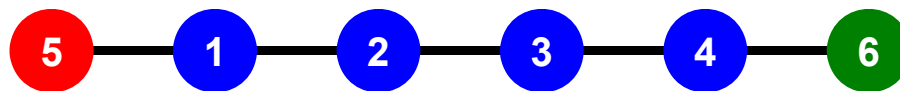
$$[\mathbf{M}] = \begin{bmatrix} D_1 & 0 & \dots & 0 & 0 \\ 0 & D_2 & & 0 & 0 \\ \dots & & \dots & & \dots \\ 0 & 0 & & D_{N-1} & 0 \\ 0 & 0 & \dots & 0 & D_N \end{bmatrix}$$

# Matrix-Vector Products

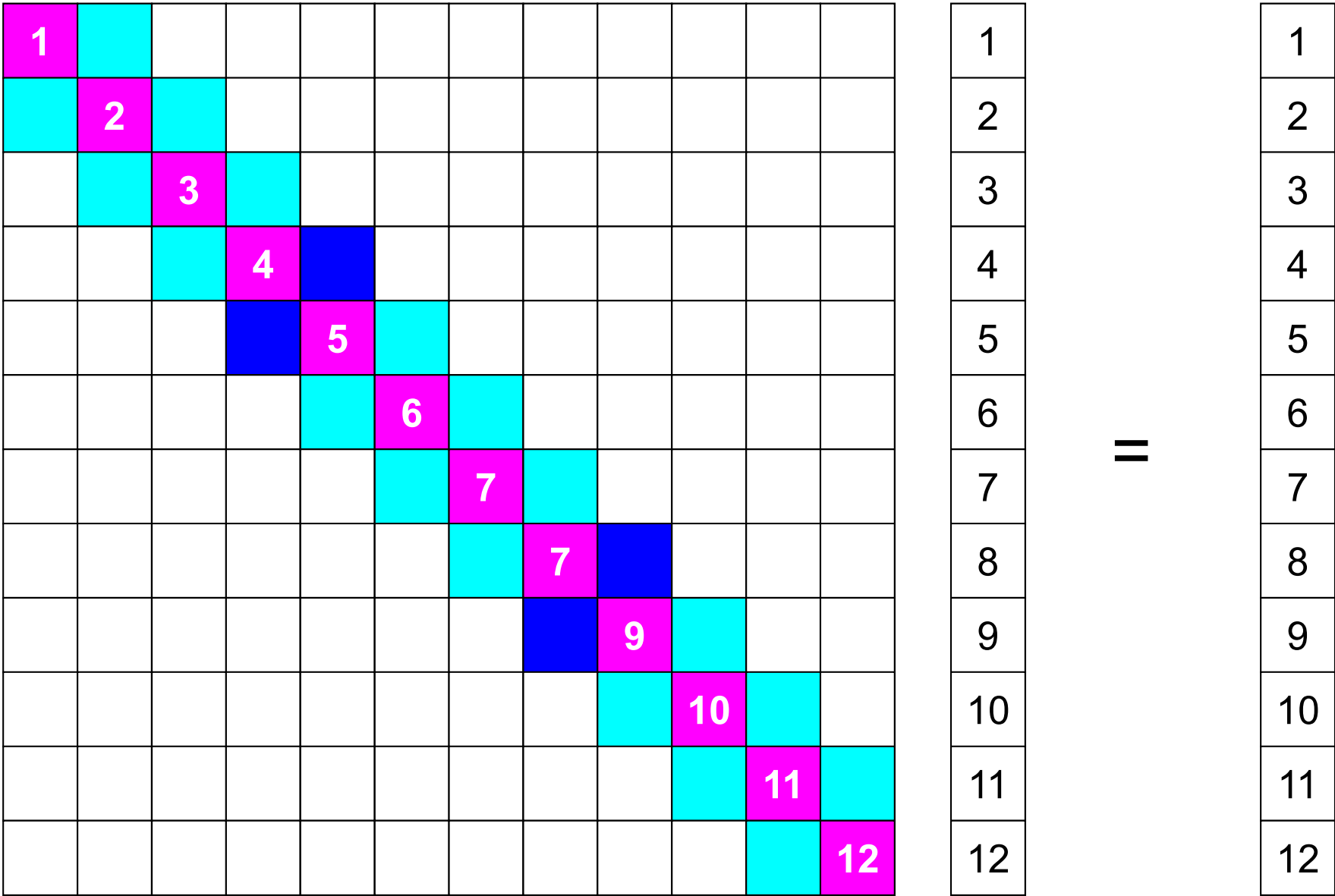
## Values at External Points: P-to-P Communication

```
!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
```

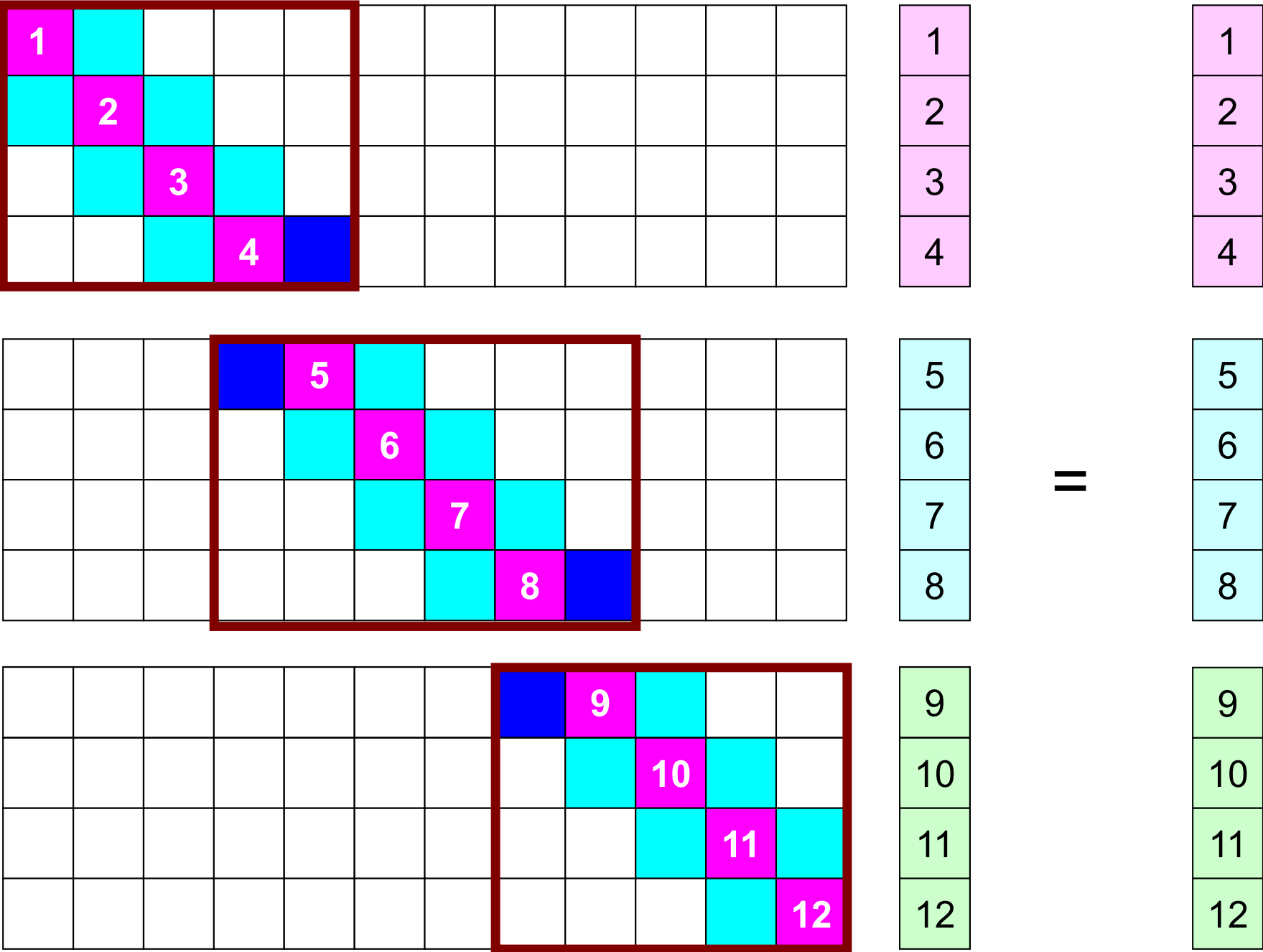


# Mat-Vec Products: Local Op. Possible

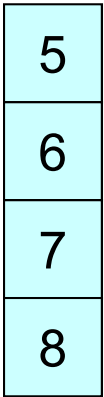
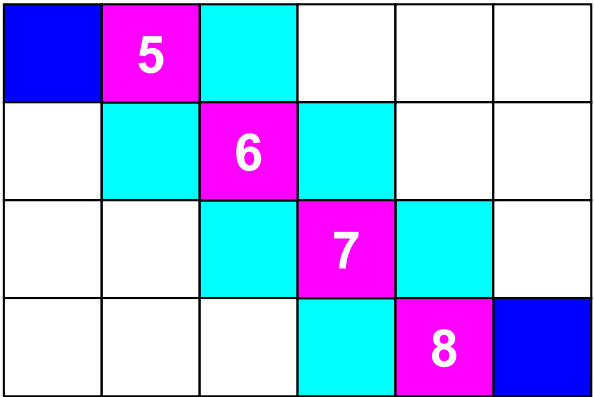
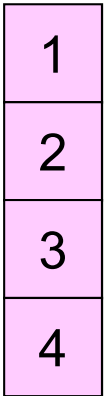
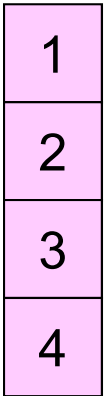
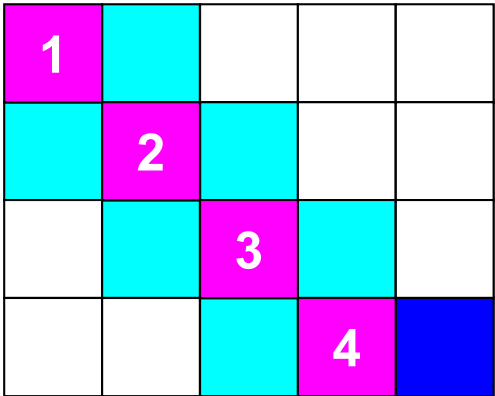




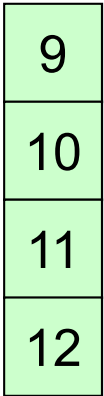
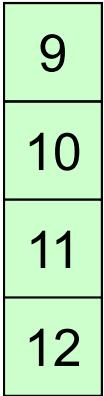
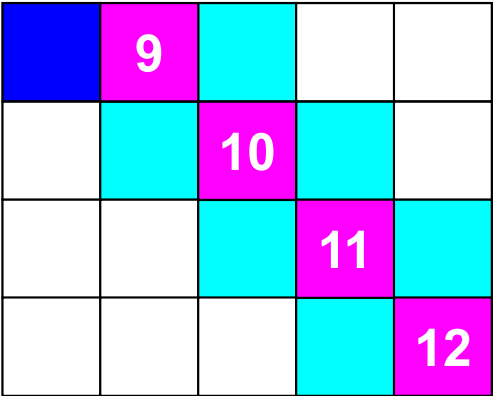
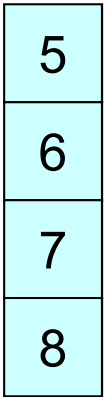
# Mat-Vec Products: Local Op. Possible



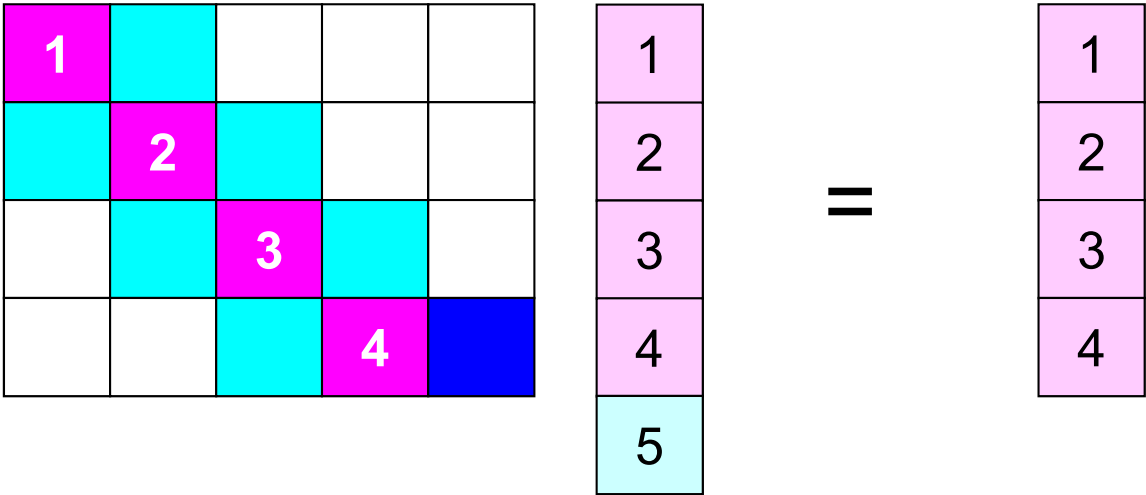
# Mat-Vec Products: Local Op. Possible



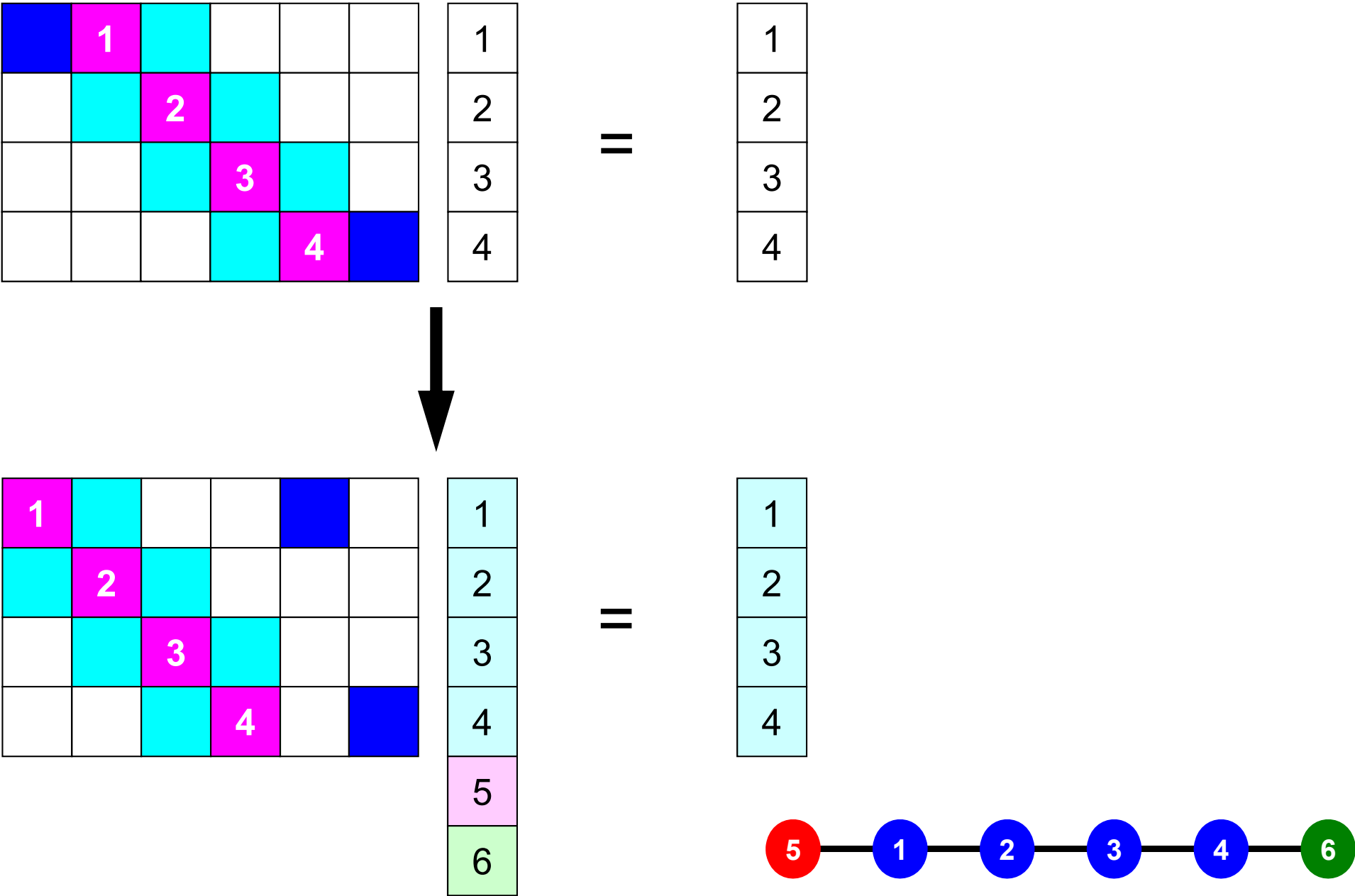
=



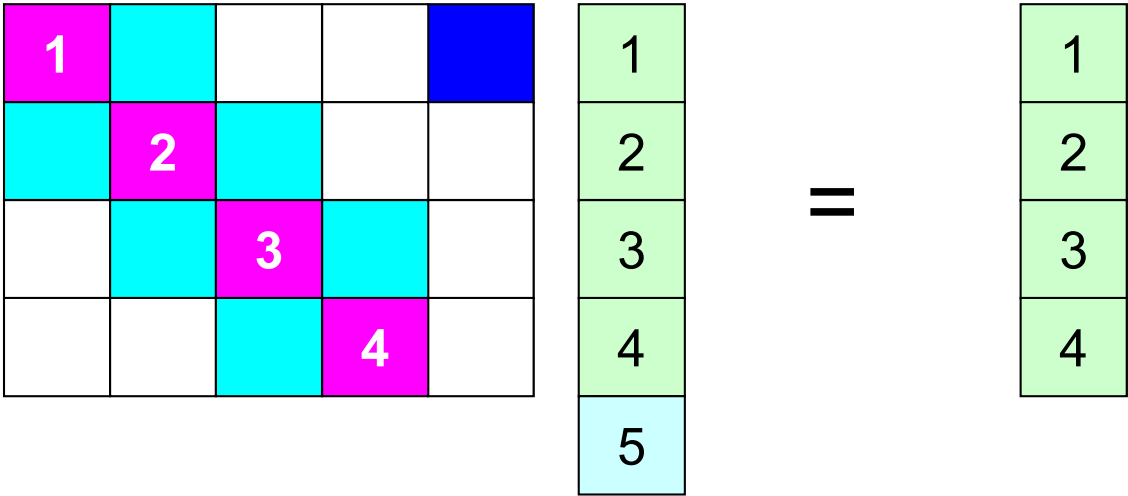
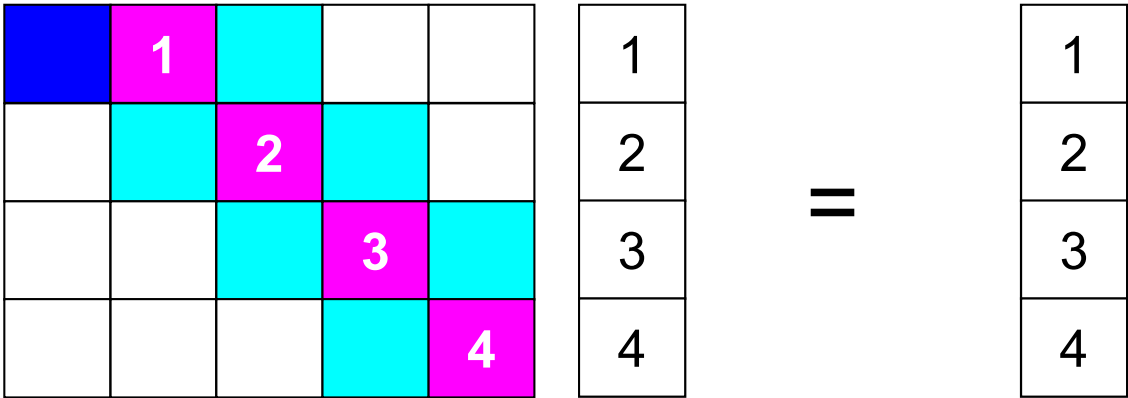
# Mat-Vec Products: Local Op. #0



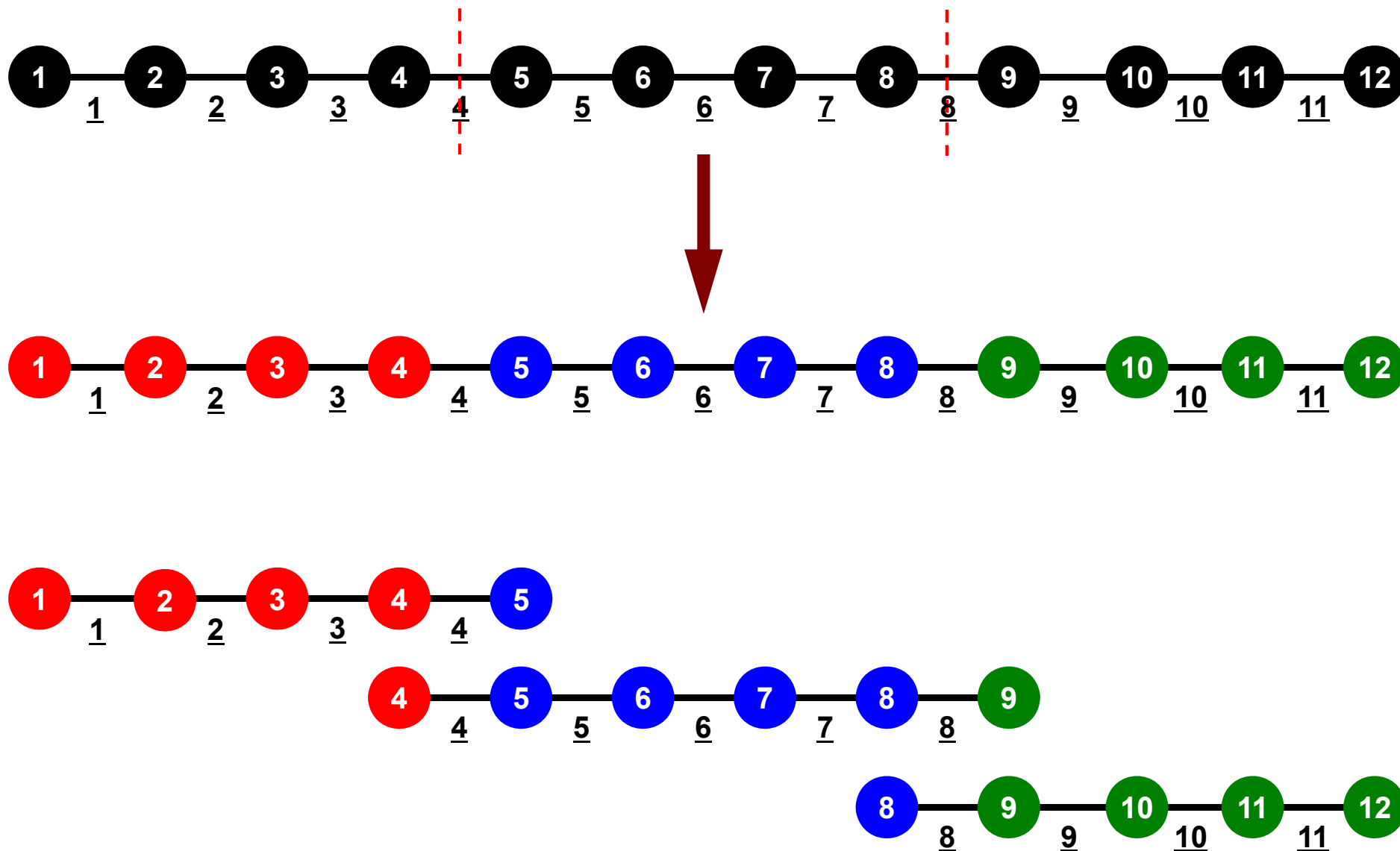
# Mat-Vec Products: Local Op. #1



# Mat-Vec Products: Local Op. #2

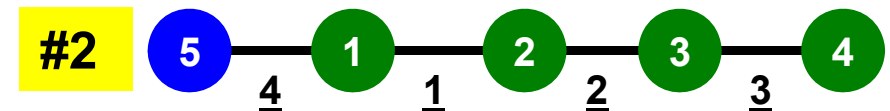
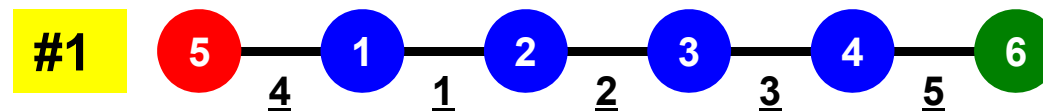
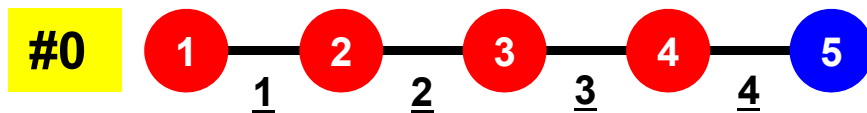


# 1D FEM: 12 nodes/11 elem's/3 domains



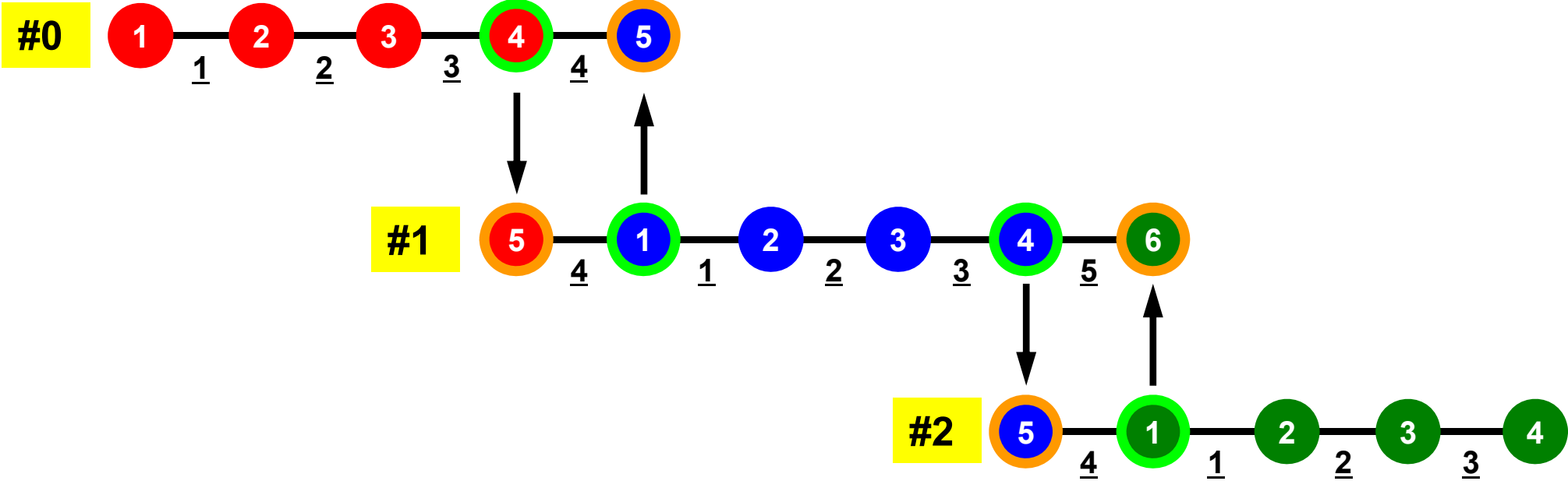
# 1D FEM: 12 nodes/11 elem's/3 domains

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



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

Internal/External Nodes



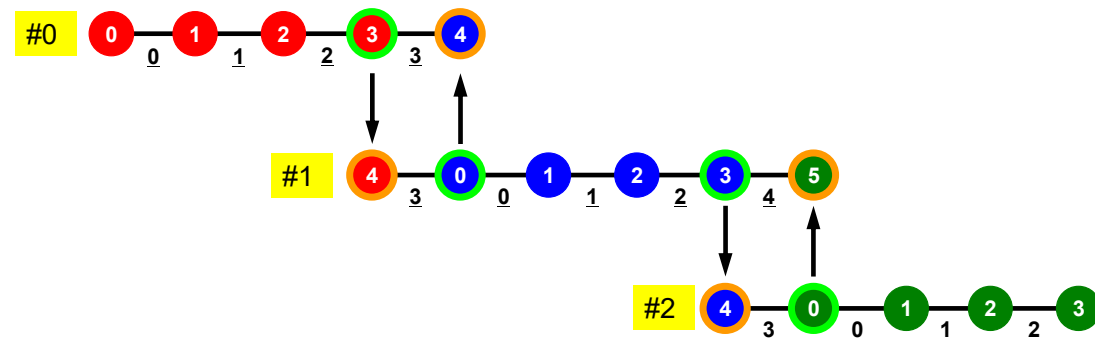


# What is Peer-to-Peer Communication ?

- Collective Communication
  - MPI\_Reduce, MPI\_Scatter/Gather etc.
  - Communications with all processes in the communicator
  - Application Area
    - BEM, Spectral Method, MD: global interactions are considered
    - Dot products, MAX/MIN: Global Summation & Comparison

- Peer-toPeer/Point-to-Point

- MPI\_Send, MPI\_Receive
- Communication with limited processes
  - Neighbors
- Application Area
  - FEM, FDM: Localized Method



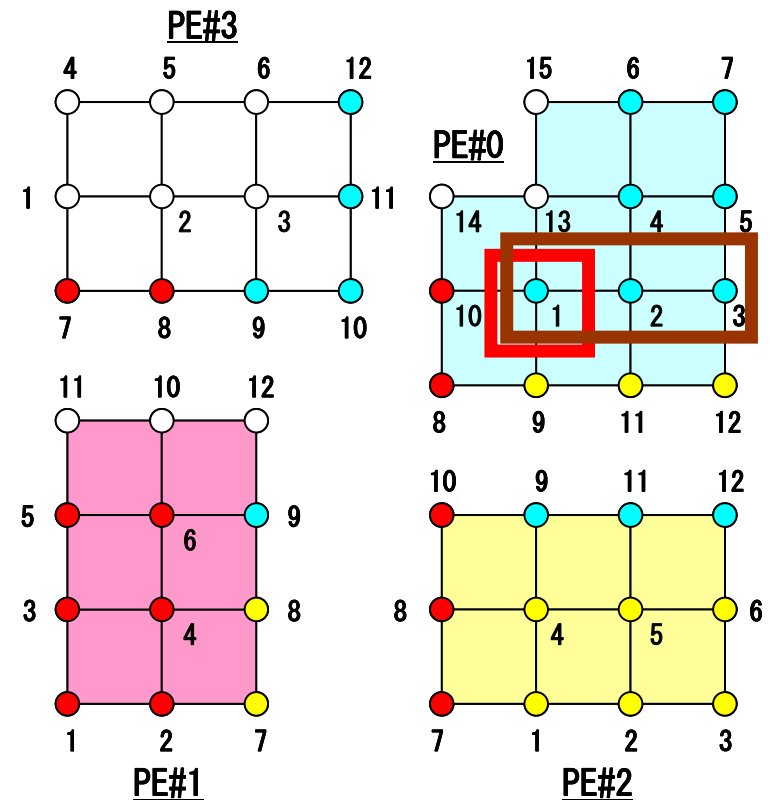
# SEND: sending from boundary nodes

Send continuous data to send buffer of neighbors

- **MPI\_Isend**

(**sendbuf**, **count**, **datatype**, **dest**, **tag**, **comm**, **request**)

- **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



# 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            0            communication request array used in MPI\_Waitall
  - **ierr**         I            0            completion code

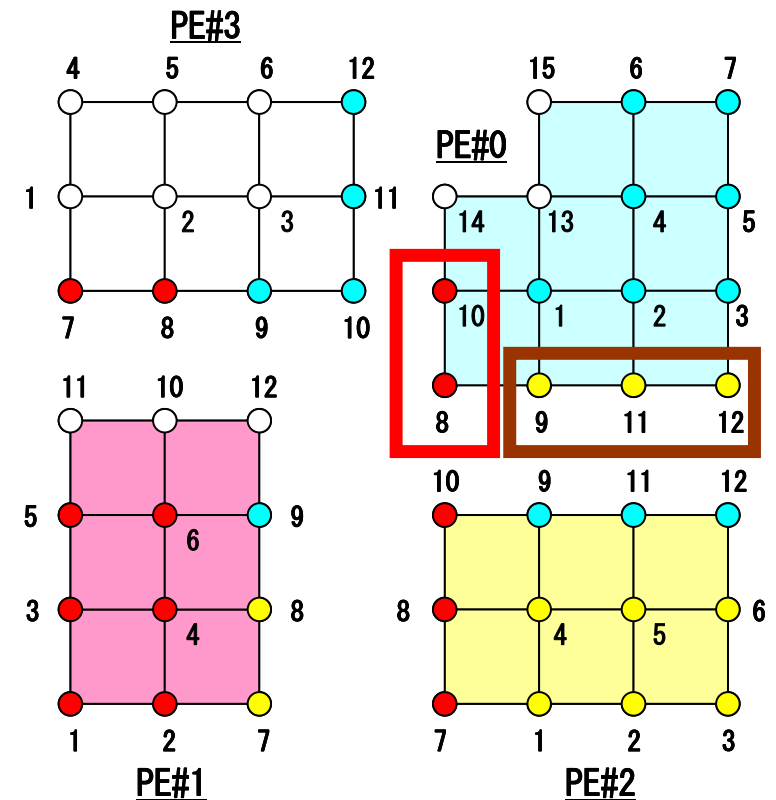
# RECV: receiving to external nodes

Recv. continuous data to recv. buffer from neighbors

- MPI\_Irecv

(recvbuf, count, datatype, dest, tag, comm, request)

- 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



# 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)**

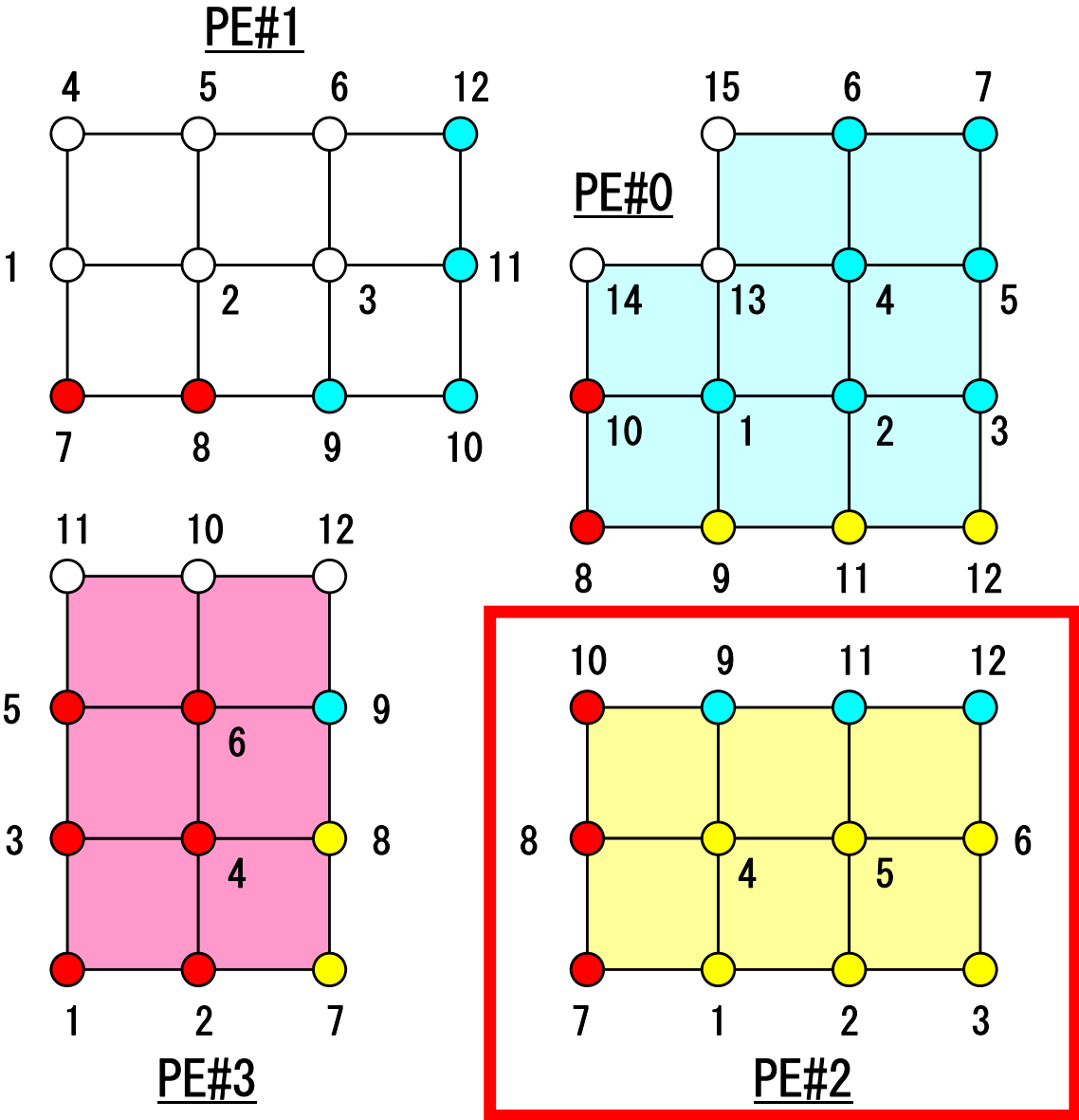
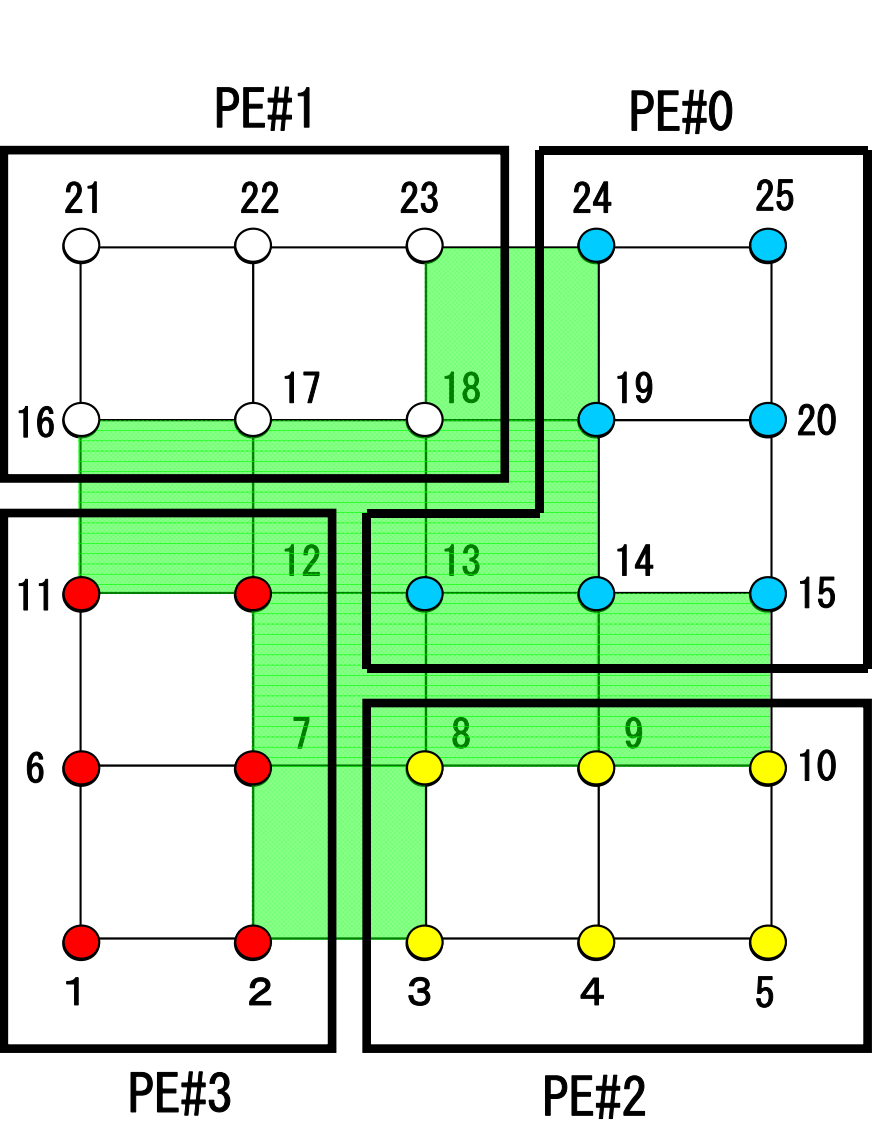
|                                                                                                                                            |        |   |                                           |
|--------------------------------------------------------------------------------------------------------------------------------------------|--------|---|-------------------------------------------|
| – <u>recvbuf</u>                                                                                                                           | choice | I | starting address of receiving buffer      |
| – <u>count</u>                                                                                                                             | I      | I | number of elements in receiving buffer    |
| – <u>datatype</u>                                                                                                                          | I      | I | data type of elements of receiving buffer |
| – <u>source</u>                                                                                                                            | I      | I | rank of source                            |
| – <u>tag</u>                                                                                                                               | 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),                                                                                              |        |   |                                           |
| – <u>comm</u>                                                                                                                              | I      | I | communicator                              |
| – <u>request</u>                                                                                                                           | I      | 0 | communication request used in MPI_Waitall |
| – <u>ierr</u>                                                                                                                              | I      | 0 | completion code                           |

# MPI\_WAITALL

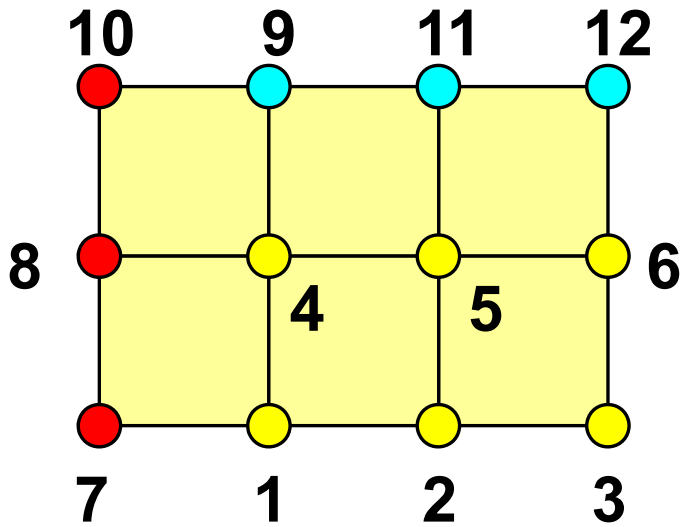
- **MPI\_Waitall** blocks until all comm's, associated with request in the array, complete. It is used for synchronizing MPI\_Isend and MPI\_Irecv in this class.
- At sending phase, contents of sending buffer cannot be modified before calling corresponding **MPI\_Waitall**. At receiving phase, contents 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            0            array of status objects  
MPI\_STATUS\_SIZE: defined in 'mpif.h', 'mpi.h'
  - ierr       I            0            completion code

# Node-based Partitioning

internal nodes - elements - external nodes



# Description of Distributed Local Data

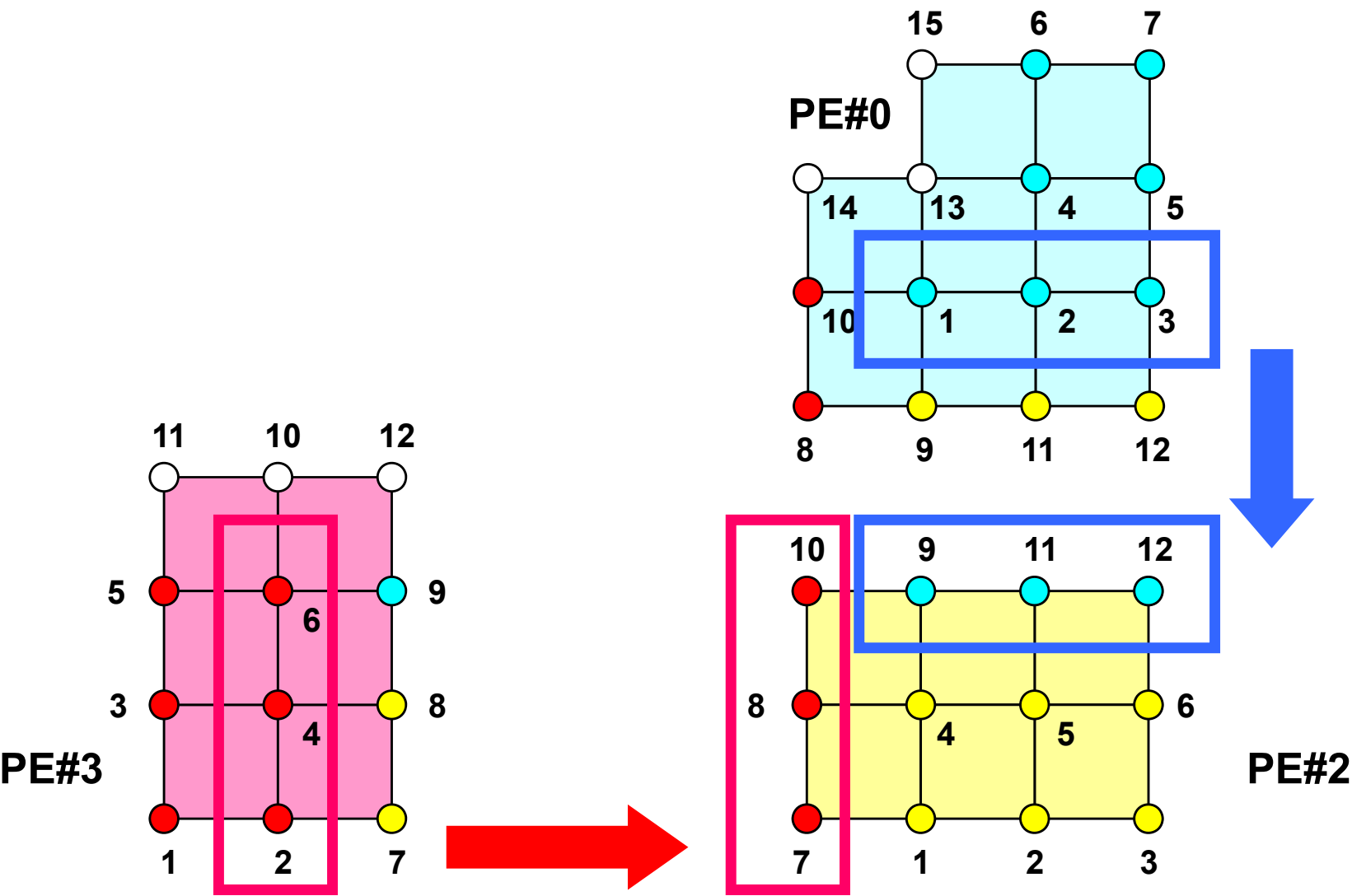


- Internal/External Points
  - Numbering: Starting from internal pts, then external pts after that
- Neighbors
  - Shares overlapped meshes
  - Number and ID of neighbors
- External Points
  - From where, how many, and which external points are received/imported ?
- Boundary Points
  - To where, how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



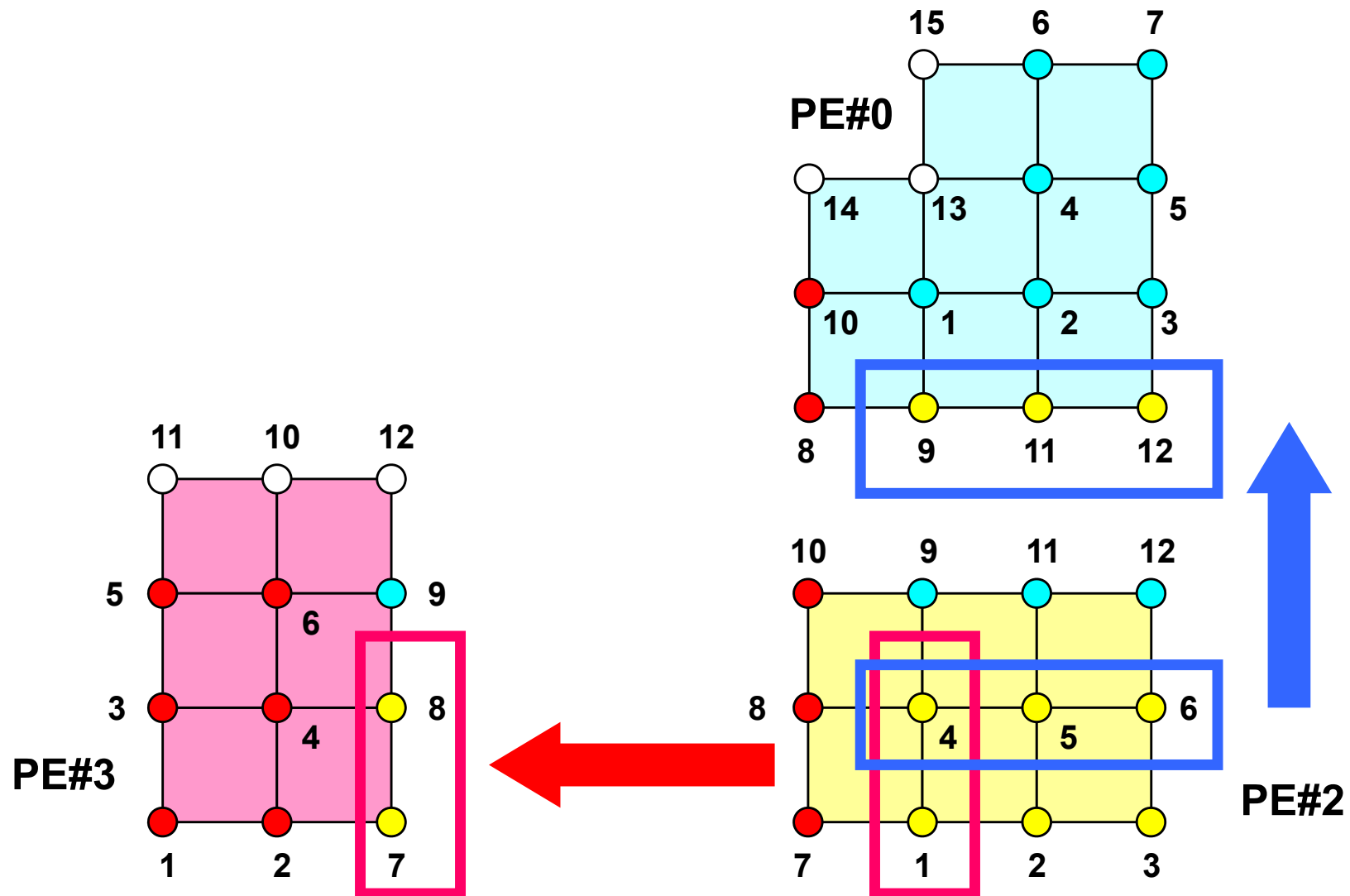
# External Nodes (外点) : RECEIVE

PE#2 : receive information for “external nodes”



# Boundary Nodes (境界点) : SEND

PE#2 : send information on “boundary nodes”



# 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