

Exercise S1

C

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Exercise S1 (1/2)

- Problem S1-1
 - Read local files <\$O-S1>/a1.0~a1.3, <\$O-S1>/a2.0~a2.3.
 - Develop codes which calculate norm $\|x\|_2$ of global vector for each case.
 - <\$O-S1>file.c, <\$T-S1>file2.c
- Problem S1-2
 - Read local files <\$O-S1>/a2.0~a2.3.
 - Develop a code which constructs “global vector” using MPI_Allgatherv.

Exercise S1 (2/2)

- Problem S1-3
 - Develop parallel program which calculates the following numerical integration using “trapezoidal rule” by MPI_Reduce, MPI_Bcast etc.
 - Measure computation time, and parallel performance

$$\int_0^1 \frac{4}{1+x^2} dx$$

Copying files on Odyssey

Fortran

```
>$ cd /work/gt36/t36XXX/pFEM  
>$ moule load fj  
>$ cp /work/gt00/z30088/pFEM/F/s1r-f.tar .  
>$ tar xvf s1r-f.tar
```

C

```
>$ cd /work/gt36/t36XXX/pFEM  
>$ module load fj  
>$ cp /work/gt00/z30088/pFEM/C/s1r-c.tar .  
>$ tar xvf s1r-c.tar
```

Confirm directory

```
>$ ls  
    mpi  
>$ cd mpi/S1-ref
```

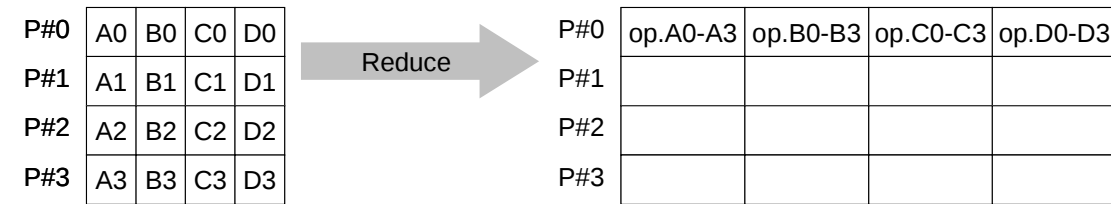
This directory is called as **<\$0-S1r>**.

<\$0-S1r> = <\$0-TOP>/mpi/S1-ref

S1-1: Reading Local Vector, Calc. Norm

- Problem S1-1
 - Read local files <\$O-S1>/a1.0~a1.3, <\$O-S1>/a2.0~a2.3.
 - Develop codes which calculate norm $\|x\|_2$ of global vector for each case.
- Use MPI_Allreduce (or MPI_Reduce)
- Advice
 - Checking each component of variables and arrays !

MPI_Reduce



- Reduces values on all processes to a single value
 - Summation, Product, Max, Min etc.
- **MPI_Reduce (sendbuf, recvbuf, count, datatype, op, root, comm)**
 - **sendbuf** choice I starting address of send buffer
 - **recvbuf** choice 0 starting address receive buffer
 - type is defined by "**datatype**"
 - **count** int I number of elements in send/receive buffer
 - **datatype** MPI_Datatype I data type of elements of send/recive buffer
 - FORTTRAN MPI_INTEGER, MPI_REAL, MPI_DOUBLE_PRECISION, MPI_CHARACTER etc.
 - C MPI_INT, MPI_FLOAT, MPI_DOUBLE, MPI_CHAR etc
 - **op** MPI_Op 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** int I rank of root process
 - **comm** MPI_Comm I communicator

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.

“op” of MPI_Reduce/Allreduce

MPI_Reduce

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

- **MPI_MAX, MPI_MIN** Max, Min
- **MPI_SUM, MPI_PROD** Summation, Product
- **MPI LAND** Logical AND

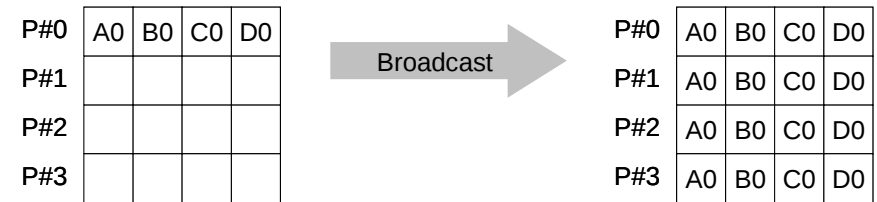
```
double X0, XSUM;
```

```
MPI_Reduce  
(&X0, &XSUM, 1, MPI_DOUBLE, MPI_SUM, 0, <comm>)
```

```
double X0[4];
```

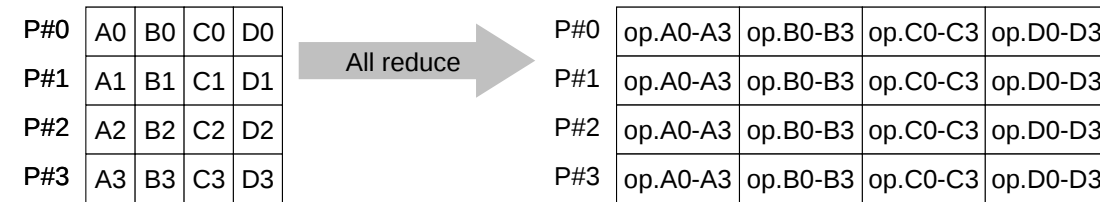
```
MPI_Reduce  
(&X0[0], &X0[2], 2, MPI_DOUBLE_PRECISION, MPI_SUM, 0, <comm>)
```

MPI_Bcast



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

MPI_Allreduce



- MPI_Reduce + MPI_Bcast
- Summation (of dot products) and MAX/MIN values are likely to be utilized in each process

- call MPI_Allreduce

(**sendbuf**, **recvbuf**, **count**, **datatype**, **op**, **comm**)

- sendbuf choice I starting address of send buffer
- recvbuf choice 0 starting address receive buffer
- type is defined by "datatype"

- count int I number of elements in send/receive buffer
- datatype MPI_Datatype I data type of elements of send/recive buffer

- op MPI_Op I reduce operation
- comm MPI_Comm I communicator

S1-1: Local Vector, Norm Calculation

Uniform Vectors (a1.*): s1-1-for_a1.c

```

#include <mpi.h>
#include <stdio.h>
#include <math.h>
#include <assert.h>

int main(int argc, char **argv){
    int i, N;
    int PeTot, MyRank;
    MPI_Comm SolverComm;
    double vec[8];
    double sum0, sum;
    char filename[80];
    FILE *fp;

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &PeTot);
    MPI_Comm_rank(MPI_COMM_WORLD, &MyRank);

    sprintf(filename, "a1.%d", MyRank);
    fp = fopen(filename, "r");
    assert(fp != NULL);

    N=8;
    for(i=0; i<N; i++){
        fscanf(fp, "%lf", &vec[i]);
    }
    sum0 = 0.0;
    for(i=0; i<N; i++){
        sum0 += vec[i] * vec[i];
    }

    MPI_Allreduce(&sum0, &sum, 1, MPI_DOUBLE, MPI_SUM, MPI_COMM_WORLD);
    sum = sqrt(sum);

    if(!MyRank) printf("%27.20E\n", sum);
    MPI_Finalize();
    return 0;
}

```

S1-1: Local Vector, Norm Calculation

Non-uniform Vectors (a2.*): s1-1-for_a2.c

```

#include <mpi.h>
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#include <assert.h>

int main(int argc, char **argv){
    int i, PeTot, MyRank, n;
    MPI_Comm SolverComm;
    double *vec, *vec2;
    int * Count, CountIndex;
    double sum0, sum;
    char filename[80];
    FILE *fp;

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &PeTot);
    MPI_Comm_rank(MPI_COMM_WORLD, &MyRank);

    sprintf(filename, "a2.%d", MyRank);
    fp = fopen(filename, "r");
    assert(fp != NULL);

    fscanf(fp, "%d", &n);
    vec = malloc(n * sizeof(double));
    for(i=0; i<n; i++){
        fscanf(fp, "%lf", &vec[i]);
    }
    sum0 = 0.0;
    for(i=0; i<n; i++){
        sum0 += vec[i] * vec[i];
    }

    MPI_Allreduce(&sum0, &sum, 1, MPI_DOUBLE, MPI_SUM, MPI_COMM_WORLD);
    sum = sqrt(sum);

    if(!MyRank) printf("%27.20E\n", sum);
    MPI_Finalize();
    return 0;}

```

S1-1: Running the Codes

FORTRAN

```
$ cd /work/gt36/t36XXX/pFEM/mpi/S1-ref  
$ module load fj  
$ mpifrtpx -Kfast s1-1-for_a1.f  
$ mpifrtpx -Kfast s1-1-for_a2.f  
  
(modify "go4.sh")  
$ pjsub go4.sh
```

C

```
$ cd /work/gt36/t36XXX/pFEM/mpi/S1-ref  
$ module load fj  
$ mpifccpx -Nclang -Kfast s1-1-for_a1.c  
$ mpifccpx -Nclang -Kfast s1-1-for_a2.c  
  
(modify "go4.sh")  
$ pjsub go4.sh
```

S1-1: Local Vector, Calc. Norm Results

Results using one core

```
a1.* 1.62088247569032590000E+03  
a2.* 1.22218492872396360000E+03
```

```
$> frtpx -Kfast dot-a1.f  
$> pjsub go1.sh
```

```
$> frtpx -Kfast dot-a2.f  
$> pjsub go1.sh
```

Results

```
a1.* 1.62088247569032590000E+03  
a2.* 1.22218492872396360000E+03
```

go1.sh

```
#!/bin/bash  
#PJM -N "test"  
#PJM -L "rscgrp=lecture6-o"  
#PJM -L node=1  
#PJM --mpi proc=1  
#PJM -L elapse=00:15:00  
#PJM -g gt36  
#PJM -j  
#PJM -e err  
#PJM -o test.lst
```

```
module load fj  
module load fjmpi
```

```
mpiexec ./a.out
```

S1-1: Local Vector, Calc. Norm

If SENDBUF=RECVBUF, what happens ?

True

```
MPI_Allreduce(&sum0, &sum, 1, MPI_DOUBLE, MPI_SUM,  
              MPI_COMM_WORLD)
```

False

```
MPI_Allreduce(&sum0, &sum0, 1, MPI_DOUBLE, MPI_SUM,  
              MPI_COMM_WORLD)
```

S1-1: Local Vector, Calc. Norm

If SENDBUF=RECVBUF, what happens ?

True

```
MPI_Allreduce(&sum0, &sum, 1, MPI_DOUBLE, MPI_SUM,  
              MPI_COMM_WORLD)
```

False

```
MPI_Allreduce(&sum0, &sum0, 1, MPI_DOUBLE, MPI_SUM,  
              MPI_COMM_WORLD)
```

True

```
MPI_Allreduce(&sumK[1], &sumK[2], 1, MPI_DOUBLE, MPI_SUM,  
              MPI_COMM_WORLD)
```

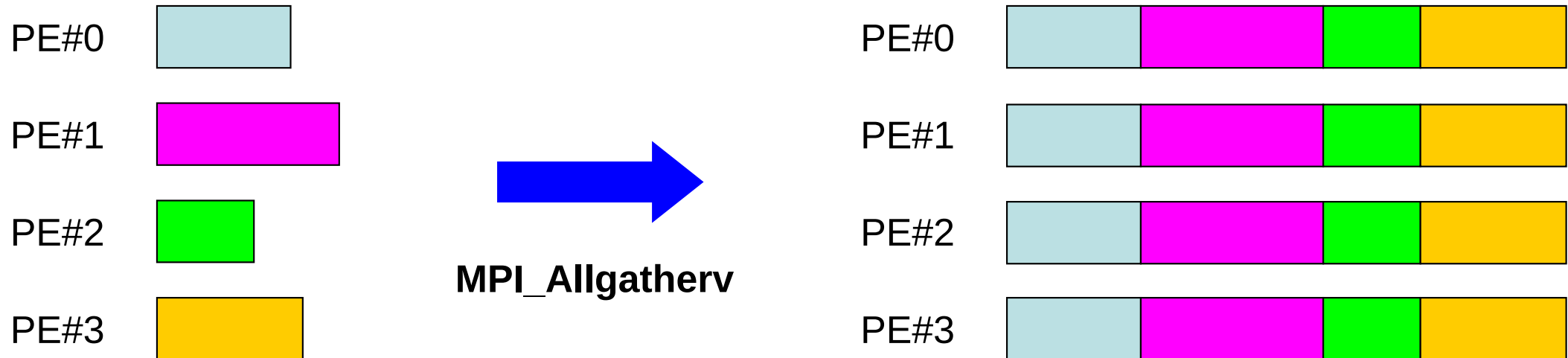
SENDBUF .ne. RECVBUF

S1-2: Local -> Global Vector

- Problem S1-2
 - Read local files <\$O-S1>/a2.0~a2.3.
 - Develop a code which constructs “global vector” using MPI_Allgatherv.

S1-2: Local -> Global Vector

MPI_Allgatherv (1/5)

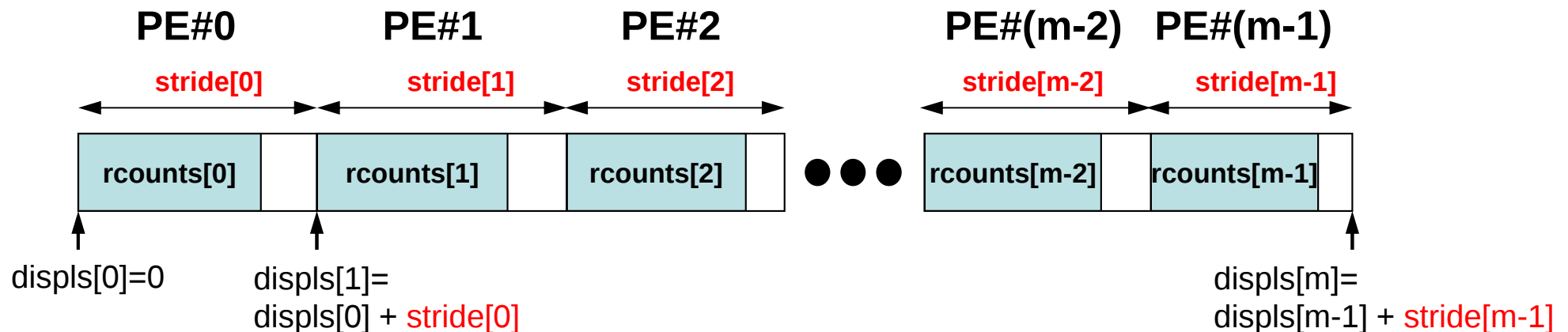


MPI_Allgather

- Variable count version of MPI_Allgather
 - creates “global data” from “local data”
- **MPI_Allgatherv (sendbuf, scount, sendtype, recvbuf, rcounts, displs, recvtype, comm)**
 - **sendbuf** choice I starting address of sending buffer
 - **scount** int I number of elements sent to each process
 - **sendtype** MPI_Datatype I data type of elements of sending buffer
 - **recvbuf** choice 0 starting address of receiving buffer
 - **rcounts** int I integer array (of length group size) containing the number of elements that are to be received from each process (array: size= PETOT)
 - **displs** int I integer array (of length group size). Entry *i* specifies the displacement (relative to recvbuf) at which to place the incoming data from process *i* (array: size= PETOT+1)
 - **recvtype** MPI_Datatype I data type of elements of receiving buffer
 - **comm** MPI_Comm I communicator

MPI_Allgatherv (cont.)

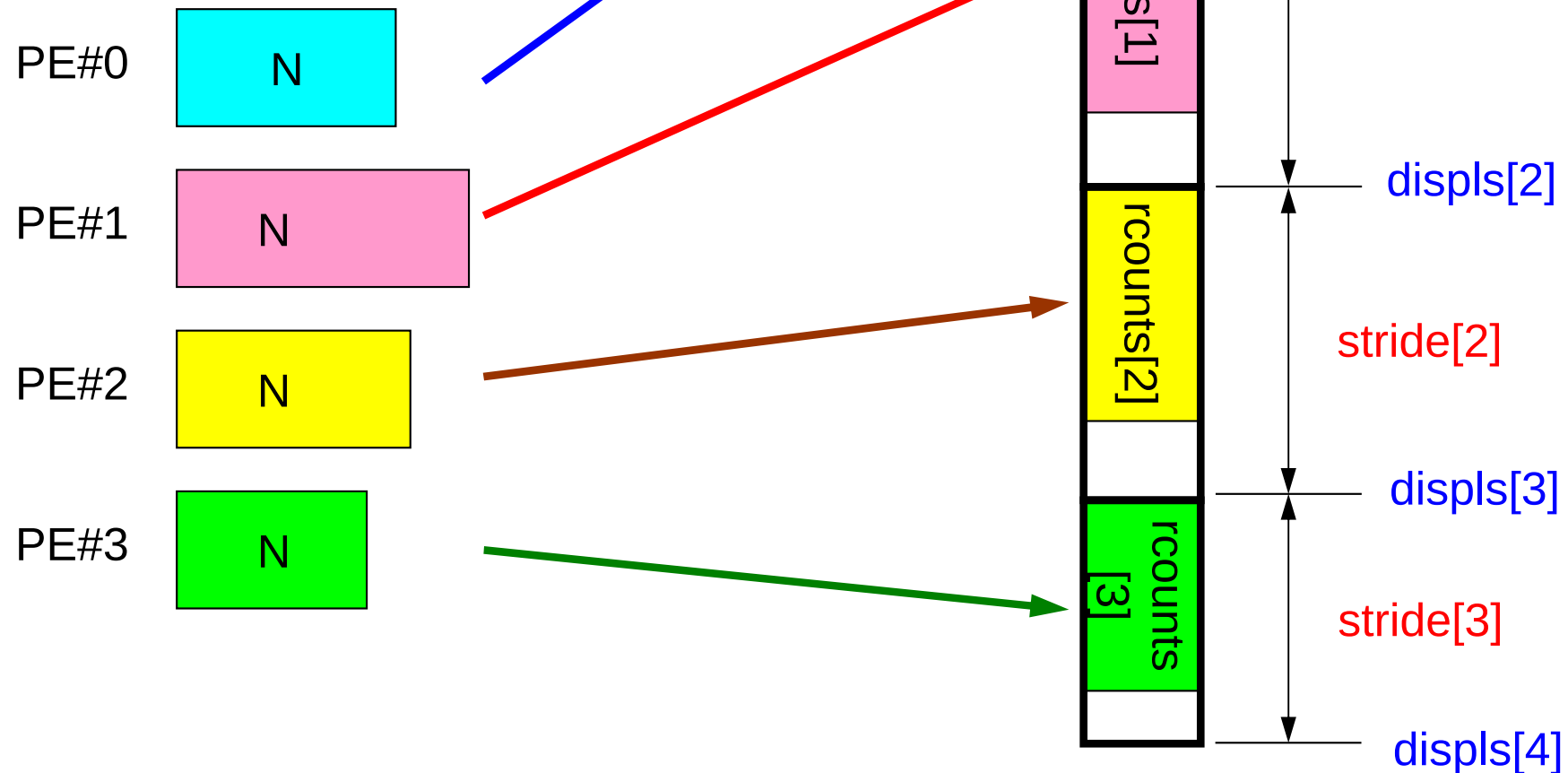
- **MPI_Allgatherv** (sendbuf, scount, sendtype, recvbuf, rcounts, displs, recvtype, comm)
 - **rcounts** int I integer array (of length group size) containing the number of elements that are to be received from each process (array: size= PETOT)
 - **displs** int I integer array (of length group size). Entry i specifies the displacement (relative to recvbuf) at which to place the incoming data from process i (array: size= PETOT+1)
 - These two arrays are related to size of final “global data”, therefore each process requires information of these arrays (rcounts, displs)
 - Each process must have same values for all components of both vectors
 - Usually, **stride(i)=rcounts(i)**



$$\text{size}[\text{recvbuf}] = \text{displs}[\text{PETOT}] = \text{sum}[\text{stride}]$$

What MPI_Allgatherv is doing

Generating global data from local data

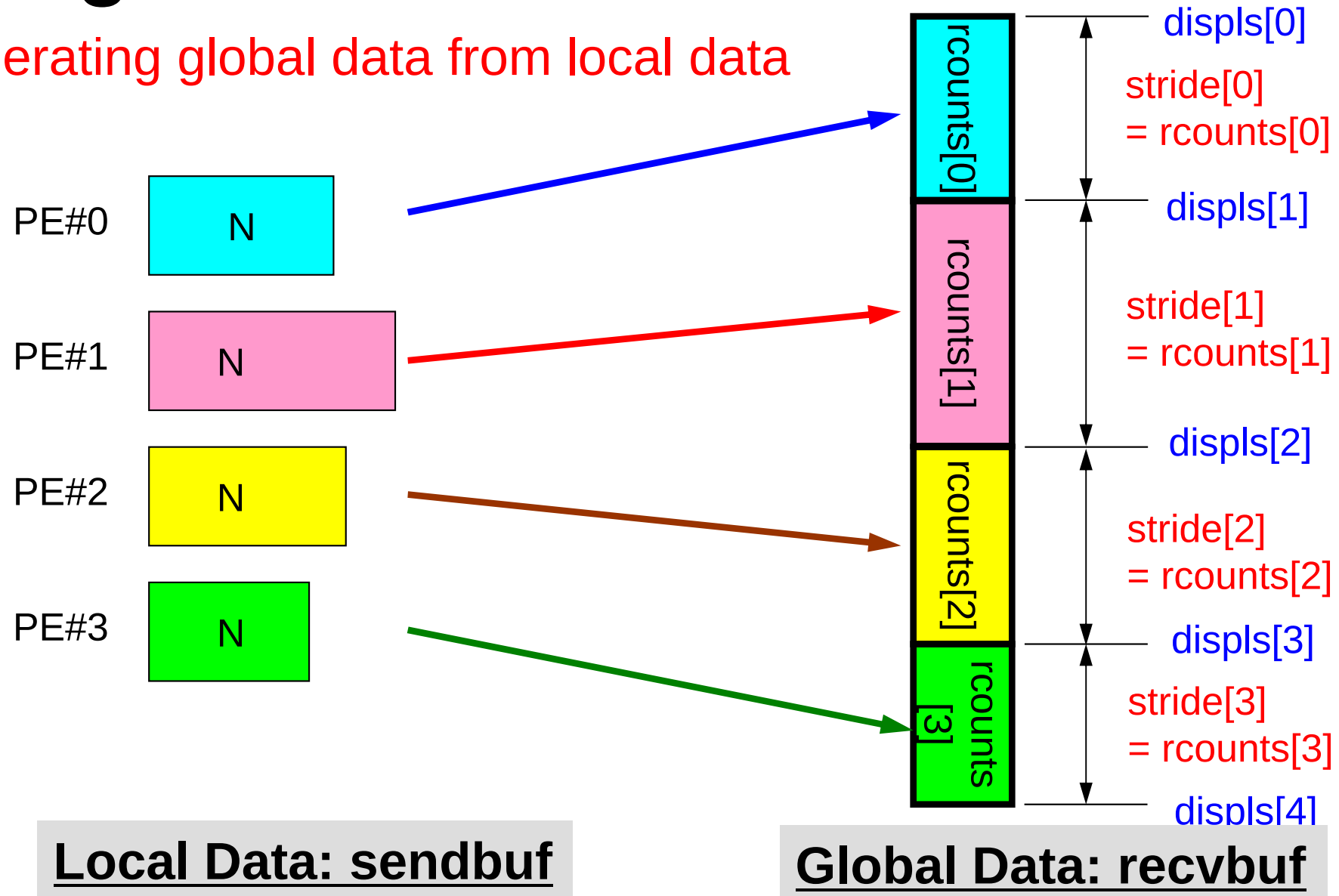


Local Data: sendbuf

Global Data: recvbuf

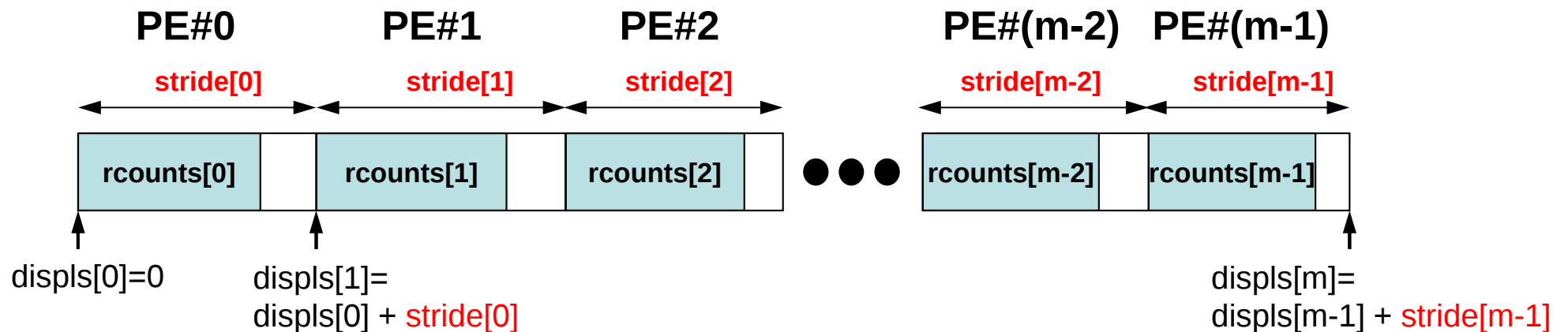
What MPI_Allgatherv is doing

Generating global data from local data



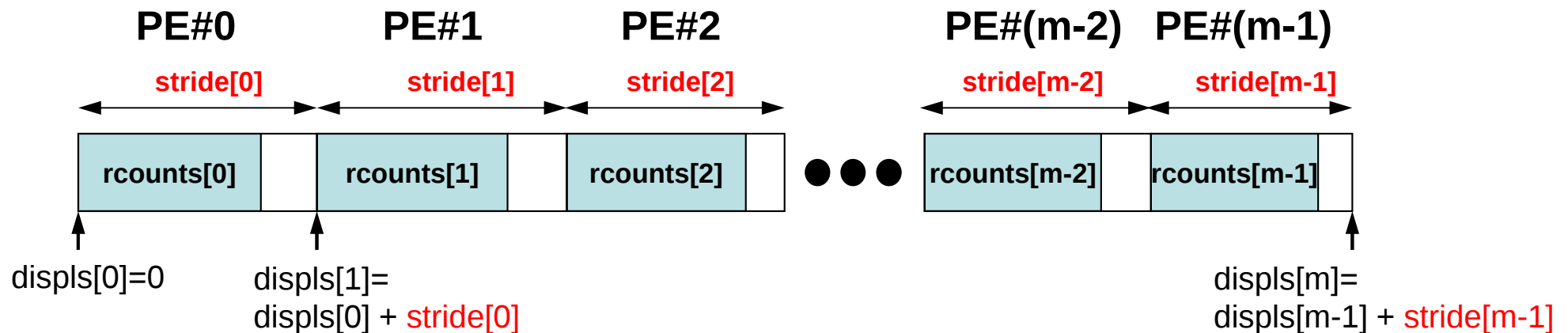
MPI_Allgatherv in detail (1/2)

- `MPI_Allgatherv (sendbuf, scount, sendtype, recvbuf, rcounts, displs, recvtype, comm)`
- **rcounts**
 - Size of message from each PE: Size of Local Data (Length of Local Vector)
- **displs**
 - Address/index of each local data in the vector of global data
 - **`displs(PETOT+1)`** = Size of Entire Global Data (Global Vector)



MPI_Allgatherv in detail (2/2)

- Each process needs information of **rcounts** & **displs**
 - "rcounts"** can be created by gathering local vector length **"N"** from each process.
 - On each process, **"displs"** can be generated from **"rcounts"** on each process.
 - $\text{stride}[i] = \text{rcounts}[i]$
 - Size of **"recvbuf"** is calculated by summation of **"rcounts"**.



$$\text{size}[\text{recvbuf}] = \text{displs}[\text{PETOT}] = \text{sum}[\text{stride}]$$

Preparation for MPI_Allgatherv

<O-S1>/agv.c

- “Generating global vector from “a2.0”~”a2.3”.
- Length of the each vector is 8, 5, 7, and 3, respectively. Therefore, size of final global vector is 23 (= 8+5+7+3).

a2.0~a2.3

PE#0

8

101.0
103.0
105.0
106.0
109.0
111.0
121.0
151.0

PE#1

5

201.0
203.0
205.0
206.0
209.0

PE#2

7

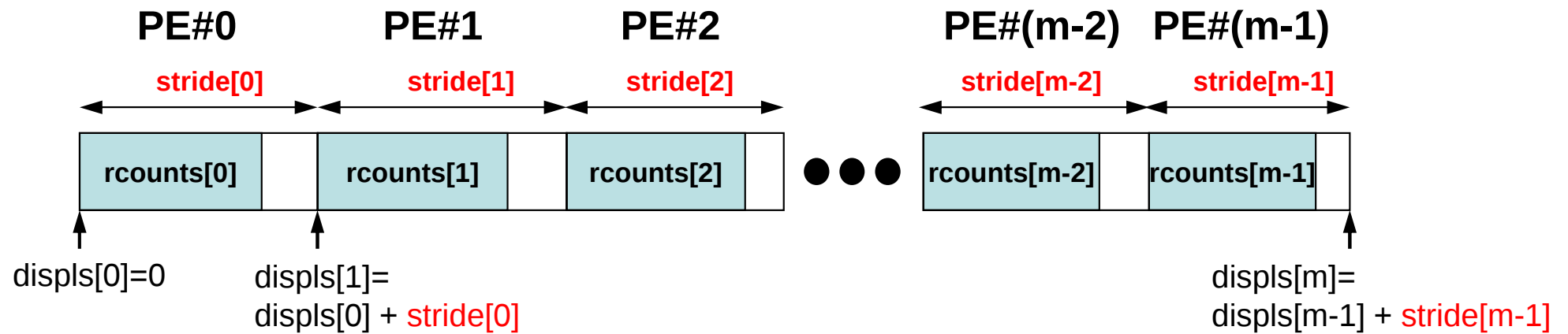
301.0
303.0
305.0
306.0
311.0
321.0
351.0

PE#3

3

401.0
403.0
405.0

S1-2: Local -> Global Vector



`size[recvbuf]= displs[PETOT]= sum[stride]`

- Read local vectors
- Create “rcounts” and “displs”
- Prepare “recvbuf”
- Do “Allgatherv”

S1-2: Local -> Global Vector (1/2)

s1-2.c

```

#include <mpi.h>
#include <stdio.h>
#include <stdlib.h>
#include <math.h>
#include <assert.h>

int main(int argc, char **argv){
    int i, PeTot, MyRank, n;
    MPI_Comm SolverComm;
    double *vec, *vec2, *vecg;
    int *Rcounts, *Displs;
    double sum0, sum;
    char filename[80];
    FILE *fp;

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &PeTot);
    MPI_Comm_rank(MPI_COMM_WORLD, &MyRank);

    sprintf(filename, "a2.%d", MyRank);
    fp = fopen(filename, "r");
    assert(fp != NULL);

    fscanf(fp, "%d", &n);
    vec = malloc(n * sizeof(double));
    for(i=0; i<n; i++){
        fscanf(fp, "%lf", &vec[i]);
    }

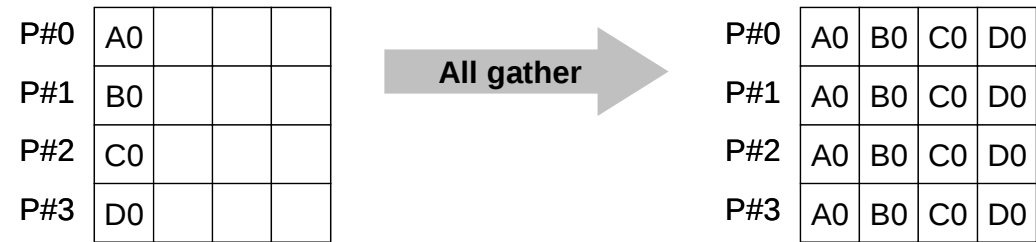
    Rcounts = calloc(PeTot, sizeof(int));
    Displs = calloc(PeTot+1, sizeof(int));

    MPI_Allgather(&n, 1, MPI_INT, Rcounts, 1, MPI_INT, MPI_COMM_WORLD);

```

“Rcounts”
vector length at each PE

MPI_Allgather



- MPI_Gather+MPI_Bcast
 - Gathers data from all tasks and distribute the combined data to all tasks
- MPI_Allgather (sendbuf, scount, sendtype, recvbuf, rcount, recvtype, comm)
 - sendbuf choice I starting address of sending buffer
 - scount int I number of elements sent to each process
 - sendtype MPI_Datatype I data type of elements of sending buffer
 - recvbuf choice 0 starting address of receiving buffer
 - rcount int I number of elements received from the root process
 - recvtype MPI_Datatype I data type of elements of receiving buffer
 - comm MPI_Comm I communicator

S1-2: Local -> Global Vector (2/2)

s1-2.c

```

Displs[0]=0;
for(i=0;i<PeTot;i++){
    Displs[i+1] = Displs[i] + Rcounts[i];
}

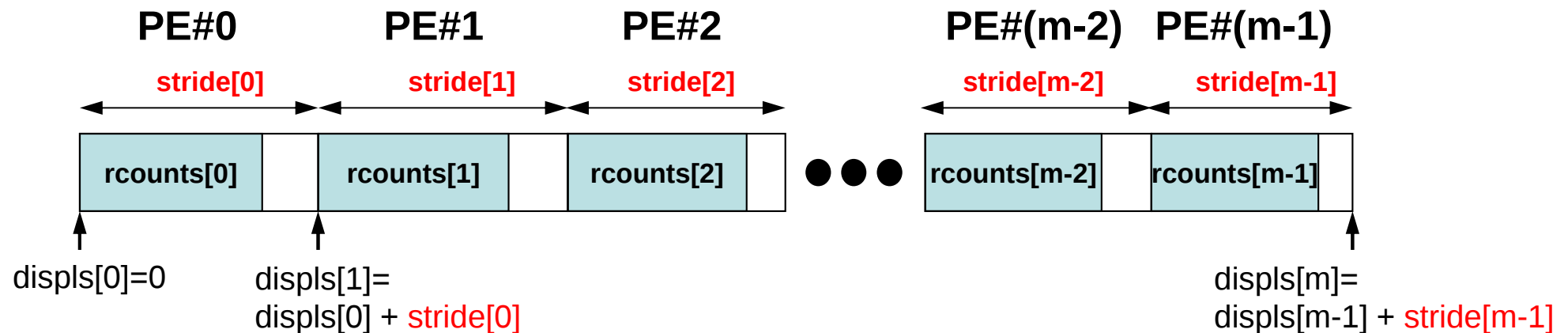
```

Creating "Displs"

```

vecg = calloc(Displs[PeTot], sizeof(double));
MPI_Allgatherv(vec, n, MPI_DOUBLE, vecg, Rcounts, Displs, MPI_DOUBLE, MPI_COMM_WORLD);
for(i=0;i<Displs[PeTot];i++){
    printf("%8.2f", vecg[i]);
}
printf("\n");
MPI_Finalize();
return 0;
}

```



`size[recvbuf]= displs[PETOT]= sum[stride]`

S1-2: Local -> Global Vector (2/2)

s1-2.c

```
Displs[0]=0;
for(i=0;i<PeTot;i++){
    Displs[i+1] = Displs[i] + Rcounts[i];
}
```

```
vecg = calloc(Displs[PeTot], sizeof(double));
```

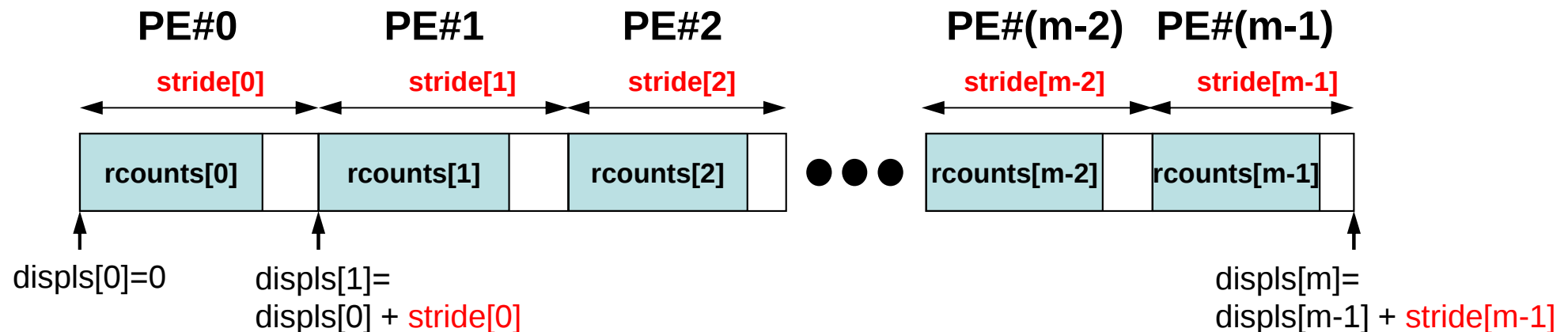
“recvbuf”

```
MPI_Allgatherv(vec, n, MPI_DOUBLE, vecg, Rcounts, Displs, MPI_DOUBLE, MPI_COMM_WORLD);
```

```
for(i=0;i<Displs[PeTot];i++){
    printf("%8.2f", vecg[i]);
}
printf("\n");
```

```
MPI_Finalize();
return 0;
```

```
}
```



$\text{size}[\text{recvbuf}] = \text{displs}[\text{PETOT}] = \text{sum}[\text{stride}]$

S1-2: Local -> Global Vector (2/2)

s1-2.c

```

Displs[0]=0;
for(i=0;i<PeTot;i++){
    Displs[i+1] = Displs[i] + Rcounts[i];
}

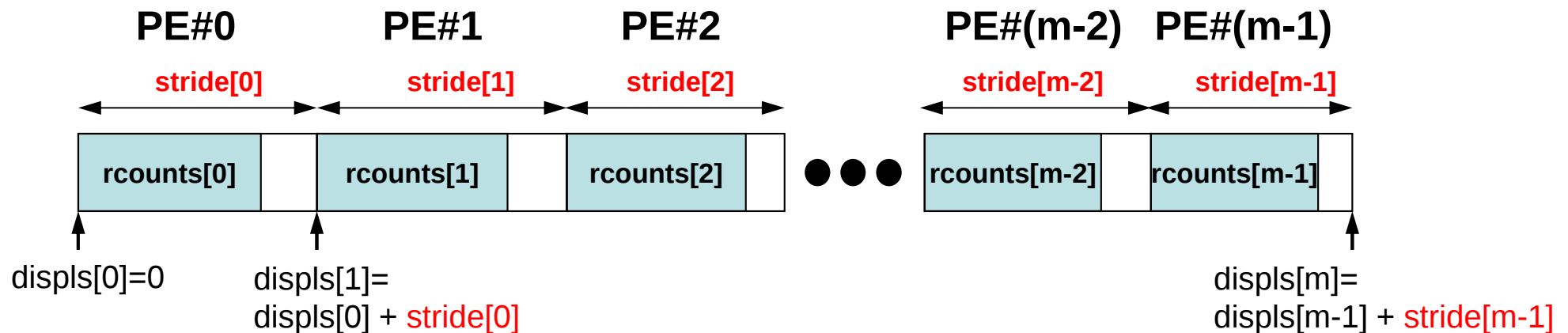
vecg = calloc(Displs[PeTot], sizeof(double));

MPI_Allgatherv(vec, n, MPI_DOUBLE, vecg, Rcounts, Displs, MPI_DOUBLE, MPI_COMM_WORLD);

for(i=0;i<Displs[PeTot];i++){
    printf("%8.2f", vecg[i]);
}
printf("\n");

MPI_Finalize();
return 0;
}

```



$\text{size}[\text{recvbuf}] = \text{displs}[\text{PETOT}] = \text{sum}[\text{stride}]$

S1-2: Running the Codes

FORTRAN

```
$ cd /work/gt36/t36XXX/pFEM/mpi/S1-ref  
$ module load fj  
$ mpifrtpx -Kfast s1-2.f  
  
(modify "go4.sh")  
$ pjsub go4.sh
```

C

```
$ cd /work/gt36/t36XXX/pFEM/mpi/S1-ref  
$ module load fj  
$ mpifccpx -Nclang -Kfast s1-2.c  
  
(modify "go4.sh")  
$ pjsub go4.sh
```

S1-2: Results

my_rank	ID	VAL
0	1	101.
0	2	103.
0	3	105.
0	4	106.
0	5	109.
0	6	111.
0	7	121.
0	8	151.
0	9	201.
0	10	203.
0	11	205.
0	12	206.
0	13	209.
0	14	301.
0	15	303.
0	16	305.
0	17	306.
0	18	311.
0	19	321.
0	20	351.
0	21	401.
0	22	403.
0	23	405.

my_rank	ID	VAL
1	1	101.
1	2	103.
1	3	105.
1	4	106.
1	5	109.
1	6	111.
1	7	121.
1	8	151.
1	9	201.
1	10	203.
1	11	205.
1	12	206.
1	13	209.
1	14	301.
1	15	303.
1	16	305.
1	17	306.
1	18	311.
1	19	321.
1	20	351.
1	21	401.
1	22	403.
1	23	405.

my_rank	ID	VAL
2	1	101.
2	2	103.
2	3	105.
2	4	106.
2	5	109.
2	6	111.
2	7	121.
2	8	151.
2	9	201.
2	10	203.
2	11	205.
2	12	206.
2	13	209.
2	14	301.
2	15	303.
2	16	305.
2	17	306.
2	18	311.
2	19	321.
2	20	351.
2	21	401.
2	22	403.
2	23	405.

my_rank	ID	VAL
3	1	101.
3	2	103.
3	3	105.
3	4	106.
3	5	109.
3	6	111.
3	7	121.
3	8	151.
3	9	201.
3	10	203.
3	11	205.
3	12	206.
3	13	209.
3	14	301.
3	15	303.
3	16	305.
3	17	306.
3	18	311.
3	19	321.
3	20	351.
3	21	401.
3	22	403.
3	23	405.

S1-3: Integration by Trapezoidal Rule

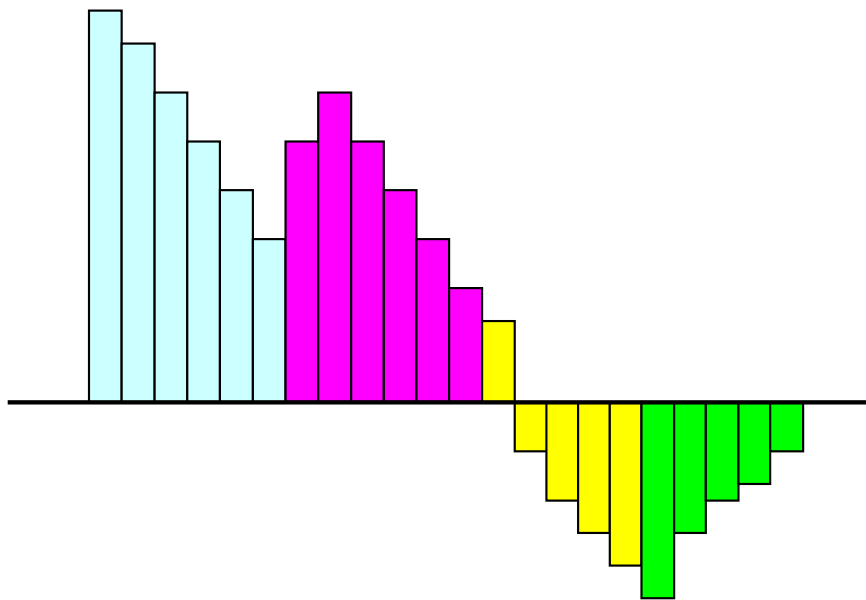
- Problem S1-3
 - Develop parallel program which calculates the following numerical integration using “trapezoidal rule” by MPI_Reduce, MPI_Bcast etc.
 - Measure computation time, and parallel performance

$$\int_0^1 \frac{4}{1+x^2} dx$$

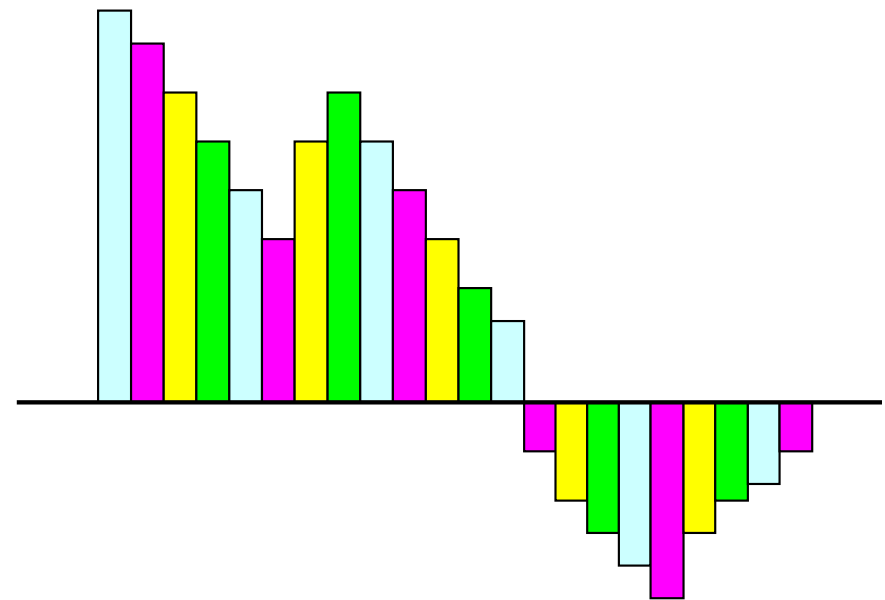
S1-3: Integration by Trapezoidal Rule

Two Types of Load Distribution

Type-A



Type-B



$$\frac{1}{2} \Delta x \left(f_1 + f_{N+1} + \sum_{i=2}^N 2 f_i \right) \text{ corresponds to "Type-A".}$$

S1-3: Integration by Trapezoidal Rule

TYPE-A (1/2): s1-3a.c

```
#include <stdio.h>
#include <stdlib.h>
#include <assert.h>
#include <math.h>
#include "mpi.h"

int main(int argc, char **argv){
    int i;
    double TimeStart, TimeEnd, sum0, sum, dx;
    int PeTot, MyRank, n, int *index;
    FILE *fp;

    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &PeTot);
    MPI_Comm_rank(MPI_COMM_WORLD, &MyRank);

    index = calloc(PeTot+1, sizeof(int));
    fp = fopen("input.dat", "r");
    fscanf(fp, "%d", &n);
    fclose(fp);
    if(MyRank==0) printf("%s%8d\n", "N=", n);
    dx = 1.0/n;

    for(i=0; i<=PeTot; i++){
        index[i] = ((long long)i * n)/PeTot;
    }
}
```

“N (number of segments) “ is specified in “input.dat”

PE#0

PE#1

PE#2

● ● ●

PE#(PETOT-1)

index[0]

index[1]

index[2]

index[3]

index[PETOT-1]

index[PeTot]
=N

S1-3: Integration by Trapezoidal Rule

TYPE-A (2/2): s1-3a.c

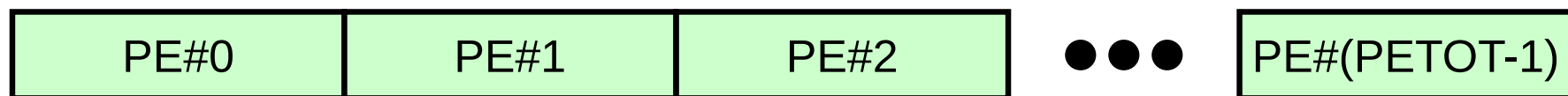
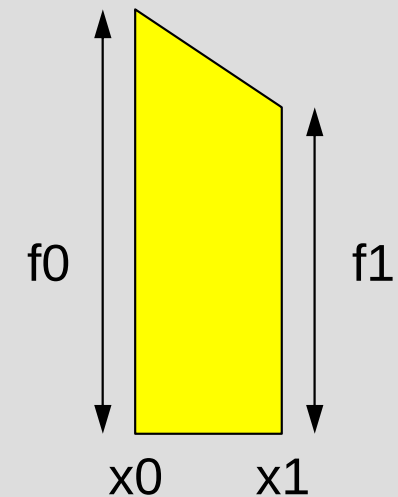
```
TimeS = MPI_Wtime();
sum0 = 0.0;
for(i=index[MyRank]; i<index[MyRank+1]; i++)
{
    double x0, x1, f0, f1;
    x0 = (double)i * dx;
    x1 = (double)(i+1) * dx;
    f0 = 4.0/(1.0+x0*x0);
    f1 = 4.0/(1.0+x1*x1);
    sum0 += 0.5 * (f0 + f1) * dx;
}
```

```
MPI_Reduce(&sum0, &sum, 1, MPI_DOUBLE, MPI_SUM, 0, MPI_COMM_WORLD);
TimeE = MPI_Wtime();
```

```
if(!MyRank) printf("%24.16f%24.16f%24.16f¥n", sum, 4.0*atan(1.0), TimeE - TimeS);
```

```
MPI_Finalize();
return 0;
```

```
}
```



index[0]

index[1]

index[2]

index[3]

index[PETOT-1]

index[PeTot]
=N

S1-3: Integration by Trapezoidal Rule

TYPE-B: s1-3b.c

```

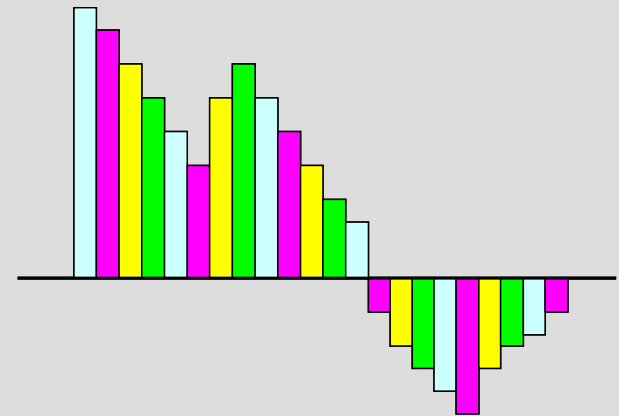
TimeS = MPI_Wtime();
sum0 = 0.0;
for(i=MyRank; i<n; i+=PeTot)
{
    double x0, x1, f0, f1;
    x0 = (double)i * dx;
    x1 = (double)(i+1) * dx;
    f0 = 4.0/(1.0+x0*x0);
    f1 = 4.0/(1.0+x1*x1);
    sum0 += 0.5 * (f0 + f1) * dx;
}

MPI_Reduce(&sum0, &sum, 1, MPI_DOUBLE, MPI_SUM, 0, MPI_COMM_WORLD);
TimeE = MPI_Wtime();

if(!MyRank) printf("%24.16f%24.16f%24.16f\n", sum, 4.0*atan(1.0), TimeE-TimeS);

MPI_Finalize();
return 0;
}

```



S1-3: Running the Codes

```
$ cd /work/gt36/t36XXX/pFEM/mpi/S1-ref  
$ module load fj  
$ mpifrtpx -Kfast s1-3a.f -o s13a  
$ mpifrtpx -Kfast s1-3b.f -o s13b
```

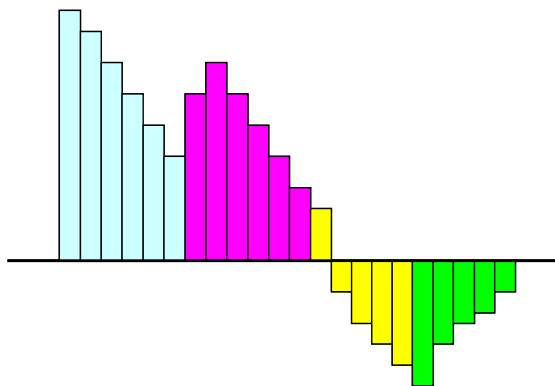
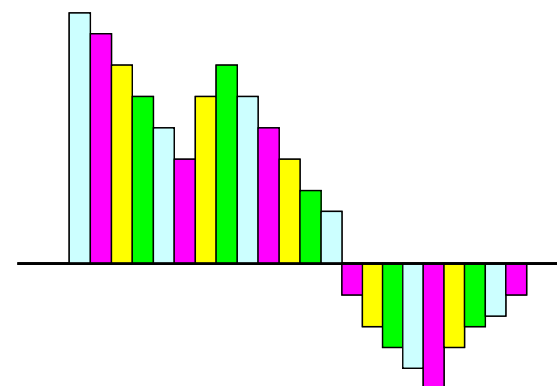
FORTTRAN

```
(modify "XX.sh")  
$ pjsub XX.sh
```

```
$ cd /work/gt36/t36XXX/pFEM/mpi/S1-ref  
$ module load fj  
$ mpifccpx -Nclang -Kfast s1-3a.c -o s13a  
$ mpifccpx -Nclang -Kfast s1-3b.c -o s13b
```

C

```
(modify "XX.sh")  
$ pjsub XX.sh
```

Type-A**Type-B**

a384.sh: 8-nodes, 384-cores

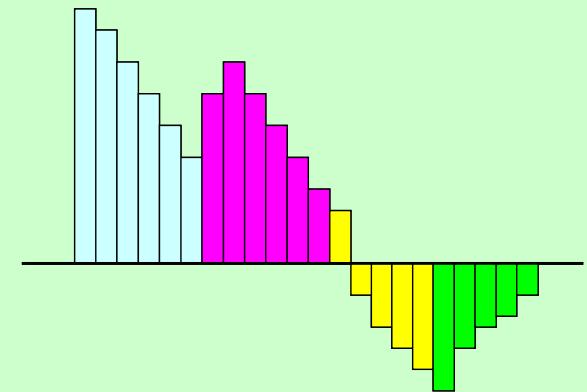
```
#!/bin/sh
#PJM -N "m384"
#PJM -L rscgrp=lecture6-o
#PJM -L node=8
#PJM --mpi proc=384
#PJM -L elapse=00:15:00
#PJM -g gt36
#PJM -j
#PJM -e err
#PJM -o z384.lst
```

```
module load fj
module load fjmp
```

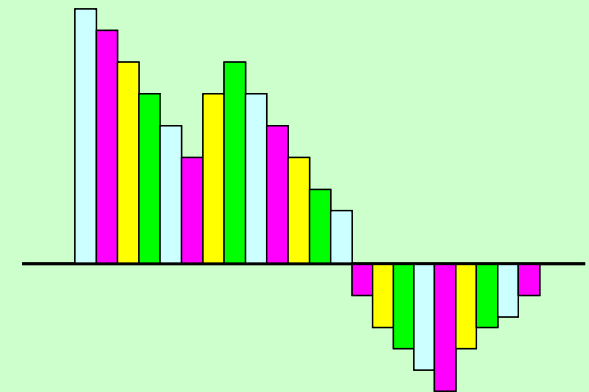
```
mpiexec ./s13a
mpiexec ./s13b
mpiexec numactl -l ./s13a
mpiexec numactl -l ./s13b
```

384/8= 48 cores/node

Type-A



Type-B



numactl -l/--localalloc for utilizing local memory (no effects)

a012.sh

```
#!/bin/sh
#PJM -N "test"
#PJM -L rscgrp=lecture6-o
#PJM -L node=1
#PJM --mpi proc=12
#PJM -L elapse=00:15:00
#PJM -g gt36
#PJM -j
#PJM -e err
#PJM -o test.lst
```

```
module load fj
module load fjmpi
mpiexec ./a.out
mpiexec numactl -l ./a.out
```

a048.sh

```
#!/bin/sh
#PJM -N "test"
#PJM -L rscgrp=lecture6-o
#PJM -L node=1
#PJM --mpi proc=48
#PJM -L elapse=00:15:00
#PJM -g gt36
#PJM -j
#PJM -e err
#PJM -o test.lst
```

```
module load fj
module load fjmpi
mpiexec ./a.out
mpiexec numactl -l ./a.out
```

a384.sh

```
#!/bin/sh
#PJM -N "test"
#PJM -L rscgrp=lecture6-o
#PJM -L node=8
#PJM --mpi proc=384
#PJM -L elapse=00:15:00
#PJM -g gt36
#PJM -j
#PJM -e err
#PJM -o test.lst
```

```
module load fj
module load fjmpi
mpiexec ./a.out
mpiexec numactl -l ./a.out
```

a576.sh

```
#!/bin/sh
#PJM -N "test"
#PJM -L rscgrp=lecture6-o
#PJM -L node=12
#PJM --mpi proc=576
#PJM -L elapse=00:15:00
#PJM -g gt36
#PJM -j
#PJM -e err
#PJM -o test.lst
```

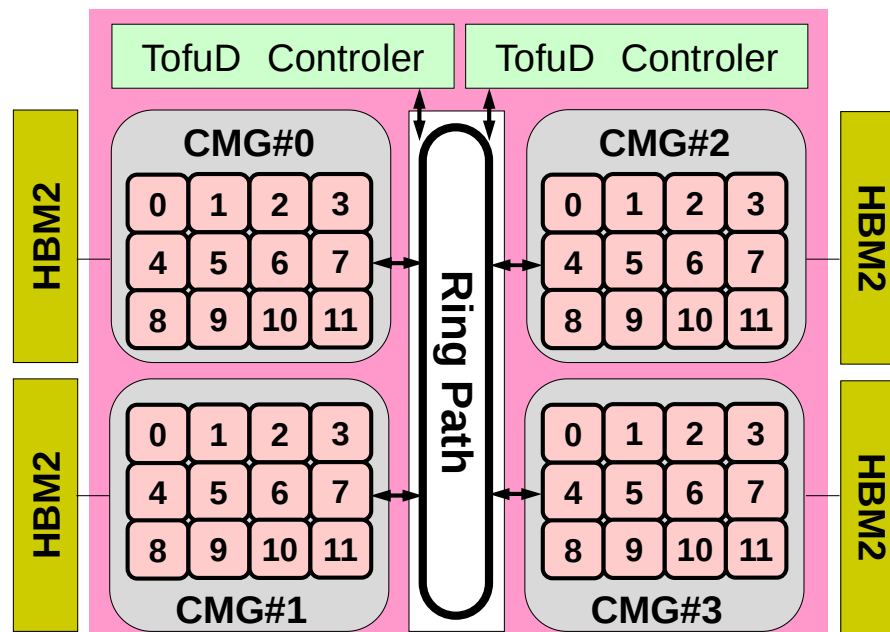
```
module load fj
module load fjmpi
mpiexec ./a.out
mpiexec numactl -l ./a.out
```

numactl -l/--localalloc for utilizing local memory (no effects)

Number of Processes

#PJM	-L	node=1;	#PJM	--mpi	proc= 1	1-node, 1-proc, 1-proc/n
#PJM	-L	node=1;	#PJM	--mpi	proc= 4	1-node, 4-proc, 4-proc/n
#PJM	-L	node=1;	#PJM	--mpi	proc=12	1-node, 12-proc, 12-proc/n
#PJM	-L	node=1;	#PJM	--mpi	proc=24	1-node, 24-proc, 24-proc/n
#PJM	-L	node=1;	#PJM	--mpi	proc=48	1-node, 48-proc, 48-proc/n

#PJM	-L	node= 4;	#PJM	--mpi	proc=192	4-node, 192-proc, 48-proc/n
#PJM	-L	node= 8;	#PJM	--mpi	proc=384	8-node, 384-proc, 48-proc/n
#PJM	-L	node=12;	#PJM	--mpi	proc=576	12-node, 576-proc, 48-proc/n



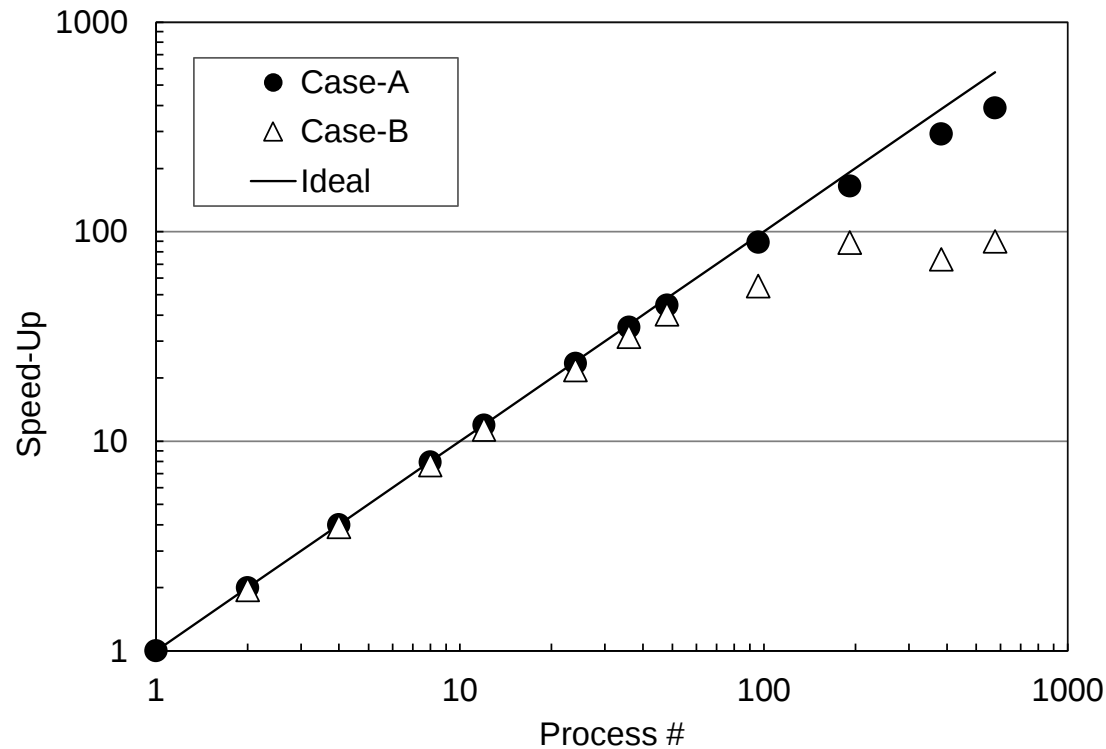
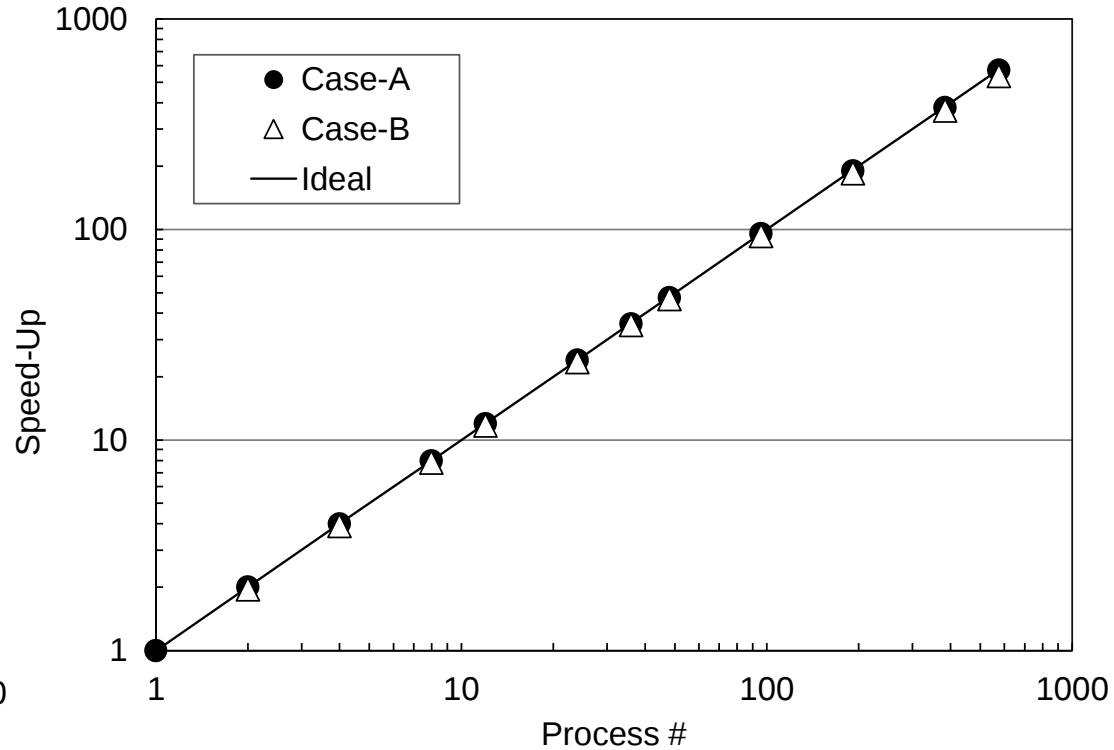
S1-3: Performance on Odyssey

- Based on results (sec.) using a single core
- The best case for 5-runs is selected
- Type A/B
 - Type-A is better, especially for smaller cases
 - Type-B is very slow for C language
- Strong Scaling
 - Entire problem size fixed
 - $1/N$ comp. time using N -x cores
- Weak Scaling
 - Problem size/core is fixed
 - Comp. time is kept constant for N -x scale problems using N -x cores

Strong Scaling: ~ 12-nodes, 576-cores

Speed-Up, Fortran

Performance of Type-A with 1-core= 1.00

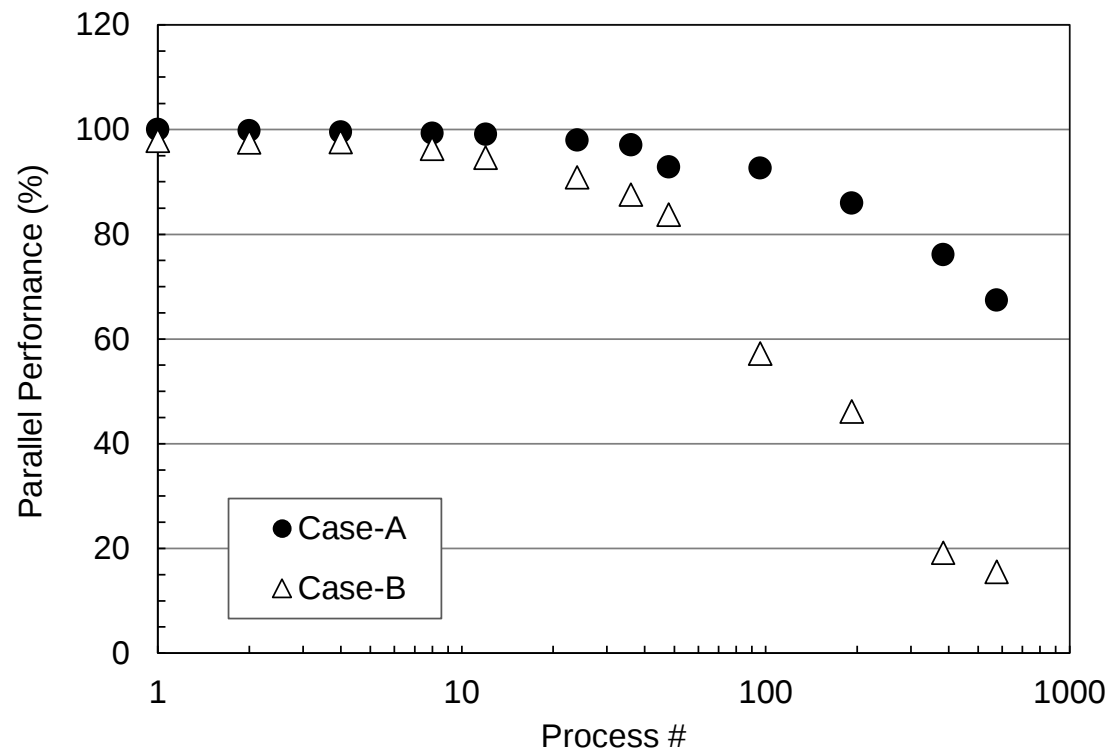
 $N=2 \times 10^7$  **$N=2 \times 10^9$** 

Strong Scaling: ~12-nodes, 576-cores

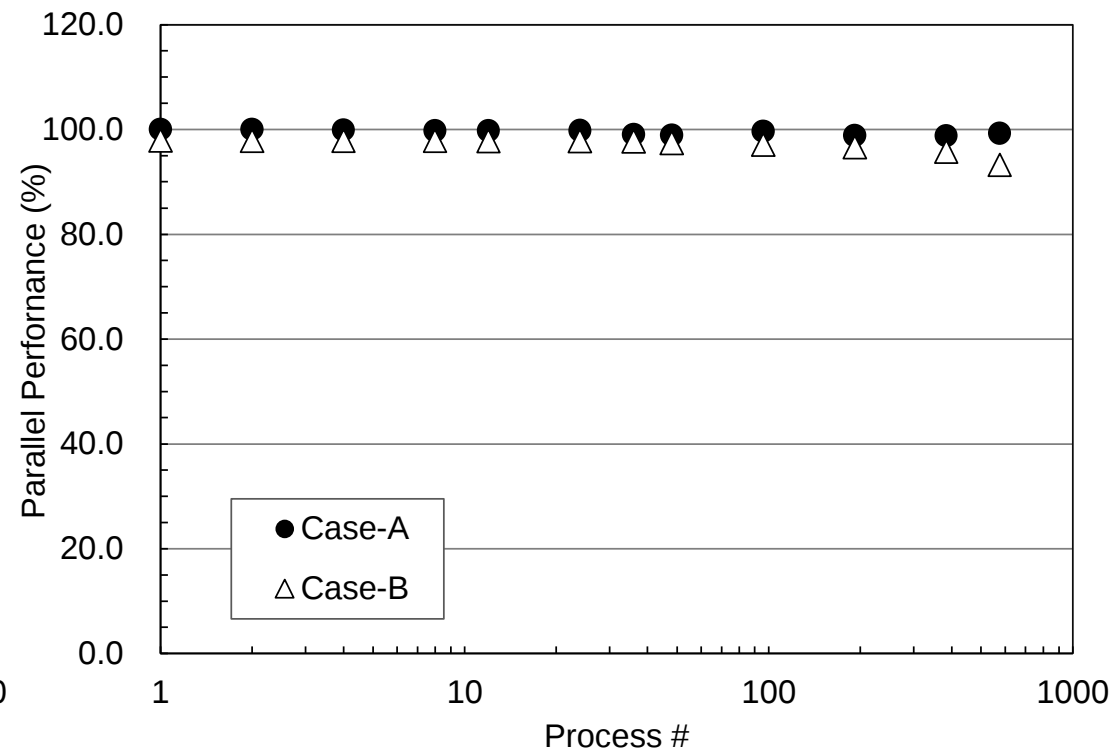
Parallel Performance, Fortran

based on performance of Type-A with 1-core

$N=2 \times 10^7$



$N=2 \times 10^9$



Parallel Performance

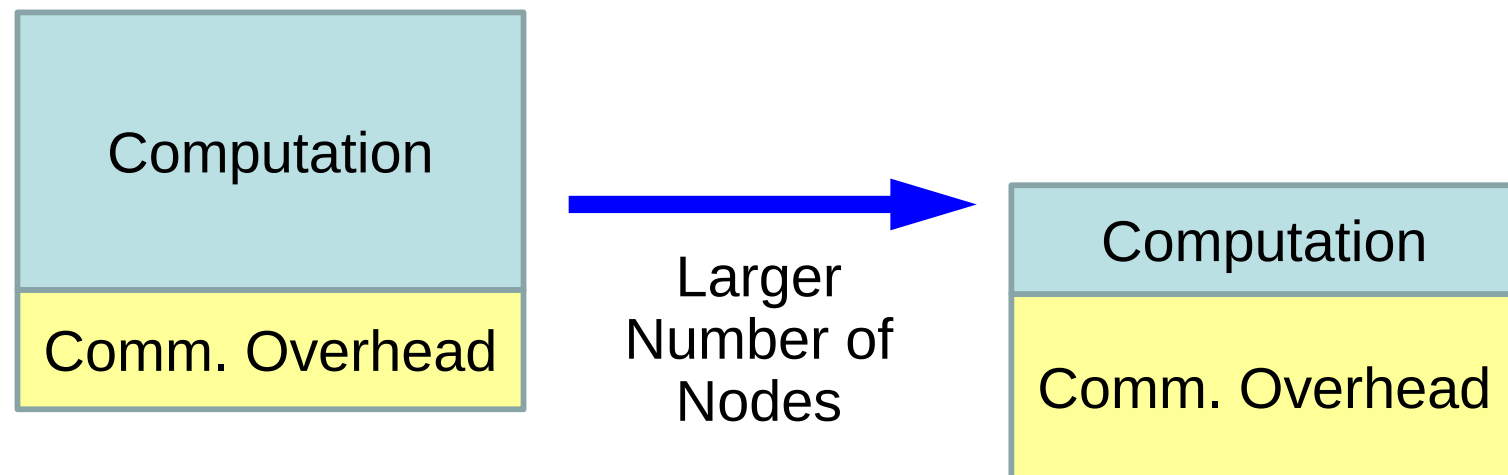
Number of PE's	Computation Time (sec)	Parallel Performance(%) (based on performance with 1PE)
1	100	-
100	1.00	100%
100	1.50	66.7% $= (1.00/1.50) \times 100$

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.
 - Granularity (粒度): Problem Size/PE



Parallel Performance

Number of PE's	Computation Time (sec)	Parallel Performance(%) (based on performance with 1PE)
1	100	-
100	1.00	100%
100	1.50	66.7% $= (1.00/1.50) \times 100$