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Message Passing Interface (MPI)

- MPI: Standard for managing communications (data and instructions) between different nodes/computers.
- Implementations: OpenMPI, MPICH2, Cray MPT, . . .
- Enables parallelism on distributed machines.
- Communicator: manages communications between processors.



MPI Operations (1 of 3)

 Managing a Communicator: Create and destroy communicators.

```
init() — initialize communicator
finalize() — shut down communicator(s)
```

 Rank query: determine the processor's position in the communicator.

```
comm.rank() — "who am I?"
comm.size() — "how many of us are there?"
```

• **Printing**: Printing output from various ranks.

```
comm.print(x)
comm.cat(x)
```

WARNING: only use these functions on *results*, never on yet-to-be-computed things.



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Quick Example 1

Rank Query: 1_rank.r

```
library(pbdMPI, quiet = TRUE)
init()

my.rank <- comm.rank()
comm.print(my.rank, all.rank=TRUE)

finalize()</pre>
```

Execute this script via:



Quick Example 2

Hello World: 2_hello.r

```
library(pbdMPI, quiet=TRUE)
init()

comm.print("Hello, world")

comm.print("Hello again", all.rank=TRUE, quiet=TRUE)

finalize()
```

Execute this script via:

mpirun -np 2 Rscript 2_hello.r

```
COMM.RANK = 0
[1] "Hello, world"
[1] "Hello again"
[1] "Hello again"
```

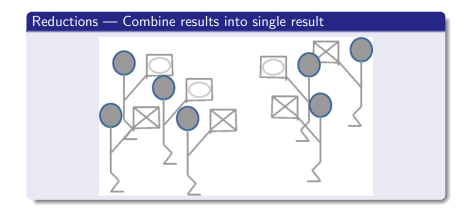


MPI Operations

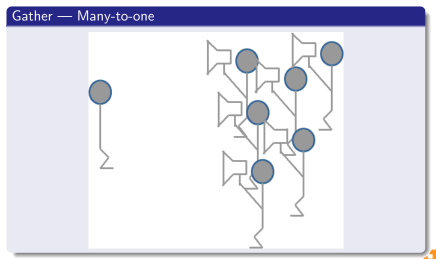
- Reduce
- Gather
- Broadcast
- Barrier



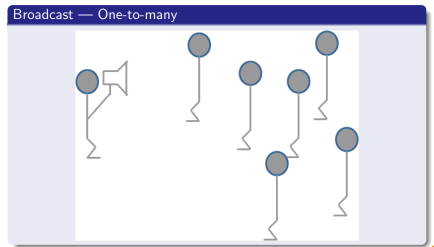
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Barrier — Synchronization Barrier Barrier Barrier



MPI Operations (2 of 3)

- Reduction: each processor has a number x; add all of them up, find the largest/smallest,
 reduce(x, op='sum') reduce to one allreduce(x, op='sum') reduce to all
- Gather: each processor has a number; create a new object on some processor containing all of those numbers.
 gather(x) — gather to one allgather(x) — gather to all
- Broadcast: one processor has a number x that every other processor should also have.
 bcast(x)
- Barrier: "computation wall"; no processor can proceed until all processors can proceed.
 barrier()



Quick Example 3

```
Reduce and Gather: 3_gt.r
```

```
library(pbdMPI, quiet = TRUE)
2 init()
  comm.set.seed(diff=TRUE)
5
  n <- sample(1:10, size=1)
  gt <- gather(n)
  comm.print(unlist(gt))
10
  sm <- allreduce(n, op='sum')</pre>
  comm.print(sm, all.rank=T)
13
14 finalize()
```

Execute this script via:

```
1 mpirun -np 2 Rscript 3_gt.r
                                        COMM.RANK = O
                                      2 [1] 2 8
                                      3 COMM.RANK = O
                                      4 [1] 10
                                      5 COMM . RANK = 1
                                      6 [1] 10
```



Quick Example 4

Broadcast: 4_bcast.r

```
library(pbdMPI, quiet=T)
init()

if (comm.rank()==0){
    x <- matrix(1:4, nrow=2)
} else {
    x <- NULL
}

or y <- bcast(x, rank.source=0)

comm.print(y, rank=1)

finalize()</pre>
```

Execute this script via:

```
mpirun -np 2 Rscript 4_bcast.r
```

```
COMM.RANK = 1
[,1] [,2]
[1,1] 1 3
[2,1] 2 4
```



15

17 18 finalize()

Reduce, Gather, Broadcast, and Barrier

Quick Example 5 Barrier: 5_barrier.r library(pbdMPI, quiet = TRUE) init() .SPMD.CT\$msg.barrier <- TRUE .SPMD.CT\$print.quiet <- TRUE for (rank in 1:comm.size()-1){ if (comm.rank() == rank){ cat(paste("Hello from process", rank+1, "of", comm.size(), "\n")) 10 barrier() 11 12 13 comm.cat("\n")

```
Execute this script via:
```

```
Sample Output:
```

```
mpirun -np 2 Rscript 5_barrier.r
                                  1 Hello from process 1
                                         of 2
                                  2 Hello from process 2
                                         of 2
```

comm.cat(paste("Hello from process", comm.rank()+1, "of", comm.size(), "\n"), all.rank=TRUE)



Quick Example 6

```
Timing: 6_timer.r
```

```
library(pbdMPI, quiet=TRUE)
2 init()
  comm.set.seed(diff=T)
  test <- function(timed)
    ltime <- system.time(timed)[3]</pre>
8
9
10
    mintime <- allreduce(ltime, op='min')
    maxtime <- allreduce(ltime, op='max')
11
12
    meantime <- allreduce(ltime, op='sum')/comm.size()
13
14
    return (data.frame (min=mintime, mean=meantime,
        max=maxtime))
15 }
16
  times <- test(rnorm(1e6)) # ~7.6MiB of data
  comm.print(times)
19
  finalize()
```

Execute this script via:

```
mpirun -np 2 Rscript 6_timer.r
```

```
1 min mean max
2 1 0.17 0.173 0.176
```



MPI Operations (3 of 3)

- Random Seeds: Random number seeds. comm.set.seed(diff=TRUE) — Independent streams via the rlecuyer package. comm.set.seed(seed=1234, diff=FALSE) — All
 - processors use the same seed.
- *ply: Functions in the *ply family. pbdApply(X, MARGIN, FUN, ...) — analogue of apply() pbdLapply(X, FUN, ...) — analogue of lapply() pbdSapply(X, FUN, ...) — analogue of sapply()



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Quick Comments for Using pbdMPI

Start by loading the package:

```
1 library(pbdMPI, quiet = TRUE)
```

② Always initialize before starting and finalize when finished:

```
1 init()
2
3 # ...
4
5 finalize()
```

