

# Package ‘pbdMPI’

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**Title** Programming with Big Data -- Interface to MPI

**Depends** R (>= 3.5.0), methods

**Imports** rlecuyer, float

**LazyLoad** yes

**Description** An efficient interface to MPI by utilizing S4

classes and methods with a focus on Single Program/Multiple Data ('SPMD')

parallel programming style, which is intended for batch parallel execution.

**SystemRequirements** OpenMPI (>= 1.5.4) on Solaris, Linux, Mac, and FreeBSD. MS-MPI (Microsoft MPI v7.1 (SDK) and Microsoft HPC Pack 2012 R2 MS-MPI Redistributable Package) on Windows.

**License** Mozilla Public License 2.0

**URL** <https://pbdr.org/>

**BugReports** <https://github.com/snoweye/pbdMPI/issues>

**MailingList** Please send questions and comments regarding pbDR to RBigData@gmail.com

**NeedsCompilation** yes

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Sebastien Lamy de la Chapelle [aut] (fix check type for send/recv long vectors)

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## Description

pbdMPI provides an efficient interface to MPI by utilizing S4 classes and methods with a focus on Single Program/Multiple Data (SPMD) parallel programming style, which is intended for batch parallel execution.

## Details

Package:	pbdMPI
Type:	Package
License:	Mozilla Public License 2.0
LazyLoad:	yes

This package requires an MPI library (OpenMPI, MPICH2, or LAM/MPI). The install command (with OpenMPI library) is

```
> tar zxvf pbdMPI_0.1-0.tar.gz
> R CMD INSTALL pbdMPI
```

Other arguments include

Argument	Default
--with-mpi-type	OPENMPI
--with-mpi-include	<code> \${MPI_ROOT}/include</code>
--with-mpi-libpath	<code> \${MPI_ROOT}/lib</code>
--with-mpi	<code> \${MPI_ROOT}</code>

where  `${MPI_ROOT}` is the path to the MPI root. See the package source file pbdMPI/configure for details.

After loading `library(pbdMPI)`, the standard process starts from `init()` which set two global variables `.comm.size` and `.comm.rank`. The standard process should end with `finalize()`.

Most functions are assumed to run in SPMD, i.e. in batch mode. Ideally, most codes run with

`mpiexec` and `Rscript`, together, such as

```
> mpiexec -np 2 Rscript some_code.r
```

where `some_code.r` contains whole SPMD program.

The package source files provide several examples based on **pbdMPI**, such as

Directory	Examples
<code>pbdMPI/inst/examples/test_spmd/</code>	major SPMD functions
<code>pbdMPI/inst/examples/test_rmpi/</code>	analog to <b>Rmpi</b>
<code>pbdMPI/inst/examples/test_parallel/</code>	analog to <b>parallel</b>
<code>pbdMPI/inst/examples/test_performance/</code>	performance tests
<code>pbdMPI/inst/examples/test_s4/</code>	S4 extension
<code>pbdMPI/inst/examples/test_cs/</code>	client/server examples
<code>pbdMPI/inst/examples/test_long_vector/</code>	long vector examples

where `test_long_vector` needs to recompile with setting

```
#define MPI_LONG_DEBUG 1
in pbdMPI/src/pkg_constant.h.
```

The current version is mainly written and tested under OpenMPI environments in Linux system (xubuntu-11.04). Also, it is tested under MPICH2 environments in Windows 7 system. It is expected to be fine for other MPI libraries and other OS platforms.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostroumov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

`allgather()`, `allreduce()`, `bcast()`, `gather()`, `reduce()`, `scatter()`.

## Examples

```
## Not run:
### Under command mode, run the demo with 2 processors by
### (Use Rscript.exe for windows system)
# mpiexec -np 2 Rscript -e "demo(allgather,'pbdMPI',ask=F,echo=F)"
# mpiexec -np 2 Rscript -e "demo(allreduce,'pbdMPI',ask=F,echo=F)"
# mpiexec -np 2 Rscript -e "demo(bcast,'pbdMPI',ask=F,echo=F)"
# mpiexec -np 2 Rscript -e "demo(gather,'pbdMPI',ask=F,echo=F)"
# mpiexec -np 2 Rscript -e "demo(reduce,'pbdMPI',ask=F,echo=F)"
# mpiexec -np 2 Rscript -e "demo(scatter,'pbdMPI',ask=F,echo=F)"
### Or
```

```
# execmpi("demo(allgather,'pbMPI',ask=F,echo=F)", nranks = 2L)
# execmpi("demo(allreduce,'pbMPI',ask=F,echo=F)", nranks = 2L)
# execmpi("demo(bcast,'pbMPI',ask=F,echo=F)", nranks = 2L)
# execmpi("demo(gather,'pbMPI',ask=F,echo=F)", nranks = 2L)
# execmpi("demo(reduce,'pbMPI',ask=F,echo=F)", nranks = 2L)
# execmpi("demo(scatter,'pbMPI',ask=F,echo=F)", nranks = 2L)

## End(Not run)
```

**allgather-method***All Ranks Gather Objects from Every Rank***Description**

This method lets all ranks gather objects from every rank in the same communicator. The default return is a list of length equal to `comm.size(comm)`.

**Usage**

```
allgather(x, x.buffer = NULL, x.count = NULL, displs = NULL,
          comm = .pb_env$SPMD.CT$comm,
          unlist = .pb_env$SPMD.CT$unlist)
```

**Arguments**

<code>x</code>	an object to be gathered from all ranks.
<code>x.buffer</code>	a buffer to hold the return object which probably has ‘size of <code>x</code> ’ times ‘ <code>comm.size(comm)</code> ’ with the same type of <code>x</code> .
<code>x.count</code>	a vector of length ‘ <code>comm size</code> ’ containing all object lengths.
<code>displs</code>	<code>c(0L,cumsum(x.count))</code> by default.
<code>comm</code>	a communicator number.
<code>unlist</code>	if <code>unlist</code> the return.

**Details**

All `x` on all ranks are likely presumed to have the same size and type.

`x.buffer`, `x.count`, and `displs` can be `NULL` or unspecified. If specified, the type should be one of integer, double, or raw specified correctly according to the type of `x`.

If `x.count` is specified, then the `spmd.allgatherv.*()` is called.

**Value**

A list of length `comm.size(comm)` is returned by default.

## Methods

For calling `spmd.allgather.*()`:

```
signature(x = "ANY", x.buffer = "missing", x.count = "missing")
signature(x = "integer", x.buffer = "integer", x.count = "missing")
signature(x = "numeric", x.buffer = "numeric", x.count = "missing")
signature(x = "raw", x.buffer = "raw", x.count = "missing")
```

For calling `spmd.allgatherv.*`:

```
signature(x = "ANY", x.buffer = "missing", x.count = "integer")
signature(x = "ANY", x.buffer = "ANY", x.count = "integer")
signature(x = "integer", x.buffer = "integer", x.count = "integer")
signature(x = "numeric", x.buffer = "numeric", x.count = "integer")
signature(x = "raw", x.buffer = "raw", x.count = "integer")
```

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[gather\(\)](#), [allreduce\(\)](#), [reduce\(\)](#).

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.file <- tempfile()
cat("
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
y <- allgather(matrix(x, nrow = 1))
comm.print(y)
```

```

y <- allgather(x, double(N * .comm.size))
comm.print(y)

### Finish.
finalize()
", file = spmd.file)
pbMPI::execmpi(spmd.file = spmd.file, nranks = 2L)

```

**allreduce-method***All Ranks Receive a Reduction of Objects from Every Rank***Description**

This method lets all ranks receive a deduction of objects from every rank in the same communicator based on a given operation. The default return is an object like the input.

**Usage**

```
allreduce(x, x.buffer = NULL, op = .pbEnv$SPMD.CT$op,
          comm = .pbEnv$SPMD.CT$comm)
```

**Arguments**

x	an object to be gathered from all ranks.
x.buffer	a buffer to hold the return object which probably has x with the same type of x.
op	a reduction operation applied to combine all x.
comm	a communicator number.

**Details**

All x on all ranks are likely presumed to have the same size and type.

x.buffer can be NULL or unspecified. If specified, the type should be one of integer, double, or raw specified correctly according to the type of x.

**Value**

The reduced object of the same type as x is returned by default.

**Methods**

For calling spmd.allreduce.\*:

```

signature(x = "ANY", x.buffer = "missing")
signature(x = "integer", x.buffer = "integer")
signature(x = "numeric", x.buffer = "numeric")
signature(x = "logical", x.buffer = "logical")
signature(x = "float32", x.buffer = "float32")

```

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[allgather\(\)](#), [gather\(\)](#), [reduce\(\)](#).

**Examples**

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
y <- allreduce(matrix(x, nrow = 1), op = "sum")
comm.print(y)

y <- allreduce(x, double(N), op = "prod")
comm.print(y)

comm.set.seed(1234, diff = TRUE)
x <- as.logical(round(runif(N)))
y <- allreduce(x, logical(N), op = "land")
comm.print(y)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code = spmd.code, nranks = 2L)
```

## Description

These functions make calls to MPI\_Alltoall() and MPI\_Alltoallv().

## Usage

```
spmd.alltoall.integer(x.send, x.recv, send.count, recv.count,
                      comm = .pbd_env$SPMD.CT$comm)
spmd.alltoall.double(x.send, x.recv, send.count, recv.count,
                      comm = .pbd_env$SPMD.CT$comm)
spmd.alltoall.raw(x.send, x.recv, send.count, recv.count,
                      comm = .pbd_env$SPMD.CT$comm)

spmd.alltoallv.integer(x.send, x.recv, send.count, recv.count,
                       sdispls, rdispls, comm = .pbd_env$SPMD.CT$comm)
spmd.alltoallv.double(x.send, x.recv, send.count, recv.count,
                       sdispls, rdispls, comm = .pbd_env$SPMD.CT$comm)
spmd.alltoallv.raw(x.send, x.recv, send.count, recv.count,
                       sdispls, rdispls, comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

x.send	an object to send.
x.recv	an object to receive
send.count	send counter
recv.count	recv counter
sdispls	send displs
rdispls	recv displs
comm	a communicator number.

## Details

These are very low level functions. Use with cautions. Neither S4 method nor long vector is supported.

## Value

These are very low level functions. Use with cautions. Neither S4 method nor long vector is supported.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[allgather\(\)](#), [allgatherv\(\)](#).

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript --vanilla [...].r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
n <- as.integer(2)
x <- 1:(.comm.size * n)
comm.cat(\"Original x:\n\", quiet = TRUE)
comm.print(x, all.rank = TRUE)

x <- as.integer(x)
y <- spmd.alltoall.integer(x, integer(length(x)), n, n)
comm.cat(\"\\nAlltoall y:\\n\", quiet = TRUE)
comm.print(y, all.rank = TRUE)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

**Description**

The functions are parallel versions of apply and lapply functions.

**Usage**

```
pbdApply(X, MARGIN, FUN, ..., pbd.mode = c("mw", "spmd", "dist"),
         rank.source = .pbd_env$SPMD.CT$rank.root,
         comm = .pbd_env$SPMD.CT$comm,
         barrier = TRUE)
pbdLapply(X, FUN, ..., pbd.mode = c("mw", "spmd", "dist"),
          rank.source = .pbd_env$SPMD.CT$rank.root,
```

```

comm = .pbd_env$SPMD.CT$comm,
bcast = FALSE, barrier = TRUE)
pbdSapply(X, FUN, ..., simplify = TRUE, USE.NAMES = TRUE,
           pbd.mode = c("mw", "spmd", "dist"),
           rank.source = .pbd_env$SPMD.CT$rank.root,
           comm = .pbd_env$SPMD.CT$comm,
           bcast = FALSE, barrier = TRUE)

```

## Arguments

X	a matrix or array in pbdApply() or a list in pbdLapply() and pbdSapply().
MARGIN	MARGIN as in the apply().
FUN	as in the apply().
...	optional arguments to FUN.
simplify	as in the sapply().
USE.NAMES	as in the sapply().
pbd.mode	mode of distributed data X.
rank.source	a rank of source where X broadcast from.
comm	a communicator number.
bcast	if bcast to all ranks.
barrier	if barrier for all ranks.

## Details

All functions are majorly called in manager/workers mode (pbd.mode1 = "mw"), and just work the same as their serial version.

If pbd.mode = "mw", the X in rank.source (master) will be redistributed to processors (workers), then apply FUN on the new data, and results are gathered to rank.source. "In SPMD, master is one of workers." ... is also scatter() from rank.source.

If pbd.mode = "spmd", the same copy of X is supposed to exist in all processors, and original apply(), lapply(), or sapply() is operated on part of X. An allgather() or gather() call is required to aggregate results manually.

If pbd.mode = "dist", the different X is supposed to exists in all processors, i.e. 'distinct or distributed' X, and original apply(), lapply(), or sapply() is operated on the all X. An allgather() or gather() call is required to aggregate results manually.

In SPMD, it is better to split data into pieces, and X is a local matrix in all processors. Originally, apply() should be sufficient in this case.

## Value

A list or matrix will be returned.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Example for pbdApply.
N <- 100
x <- matrix((1:N) + N * .comm.rank, ncol = 10)
y <- pbdApply(x, 1, sum, pbd.mode = \"mw\")
comm.print(y)

y <- pbdApply(x, 1, sum, pbd.mode = \"spmd\")
comm.print(y)

y <- pbdApply(x, 1, sum, pbd.mode = \"dist\")
comm.print(y)

### Example for pbdApply for 3D array.
N <- 60
x <- array((1:N) + N * .comm.rank, c(3, 4, 5))
dimnames(x) <- list(lat = paste(\"lat\", 1:3, sep = "\\"),
                      lon = paste(\"lon\", 1:4, sep = "\\"),
                      time = paste(\"time\", 1:5, sep = "\\"))
comm.print(x[, , 1:2])

y <- pbdApply(x, c(1, 2), sum, pbd.mode = \"mw\")
comm.print(y)

y <- pbdApply(x, c(1, 2), sum, pbd.mode = \"spmd\")
comm.print(y)

y <- pbdApply(x, c(1, 2), sum, pbd.mode = \"dist\")
comm.print(y)

### Example for pbdLapply.
N <- 100
x <- split((1:N) + N * .comm.rank, rep(1:10, each = 10))
y <- pbdLapply(x, sum, pbd.mode = \"mw\")
comm.print(unlist(y))
```

```

y <- pbdLapply(x, sum, pbd.mode = "spmd")
comm.print(unlist(y))

y <- pbdLapply(x, sum, pbd.mode = "dist")
comm.print(unlist(y))

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)

```

**bcast-method***A Rank Broadcast an Object to Every Rank***Description**

This method lets a rank broadcast an object to every rank in the same communicator. The default return is the object.

**Usage**

```
bcast(x, rank.source = .pbd_env$SPMD.CT$rank.source,
      comm = .pbd_env$SPMD.CT$comm)
```

**Arguments**

- |             |   |
|-------------|---|
| x           | an object to be broadcast from all ranks. |
| rank.source | a rank of source where x broadcast from.  |
| comm        | a communicator number.                    |

**Details**

The same copy of x is sent to all ranks.

**Value**

Every rank has x returned.

**Methods**

For calling spmd.bcast.\*:

```

signature(x = "ANY")
signature(x = "integer")
signature(x = "numeric")
signature(x = "raw")
```

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[scatter\(\)](#).

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
## Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
x <- matrix(1:5, nrow = 1)
y <- bcast(x)
comm.print(y)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)
```

## Description

The functions provide controls to communicators.

## Usage

```
barrier(comm = .pbd_env$SPMD.CT$comm)
comm.is.null(comm = .pbd_env$SPMD.CT$comm)
comm.rank(comm = .pbd_env$SPMD.CT$comm)
comm.localrank(comm = .pbd_env$SPMD.CT$comm)
comm.size(comm = .pbd_env$SPMD.CT$comm)
comm.dup(comm, newcomm)
```

```

comm.free(comm = .pbd_env$SPMD.CT$comm)
init(set.seed = TRUE)
finalize(MPI.finalize = .pbd_env$SPMD.CT$MPI.finalize)
is.finalized()

comm.abort(errorcode = 1, comm = .pbd_env$SPMD.CT$comm)
comm.split(comm = .pbd_env$SPMD.CT$comm, color = 0L, key = 0L,
           newcomm = .pbd_env$SPMD.CT$newcomm)
comm.disconnect(comm = .pbd_env$SPMD.CT$comm)
comm.connect(port.name, info = .pbd_env$SPMD.CT$info,
             rank.root = .pbd_env$SPMD.CT$rank.root,
             comm = .pbd_env$SPMD.CT$comm,
             newcomm = .pbd_env$SPMD.CT$newcomm)
comm.accept(port.name, info = .pbd_env$SPMD.CT$info,
            rank.root = .pbd_env$SPMD.CT$rank.root,
            comm = .pbd_env$SPMD.CT$comm,
            newcomm = .pbd_env$SPMD.CT$newcomm)

port.open(info = .pbd_env$SPMD.CT$info)
port.close(port.name)
serv.publish(port.name, serv.name = .pbd_env$SPMD.CT$serv.name,
             info = .pbd_env$SPMD.CT$info)
serv.unpublish(port.name, serv.name = .pbd_env$SPMD.CT$serv.name,
               info = .pbd_env$SPMD.CT$info)
serv.lookup(serv.name = .pbd_env$SPMD.CT$serv.name,
            info = .pbd_env$SPMD.CT$info)

intercomm.merge(intercomm = .pbd_env$SPMD.CT$intercomm,
                high = 0L, comm = .pbd_env$SPMD.CT$comm)
intercomm.create(local.comm = .pbd_env$SPMD.CT$comm,
                 local.leader = .pbd_env$SPMD.CT$rank.source,
                 peer.comm = .pbd_env$SPMD.CT$intercomm,
                 remote.leader = .pbd_env$SPMD.CT$rank.dest,
                 tag = .pbd_env$SPMD.CT$tag,
                 newintercomm = .pbd_env$SPMD.CT$newcomm)

comm.c2f(comm = .pbd_env$SPMD.CT$comm)

```

## Arguments

comm	a communicator number.
mpi.finalize	if MPI should be shutdown.
set.seed	if a random seed preset.
port.name	a port name with default maximum length 1024 characters for OpenMPI.
info	a info number.
rank.root	a root rank.
newcomm	a new communicator number.

color	control of subset assignment.
key	control of rank assignment.
serv.name	a service name.
errorcode	an error code to abort MPI.
intercomm	a intercommunicator number.
high	used to order the groups within comm.
local.comm	a local communicator number.
local.leader	the leader number of local communicator.
peer.comm	a peer communicator number.
remote.leader	the remote leader number of peer communicator.
newintercomm	a new intercommunicator number.
tag	a tag number.

## Details

Another functions are direct calls to MPI library.

`barrier()` blocks all processors until everyone call this.

`comm.is.null()` returns -1 if the array of communicators is not allocated, i.e. `init()` is not called yet. It returns 1 if the communicator is not initialized, i.e. `NULL`. It returns 0 if the communicator is initialized.

`comm.rank()` returns the processor's rank for the given `comm`.

`comm.size()` returns the total processes for the given `comm`.

`comm.dup()` duplicate a `newcomm` from `comm`.

`comm.free()` free a `comm`.

`init()` initializes a MPI world, and set two global variables `.comm.size` and `.comm.rank` in `.GlobalEnv`. A random seed will be preset by default (`Sys.getpid() + Sys.time()`) to the package **rlecuyer**.

`finalize()` frees memory and finishes a MPI world if `mpi.finalize = TRUE`. `is.finalized()` checks if MPI is already finalized.

`comm.abort()` aborts MPI.

`comm.split()` create a `newcomm` by color and key.

`comm.disconnect()` frees a `comm`.

`comm.connect()` connects a `newcomm`.

`comm.accept()` accepts a `newcomm`.

`port.open()` opens a port and returns the port name.

`port.close()` closes a port by name.

`serv.publish()` publishes a service via `port.name`.

`serv.unpublish()` unpublishes a service via `port.name`.

`serv.lookup()` lookup the `serv.name` and returns the port name.

`intercomm.merge()` merges the `intercomm` to intracommunicator.

`intercomm.create()` creates a new `intercomm` from two peer intracommunicators.

`comm.c2f()` returns an integer for Fortran MPI support.

**Value**

Most function return an invisible state of MPI call.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples .
comm.print(.comm.size)
comm.print(.comm.rank, all.rank = TRUE)
comm.print(comm.rank(), rank.print = 1)
comm.print(comm.c2f())

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

**Description**

This method lets a rank gather objects from every rank in the same communicator. The default return is a list of length equal to ‘comm size’.

**Usage**

```
gather(x, x.buffer = NULL, x.count = NULL, displs = NULL,
       rank.dest = .pbd_env$SPMD.CT$rank.root,
       comm = .pbd_env$SPMD.CT$comm,
       unlist = .pbd_env$SPMD.CT$unlist)
```

**Arguments**

x	an object to be gathered from all ranks.
x.buffer	a buffer to hold the return object which probably has ‘size of x’ times ‘comm size’ with the same type of x.
x.count	a vector of length ‘comm size’ containing all object lengths.
displs	c(0L, cumsum(x.count)) by default.
rank.dest	a rank of destination where all x gather to.
comm	a communicator number.
unlist	if unlist the return.

**Details**

All x on all ranks are likely presumed to have the same size and type.

x.buffer, x.count, and displs can be NULL or unspecified. If specified, the type should be one of integer, double, or raw specified correctly according to the type of x.

If x.count is specified, then the spmd.gatherv.\*() is called.

**Value**

If rank.dest == comm.rank(comm), then a list of length ‘comm size’ is returned by default. Otherwise, NULL is returned.

**Methods**

For calling spmd.gather.\*():

```
signature(x = "ANY", x.buffer = "missing", x.count = "missing")
signature(x = "integer", x.buffer = "integer", x.count = "missing")
signature(x = "numeric", x.buffer = "numeric", x.count = "missing")
signature(x = "raw", x.buffer = "raw", x.count = "missing")
```

For calling spmd.gatherv.\*():

```
signature(x = "ANY", x.buffer = "missing", x.count = "integer")
signature(x = "ANY", x.buffer = "ANY", x.count = "integer")
signature(x = "integer", x.buffer = "integer", x.count = "integer")
signature(x = "numeric", x.buffer = "numeric", x.count = "integer")
signature(x = "raw", x.buffer = "raw", x.count = "integer")
```

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[gather\(\)](#), [allreduce\(\)](#), [reduce\(\)](#).

**Examples**

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
y <- gather(matrix(x, nrow = 1))
comm.print(y)
y <- gather(x, double(N * .comm.size))
comm.print(y)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)
```

**Description**

These functions are designed to get MPI and/or pbdMPI configures that were usually needed at the time of pbdMPI installation. In particular, to configure, link, and compile with ‘libmpi\*.so’ or so.

## Usage

```
get.conf(arg, arch = '', package = "pbMPI", return = FALSE)
get.lib(arg, arch, package = "pbPROF")
get.sysenv(flag)
```

## Arguments

arg	an argument to be searched in the configuration file
arch	system architecture
package	pakge name
return	to return (or print if FALSE) the search results or not
flag	a system flag that is typically used in windows environment set.

## Details

get.conf() and get.lib() are typically used by ‘pb\*/configure.ac’, ‘pb\*/src/Makevars.in’, and/or ‘pb\*/src/Makevar.win’ to find the default configurations from ‘pb\*/etc\$R\_ARCH/Makconf’. get.sysenv() is only called by ‘pbMPI/src/Makevars.win’ to obtain possible MPI dynamic/static library from the environment variable ‘MPI\_ROOT’ preset by users.

## Value

Typically, there are no return values, but the values are cat() to scrrn or stdin.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:
library(pbMPI)
if(Sys.info()["sysname"] != "Windows"){
  get.conf("MPI_INCLUDE_PATH"); cat("\n")
  get.conf("MPI_LIBPATH"); cat("\n")
  get.conf("MPI_LIBNAME"); cat("\n")
  get.conf("MPI_LIBS"); cat("\n")
} else{
  get.conf("MPI_INCLUDE", "/i386"); cat("\n")
  get.conf("MPI_LIB", "/i386"); cat("\n")

  get.conf("MPI_INCLUDE", "/x64"); cat("\n")
  get.conf("MPI_LIB", "/x64"); cat("\n")
}
```

---

```
## End(Not run)
```

---

<code>get.job.id</code>	<i>Divide Job ID by Ranks</i>
-------------------------	-------------------------------

---

### Description

This function obtains job id which can be used to divide jobs.

### Usage

```
get.jid(n, method = .pbd_env$SPMD.CT$divide.method[1], all = FALSE,
       comm = .pbd_env$SPMD.CT$comm, reduced = FALSE)
```

### Arguments

<code>n</code>	total number of jobs.
<code>method</code>	a way to divide jobs.
<code>all</code>	indicate if return all id for each processor.
<code>comm</code>	a communicator number.
<code>reduced</code>	indicate if return should be a reduced representation.

### Details

`n` is total number of jobs needed to be divided into all processors (`comm.size(comm)`), i.e. `1:n` will be split according to the rank of processor (`comm.rank(comm)`) and `method`. Job id will be returned. Currently, three possible methods are provided.

"block" will use return id's which are nearly equal size blocks. For example, 7 jobs in 4 processors will have `jid=1` for rank 0, `jid=2,3` for rank 1, `jid=4,5` for rank 2, and `jid=6,7` for rank 3.

"block0" will use return id's which are nearly equal size blocks, in the opposite direction of "block". For example, 7 jobs in 4 processors will have `jid=1,2` for rank 0, `jid=3,4` for rank 1, `jid=5,6` for rank 2, and `jid=7` for rank 3.

"cycle" will use return id's which are nearly equal size in cycle. For example, 7 jobs in 4 processors will have `jid=1,5` for rank 0, `jid=2,6` for rank 1, `jid=3,7` for rank 2, and `jid=4` for rank 3.

### Value

`get.id()` returns a vector containing job id for each individual processor if `all = FALSE`. While it returns a list containing all job id for all processor if `all = TRUE`. The list has length equal to `COMM.SIZE`.

### Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[task.pull\(\)](#).

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
comm.cat(>>> block\n", quiet = TRUE)
jid <- get.jid(7, method = "block")
comm.print(jid, all.rank = TRUE)

comm.cat(>>> cycle\n", quiet = TRUE)
jid <- get.jid(7, method = "cycle")
comm.print(jid, all.rank = TRUE)

comm.cat(>>> block (all)\n", quiet = TRUE)
alljid <- get.jid(7, method = "block", all = TRUE)
comm.print(alljid)

comm.cat(>>> cycle (all)\n", quiet = TRUE)
alljid <- get.jid(7, method = "cycle", all = TRUE)
comm.print(alljid)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)
```

## Description

This function provide global all pairs.

**Usage**

```
comm.allpairs(N, diag = FALSE, symmetric = TRUE,
              comm = .pbd_env$SPMD.CT$comm)
```

**Arguments**

N	number of elements for matching, (i, j) for all $1 \leq i, j \leq N$ .
diag	if matching the same elements, (i, i) for all i.
symmetric	if matching upper triangular elements. TRUE for $i \geq j$ only, otherwise for all (i, j).
comm	a communicator number.

**Details**

The function generates all combinations of N elements.

**Value**

The function returns a gbd matrix in row blocks with 2 columns named i and j. The number of rows is dependent on the options `diag` and `symmetric`. If `diag = TRUE` and `symmetric = FALSE`, then this case has the maximum number of rows,  $N^2$ .

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

`comm.dist()`.

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
id.matrix <- comm.allpairs(comm.size() + 1)
comm.print(id.matrix, all.rank = TRUE)
```

```
### Finish.
finalize()
"
# execmpi(spmr.code, nranks = 2L)

## End(Not run)
```

**global any and all      *Global Any and All Functions***

## Description

These functions are global any and all applying on distributed data for all ranks.

## Usage

```
comm.any(x, na.rm = FALSE, comm = .pbd_env$SPMD.CT$comm)
comm.all(x, na.rm = FALSE, comm = .pbd_env$SPMD.CT$comm)

comm.allcommon(x, comm = .pbd_env$SPMD.CT$comm,
               lazy.check = .pbd_env$SPMD.CT$lazy.check)
```

## Arguments

x	a vector.
na.rm	if NA removed or not.
comm	a communicator number.
lazy.check	if TRUE, then allreduce is used to check all ranks, otherwise, allgather is used.

## Details

These functions will apply `any()` and `all()` locally, and apply `allgather()` to get all local results from other ranks, then apply `any()` and `all()` on all local results.

`comm.allcommon()` is to check if `x` is exactly the same across all ranks. This is a vectorized operation on `x` where the input and output have the same length of vector, while `comm.any()` and `comm.all()` return a scalar.

Note that `lazy.check = TRUE` is faster as number of cores is large, but it may cause some inconsistency in some cases. `lazy.check = FALSE` is much slower, but it provides more accurate checking.

## Value

The global check values (TRUE, FALSE, NA) are returned to all ranks.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
if(comm.rank() == 0){
  a <- c(T, F, NA)
} else{
  a <- T
}

comm.any(a)
comm.all(a)
comm.any(a, na.rm = TRUE)
comm.all(a, na.rm = TRUE)

comm.allcommon(1:3)
if(comm.rank() == 0){
  a <- 1:3
} else{
  a <- 3:1
}
comm.allcommon.integer(a)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

## Description

This function redistributes a regular matrix existed in rank.soure and turns it in a gbd matrix in row blocks.

## Usage

```
comm.as.gbd(X, balance.method = .pbdr_env$SPMD.IO$balance.method,
             rank.source = .pbdr_env$SPMD.CT$rank.source,
             comm = .pbdr_env$SPMD.CT$comm)
```

## Arguments

X a regular matrix in rank.source and to be redistributed as a gbd.  
 balance.method a balance method.  
 rank.source a rank of source where elements of X scatter from.  
 comm a communicator number.

## Details

X matrix in rank.source will be redistributed as a gbd matrix in row blocks.

This function will first set NULL to X if it is not located in rank.source, then called [comm.load.balance\(\)](#) to redistribute the one located in rank.source to all other ranks.

## Value

A X.gbd will be returned.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[comm.load.balance\(\)](#), [comm.read.table\(\)](#) and [comm.write.table\(\)](#).

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
X <- matrix(1:15, ncol = 3)
X.gbd <- comm.as.gbd(X)
```

```

comm.print(X.gbd, all.rank = TRUE)

### Finish.
finalize()
"
pbMPI::execmpi(spmd.code, nranks = 2L)

```

## Description

These functions are global balance methods for gbd data.frame (or matrix) distributed in row blocks.

## Usage

```

comm.balance.info(X.gbd, balance.method = .pb_env$SPMD.IO$balance.method[1],
                  comm = .pb_env$SPMD.CT$comm)
comm.load.balance(X.gbd, bal.info = NULL,
                  balance.method = .pb_env$SPMD.IO$balance.method[1],
                  comm = .pb_env$SPMD.CT$comm)
comm.unload.balance(new.X.gbd, bal.info, comm = .pb_env$SPMD.CT$comm)

```

## Arguments

X.gbd	a gbd data.frame (or matrix).
balance.method	a balance method.
bal.info	a balance information returned from comm.balance.info(). If NULL, then this will be generated inside comm.load.balance().
new.X.gbd	a new gbd of X.gbd (may be generated from comm.load.balance()).
comm	a communicator number.

## Details

A typical use is to balance an input dataset X.gbd from `comm.read.table()`. Since by default, a two dimension data.frame is distributed in row blocks, but each processor (rank) may not (or closely) have the same number of rows. These functions redistribute the data.frame (and maybe matrix) according to the specified way in bal.info.

Currently, there are three balance methods are supported, block (uniform distributed but favor higher ranks), block0 (as block but favor lower ranks), and block.cyclic (as block cyclic with one big block in one cycle).

**Value**

`comm.balance.info()` returns a list containing balance information based on the input `X.gbd` and `balance.method`.

`comm.load.balance()` returns a new gbd `data.frame` (or `matrix`).

`comm.unload.balance()` also returns the new gbd `data.frame` back to the original `X.gbd`.

**Author(s)**

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

`comm.read.table()`, `comm.write.table()`, and `comm.as.gbd()`.

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))

### Get two gbd row-block data.frame.
da.block <- iris[get.jid(nrow(iris), method = \"block\"),]
da.block0 <- iris[get.jid(nrow(iris), method = \"block0\"),]

### Load balance one and unload it.
bal.info <- comm.balance.info(da.block0)
da.new <- comm.load.balance(da.block0)
da.org <- comm.unload.balance(da.new, bal.info)

### Check if all are equal.
comm.print(c(sum(da.new != da.block), sum(da.org != da.block0)),
           all.rank = TRUE)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

**Description**

These functions are global base functions applying on distributed data for all ranks.

**Usage**

```
comm.length(x, comm = .pbdr_env$SPMD.CT$comm)
comm.sum(..., na.rm = TRUE, comm = .pbdr_env$SPMD.CT$comm)
comm.mean(x, na.rm = TRUE, comm = .pbdr_env$SPMD.CT$comm)
comm.var(x, na.rm = TRUE, comm = .pbdr_env$SPMD.CT$comm)
comm.sd(x, na.rm = TRUE, comm = .pbdr_env$SPMD.CT$comm)
```

**Arguments**

x	a vector.
...	as in sum().
na.rm	logical, if remove NA and NaN.
comm	a communicator number.

**Details**

These functions will apply globally length(), sum(), mean(), var(), and sd().

**Value**

The global values are returned to all ranks.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdrMPI, quietly = TRUE))
```

```

init()
if(comm.size() != 2){
  comm.cat("2 processors are required.\n", quiet = TRUE)
  finalize()
}

#### Examples.
a <- 1:(comm.rank() + 1)

b <- comm.length(a)
comm.print(b)
b <- comm.sum(a)
comm.print(b)
b <- comm.mean(a)
comm.print(b)
b <- comm.var(a)
comm.print(b)
b <- comm.sd(a)
comm.print(b)

#### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)

```

**global distance function***Global Distance for Distributed Matrices***Description**

These functions globally compute distance for all ranks.

**Usage**

```
comm.dist(X.gbd, method = "euclidean", diag = FALSE, upper = FALSE,
          p = 2, comm = .pbd_env$SPMD.CT$comm,
          return.type = c("common", "gbd"))
```

**Arguments**

X.gbd	a gbd matrix.
method	as in <code>dist()</code> .
diag	as in <code>dist()</code> .
upper	as in <code>dist()</code> .
p	as in <code>dist()</code> .

<code>comm</code>	a communicator number.
<code>return.type</code>	returning type for the distance.

## Details

The distance function is implemented for a distributed matrix.

The return type `common` is only useful when the number of rows of the matrix is small since the returning matrix is  $N \times N$  for every rank where  $N$  is the total number of rows of `X.gbd` of all ranks.

The return type `gbd` returns a `gbd` matrix (distributed across all ranks, and the `gbd` matrix has 3 columns, named "`i`", "`j`", and "`value`", where  $(i, j)$  is the global indices of the  $i$ -th and  $j$ -th rows of `X.gbd`, and `value` is the corresponding distance. The  $(i, j)$  is ordered as a distance matrix.

## Value

A full distance matrix is returned from the `common` return type. Suppose  $N.gbd$  is total rows of `X.gbd`, then the distance will have  $N.gbd * (N.gbd - 1) / 2$  elements and the distance matrix will have  $N.gbd^2$  elements.

A `gbd` distance matrix with 3 columns is returned from the `gbd` return type.

## Warning

The distance or distance matrix could be huge.

## Author(s)

Wei-Chen Chen <[wccsnw@gmail.com](mailto:wccsnw@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

`comm.allpairs()` and `comm.pairwise()`.

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
comm.set.seed(123456, diff = TRUE)
```

```

X.gbd <- matrix(runif(6), ncol = 3)
dist.X.common <- comm.dist(X.gbd)
dist.X.gbd <- comm.dist(X.gbd, return.type = "gbd")

### Verify.
dist.X <- dist(do.call("rbind", allgather(X.gbd)))
comm.print(all(dist.X == dist.X.common))

### Verify 2.
dist.X.df <- do.call("rbind", allgather(dist.X.gbd))
comm.print(all(dist.X == dist.X.df[, 3]))
comm.print(dist.X)
comm.print(dist.X.df)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)

```

**global match.arg**      *Global Argument Matching*

## Description

A binding for `match.arg()` that uses `comm.stop()` rather so that the error message (if there is one) is managed according to the rules of `.pbd_env$SPMD.CT`.

## Usage

```

comm.match.arg(arg, choices, several.ok=FALSE, ...,
               all.rank = .pbd_env$SPMD.CT$print.all.rank,
               rank.print = .pbd_env$SPMD.CT$rank.source,
               comm = .pbd_env$SPMD.CT$comm,
               mpi.finalize = .pbd_env$SPMD.CT$mpi.finalize,
               quit = .pbd_env$SPMD.CT$quit)

```

## Arguments

arg,choices,several.ok	see <code>match.arg()</code>
...	ignored.
all.rank	if all ranks print (default = FALSE).
rank.print	rank for printing if not all ranks print (default = 0).
comm	communicator for printing (default = 1).
mpi.finalize	if MPI should be shutdown.
quit	if quit R when errors happen.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

global pairwise      *Global Pairwise Evaluations*

**Description**

This function provides global pairwise evaluations.

**Usage**

```
comm.pairwise(X, pairid.gbd = NULL,
              FUN = function(x, y, ...){ return(as.vector(dist(rbind(x, y), ...))) },
              ..., diag = FALSE, symmetric = TRUE, comm = .pbd_env$SPMD.CT$comm)
```

**Arguments**

X	a common matrix across ranks, or a gbd matrix. (See details.)
pairid.gbd	a pair-wise id in a gbd format. (See details.)
FUN	a function to be evaluated for given pairs.
...	extra variables for FUN.
diag	if matching the same elements, (i, i) for all i.
symmetric	if matching upper triangular elements. TRUE for i >= j only, otherwise for all (i, j).
comm	a communicator number.

**Details**

This function evaluates the objective function  $\text{FUN}(X[i,], X[j,])$  (usually distance of two elements) on any given pair  $(i, j)$  of a matrix  $X$ .

The input  $X$  should be in common across all ranks if `pairid.gbd` is provided, e.g. from `comm.pairwise()`. i.e.  $X$  is exactly the same in every ranks, but `pairid.gbd` is different and in gbd format indicating the row pair  $(i, j)$  should be evaluated. The returning gbd matrix is ordered and indexed by `pairid.gbd`.

Note that checking consistence of  $X$  across all ranks is not implemented within this function since that drops performance and may be not accurate.

The input  $X$  should be a gbd format in row major blocks (i.e.  $X.gbd$ ) if `pairid.gbd` is `NULL`. A internal pair indices will be built implicitly for evaluation. The returning gbd matrix is ordered and indexed by `X.gbd`.

**Value**

This function returns a common matrix with 3 columns named *i*, *j*, and *value*. Each *value* is the returned value and computed by `FUN(X[i,],X[j,])` where  $(i,j)$  is the global index as ordered in a distance matrix for *i*-th row and *j*-th columns.

**Author(s)**

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

`comm.pairwise()`, and `comm.dist()`.

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
comm.set.seed(123456, diff = FALSE)
X <- matrix(rnorm(10), ncol = 2)
id.matrix <- comm.allpairs(nrow(X))

### Method original.
dist.org <- dist(X)

### Method 1.
dist.common <- comm.pairwise(X, pairid.gbd = id.matrix)

### Method 2.
# if(comm.rank() != 0){
#   X <- matrix(0, nrow = 0, ncol = 4)
# }
X.gbd <- comm.as.gbd(X)    ### The other way.
dist.gbd <- comm.pairwise(X.gbd)

### Verify.
d.org <- as.vector(dist.org)
d.1 <- do.call(\"c\", allgather(dist.common[, 3]))
d.2 <- do.call(\"c\", allgather(dist.gbd[, 3]))
```

```

comm.print(all(d.org == d.1))
comm.print(all(d.org == d.2))

### Finish.
finalize()
"
# execmpi(spmr.code, nranks = 2L)

## End(Not run)

```

global print and cat    *Global Print and Cat Functions*

### Description

The functions globally print or cat a variable from specified processors, by default messages is shown on screen.

### Usage

```

comm.print(x, all.rank = .pbd_env$SPMD.CT$print.all.rank,
           rank.print = .pbd_env$SPMD.CT$rank.source,
           comm = .pbd_env$SPMD.CT$comm,
           quiet = .pbd_env$SPMD.CT$print.quiet,
           flush = .pbd_env$SPMD.CT$msg.flush,
           barrier = .pbd_env$SPMD.CT$msg.barrier,
           con = stdout(), ...)

comm.cat(..., all.rank = .pbd_env$SPMD.CT$print.all.rank,
          rank.print = .pbd_env$SPMD.CT$rank.source,
          comm = .pbd_env$SPMD.CT$comm,
          quiet = .pbd_env$SPMD.CT$print.quiet, sep = " ", fill = FALSE,
          labels = NULL, append = FALSE, flush = .pbd_env$SPMD.CT$msg.flush,
          barrier = .pbd_env$SPMD.CT$msg.barrier, con = stdout())

```

### Arguments

x	a variable to be printed.
...	variables to be cat.
all.rank	if all ranks print (default = FALSE).
rank.print	rank for printing if not all ranks print (default = 0).
comm	communicator for printing (default = 1).
quiet	FALSE for printing rank number.
sep	sep argument as in the cat() function.
fill	fill argument as in the cat() function.

labels	labels argument as in the <code>cat()</code> function.
append	labels argument as in the <code>cat()</code> function.
flush	if flush con.
barrier	if barrier con.
con	<code>stdout()</code> is the default to print message.

## Details

**Warning:** These two functions use `barrier()` to make sure the well printing process on screen, so should be called by all processors to avoid a deadlock. A typical misuse is called inside a condition check, such as `if(.comm.rank == 0) comm.cat(...)`.

`rank.print` can be a integer vector containing the ranks of processors which print messages.

## Value

A `print()` or `cat()` is called for the specified processors and the messages of the input variables is shown on screen by default.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Example.
comm.print(comm.rank(), rank.print = 1)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

global range, max, and min

*Global Range, Max, and Min Functions*

## Description

These functions are global range, max and min applying on distributed data for all ranks.

## Usage

```
comm.range(..., na.rm = FALSE, comm = .pbdr_env$SPMD.CT$comm)
comm.max(..., na.rm = FALSE, comm = .pbdr_env$SPMD.CT$comm)
comm.min(..., na.rm = FALSE, comm = .pbdr_env$SPMD.CT$comm)
```

## Arguments

...	an 'numeric' objects.
na.rm	if NA removed or not.
comm	a communicator number.

## Details

These functions will apply `range()`, `max()` and `min()` locally, and apply `allgather` to get all local results from other ranks, then apply `range()`, `max()` and `min()` on all local results.

## Value

The global values (range, max, or min) are returned to all ranks.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
```

```

if(comm.size() != 2){
  comm.cat("2 processors are required.\n", quiet = TRUE)
  finalize()
}

#### Examples.
a <- 1:(comm.rank() + 1)

b <- comm.range(a)
comm.print(b)
b <- comm.max(a)
comm.print(b)
b <- comm.min(a)
comm.print(b)

#### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)

```

## global reading

*Global Reading Functions***Description**

These functions are global reading from specified file.

**Usage**

```

comm.read.table(file, header = FALSE, sep = "", quote = "\"\"",
                dec = ".",
                na.strings = "NA", colClasses = NA, nrows = -1, skip = 0,
                check.names = TRUE, fill = !blank.lines.skip,
                strip.white = FALSE,
                blank.lines.skip = TRUE, comment.char = "#",
                allowEscapes = FALSE,
                flush = FALSE,
                fileEncoding = "", encoding = "unknown",
                read.method = .pbd_env$SPMD.IO$read.method[1],
                balance.method = .pbd_env$SPMD.IO$balance.method[1],
                comm = .pbd_env$SPMD.CT$comm)

comm.read.csv(file, header = TRUE, sep = ",", quote = "\"\",
                dec = ".", fill = TRUE, comment.char = "", ...,
                read.method = .pbd_env$SPMD.IO$read.method[1],
                balance.method = .pbd_env$SPMD.IO$balance.method[1],
                comm = .pbd_env$SPMD.CT$comm)

```

```
comm.read.csv2(file, header = TRUE, sep = ";", quote = "\"",
               dec = ",", fill = TRUE, comment.char = "", ...,
               read.method = .pbd_env$SPMD.IO$read.method[1],
               balance.method = .pbd_env$SPMD.IO$balance.method[1],
               comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

file	as in <code>read.table()</code> .
header	as in <code>read.table()</code> .
sep	as in <code>read.table()</code> .
quote	as in <code>read.table()</code> .
dec	as in <code>read.table()</code> .
na.strings	as in <code>read.table()</code> .
colClasses	as in <code>read.table()</code> .
nrows	as in <code>read.table()</code> .
skip	as in <code>read.table()</code> .
check.names	as in <code>read.table()</code> .
fill	as in <code>read.table()</code> .
strip.white	as in <code>read.table()</code> .
blank.lines.skip	as in <code>read.table()</code> .
comment.char	as in <code>read.table()</code> .
allowEscapes	as in <code>read.table()</code> .
flush	as in <code>read.table()</code> .
fileEncoding	as in <code>read.table()</code> .
encoding	as in <code>read.table()</code> .
...	as in <code>read.csv*()</code> .
read.method	either "gbd" or "common".
balance.method	balance method for <code>read.method = "gbd"</code> as <code>nrows = -1</code> and <code>skip = 0</code> are set.
comm	a communicator number.

## Details

These functions will apply `read.table()` locally and sequentially from rank 0, 1, 2, ...

By default, rank 0 reads the file only, then scatter to other ranks for small datasets (`.pbd_env$SPMD.IO$max.read.size`) in `read.method = "gbd"`. (bcast to others in `read.method = "common"`.)

As dataset size increases, the reading is performed from each ranks and read portion of rows in "gbd" format as described in **pbdDEMO** vignettes and used in **pmclust**.

`comm.load.balance()` is called for "gbd" method as as `nrows = -1` and `skip = 0` are set. Note that the default method "block" is the better way for performance in general that distributes equally and leaves residuals on higher ranks evenly. "block0" is the other way around. "block.cyclic" is only useful for converting to `ddmatrix` as in **pbdDMAT**.

**Value**

A distributed data.frame is returned.

All factors are disable and read as characters or as what data should be.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

`comm.load.balance()` and `comm.write.table()`

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))

### Check.
if(comm.size() != 2){
  comm.stop(\"2 processors are required.\")
}

### Manually distributed iris.
da <- iris[get.jid(nrow(iris)),]

### Dump data.
comm.write.table(da, file = \"iris.txt\", quote = FALSE, sep = "\\t",
                 row.names = FALSE)

### Read back in.
da.gbd <- comm.read.table(\"iris.txt\", header = TRUE, sep = "\\t",
                           quote = "\\")
comm.print(c(nrow(da), nrow(da.gbd)), all.rank = TRUE)

### Read in common.
da.common <- comm.read.table(\"iris.txt\", header = TRUE, sep = "\\t",
                           quote = "\\", read.method = "common")
comm.print(c(nrow(da.common), sum(da.common != iris)))

### Finish.
finalize()
```

```
"  
# execmpi(spmd.code, nnranks = 2L)  
## End(Not run)
```

---

**global Rprof***A Rprof Function for SPMD Routines*

---

**Description**

A Rprof function for use with parallel codes executed in the batch SPMD style.

**Usage**

```
comm.Rprof(filename = "Rprof.out", append = FALSE, interval = 0.02,  
          memory.profiling = FALSE, gc.profiling = FALSE,  
          line.profiling = FALSE, numfiles = 100L, bufsize = 10000L,  
          all.rank = .pbd_env$SPMD.CT$Rprof.all.rank,  
          rank.Rprof = .pbd_env$SPMD.CT$rank.source,  
          comm = .pbd_env$SPMD.CT$comm)
```

**Arguments**

filename	as in <a href="#">Rprof()</a> .
append	as in <a href="#">Rprof()</a> .
interval	as in <a href="#">Rprof()</a> .
memory.profiling	as in <a href="#">Rprof()</a> .
gc.profiling	as in <a href="#">Rprof()</a> .
line.profiling	as in <a href="#">Rprof()</a> .
numfiles	as in <a href="#">Rprof()</a> .
bufsize	as in <a href="#">Rprof()</a> .
all.rank	if calling Rprof on all ranks (default = FALSE).
rank.Rprof	rank for calling Rprof if all.rank = FALSE (default = 0).
comm	a communicator number.

**Details**

as in [Rprof\(\)](#).

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

`global sort`

*Global Quick Sort for Distributed Vectors or Matrices*

## Description

This function globally sorts distributed data for all ranks.

## Usage

```
comm.sort(x, decreasing = FALSE, na.last = NA,
          comm = .pbd_env$SPMD.CT$comm,
          status = .pbd_env$SPMD.CT$status)
```

## Arguments

<code>x</code>	a vector.
<code>decreasing</code>	logical. Should the sort order be increasing or decreasing?
<code>na.last</code>	for controlling the treatment of NAs. If TRUE, missing values in the data are put last; if FALSE, they are put first; if NA, they are removed.
<code>comm</code>	a communicator number.
<code>status</code>	a status number.

## Details

The distributed quick sort is implemented for this functions.

## Value

The returns are the same size of `x` but in global sorting order.

## Warning

All ranks may not have a NULL `x`.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```

## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
comm.set.seed(123456, diff = TRUE)
x <- c(rnorm(5 + .comm.rank * 2), NA)
# x <- sample(1:5, 5 + .comm.rank * 2, replace = TRUE)
comm.end.seed()

if(.comm.rank == 1){
  x <- NULL      ### Test for NULL or 0 vector
}

y <- allgather(x)
comm.print(y)

y <- comm.sort(x)
y <- allgather(y)
comm.print(y)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)

```

global stop and warning

*Global Stop and Warning Functions*

## Description

These functions are global stop and warning applying on distributed data for all ranks, and are called by experts only. These functions may lead to potential performance degradation and system termination.

## Usage

```
comm.stop(..., call. = TRUE, domain = NULL,
```

```

all.rank = .pbd_env$SPMD.CT$print.all.rank,
rank.print = .pbd_env$SPMD.CT$rank.source,
comm = .pbd_env$SPMD.CT$comm,
mpi.finalize = .pbd_env$SPMD.CT$mpi.finalize,
quit = .pbd_env$SPMD.CT$quit)

comm.warning(..., call. = TRUE, immediate. = FALSE, domain = NULL,
             all.rank = .pbd_env$SPMD.CT$print.all.rank,
             rank.print = .pbd_env$SPMD.CT$rank.source,
             comm = .pbd_env$SPMD.CT$comm)

comm.warnings(...,
              all.rank = .pbd_env$SPMD.CT$print.all.rank,
              rank.print = .pbd_env$SPMD.CT$rank.source,
              comm = .pbd_env$SPMD.CT$comm)

comm.stopifnot(..., call. = TRUE, domain = NULL,
               all.rank = .pbd_env$SPMD.CT$print.all.rank,
               rank.print = .pbd_env$SPMD.CT$rank.source,
               comm = .pbd_env$SPMD.CT$comm,
               mpi.finalize = .pbd_env$SPMD.CT$mpi.finalize,
               quit = .pbd_env$SPMD.CT$quit)

```

## Arguments

...	variables to be cat.
call.	see stop() and warnings().
immediate.	see stop() and warnings().
domain	see stop() and warnings().
all.rank	if all ranks print (default = FALSE).
rank.print	rank for printing if not all ranks print (default = 0).
comm	communicator for printing (default = 1).
mpi.finalize	if MPI should be shutdown.
quit	if quit R when errors happen.

## Details

These functions will respectively apply stop(), warning(), warnings(), and stopifnot() locally.

## Value

comm.stop() and comm.stopifnot() terminate all ranks, comm.warning() returns messages, and comm.warnings() print the message.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
if(comm.size() != 2){
  comm.cat(\"2 processors are required.\n\", quiet = TRUE)
  finalize()
}

### Examples.
comm.warning(\"test warning.\n\")
comm.warnings()
comm.stop(\"test stop.\n\")
comm.stopifnot(1 == 2)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

**Description**

A timing function for use with parallel codes executed in the batch SPMD style.

**Usage**

```
comm.timer(timed, comm = .pbd_env$SPMD.CT$comm)
```

**Arguments**

- timed** expression to be timed.  
**comm** a communicator number.

**Details**

Finds the min, mean, and max execution time across all independent processes executing the operation **timed**.

**Author(s)**

Drew Schmidt.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**global which, which.max, and which.min**  
*Global Which Functions*

**Description**

These functions are global which, which.max and which.min applying on distributed data for all ranks.

**Usage**

```
comm.which(x, arr.ind = FALSE, useNames = TRUE,
           comm = .pbdr_env$SPMD.CT$comm)
comm.which.max(x, comm = .pbdr_env$SPMD.CT$comm)
comm.which.min(x, comm = .pbdr_env$SPMD.CT$comm)
```

**Arguments**

- x** a 'logical' vector or array as in **which()**, or an 'numeric' objects in **which.max()** and **which.min()**.  
**arr.ind** logical, as in **which()**.  
**useNames** logical, as in **which()**.  
**comm** a communicator number.

**Details**

These functions will apply **which()**, **which.max()** and **which.min()** locally, and apply **allgather()** to get all local results from other ranks.

## Value

The global values (which(), which.max(), or which.min()) are returned to all ranks. comm.which() returns with two columns, 'rank id' and 'index of TRUE'. comm.which.max() and comm.which.min() return with three values, 'the \_smallest\_ rank id', 'index of the \_first\_ maximum or minimum', and 'max/min value of x'.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[comm.read.table\(\)](#)

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
if(comm.size() != 2){
  comm.cat(\"2 processors are required.\n\", quiet = TRUE)
  finalize()
}

### Examples.
a <- 1:(comm.rank() + 1)

b <- comm.which(a == 2)
comm.print(b)
b <- comm.which.max(a)
comm.print(b)
b <- comm.which.min(a)
comm.print(b)

### Finish.
finalize()
"
# execmpi(spmd.code, nnranks = 2L)

## End(Not run)
```

## Description

These functions are global writing applying on distributed data for all ranks.

## Usage

```
comm.write(x, file = "data", ncolumns = if(is.character(x)) 1 else 5,
           append = FALSE, sep = " ", comm = .pbd_env$SPMD.CT$comm)
comm.write.table(x, file = "", append = FALSE, quote = TRUE, sep = " ",
                 eol = "\n", na = "NA", dec = ".", row.names = TRUE,
                 col.names = TRUE, qmethod = c("escape", "double"),
                 fileEncoding = "", comm = .pbd_env$SPMD.CT$comm)

comm.write.csv(..., comm = .pbd_env$SPMD.CT$comm)
comm.write.csv2(..., comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

x	as in <code>write()</code> or <code>write.table()</code> .
file	as in <code>write()</code> or <code>write.table()</code> .
ncolumns	as in <code>write*()</code> .
append	as in <code>write*()</code> .
sep	as in <code>write*()</code> .
quote	as in <code>write*()</code> .
eol	as in <code>write*()</code> .
na	as in <code>write*()</code> .
dec	as in <code>write*()</code> .
row.names	as in <code>write*()</code> .
col.names	as in <code>write*()</code> .
qmethod	as in <code>write*()</code> .
fileEncoding	as in <code>write*()</code> .
...	as in <code>write*()</code> .
comm	a communicator number.

## Details

These functions will apply `write*()` locally and sequentially from rank 0, 1, 2, ...

By default, rank 0 makes the file, and rest of ranks append the data.

**Value**

A file will be returned.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

`comm.load.balance()` and `comm.read.table()`

**Examples**

```
## Not run:  
### Save code in a file "demo.r" and run with 2 processors by  
### SHELL> mpiexec -np 2 Rscript demo.r  
  
spmd.code <- "  
### Initial.  
suppressMessages(library(pbdMPI, quietly = TRUE))  
init()  
if(comm.size() != 2){  
  comm.cat(\"2 processors are required.\n\", quiet = TRUE)  
  finalize()  
}  
  
### Examples.  
comm.write((1:5) + comm.rank(), file = \"test.txt\")  
  
### Finish.  
finalize()  
"  
# execmpi(spmd.code, nranks = 2L)  
  
## End(Not run)
```

---

**Description**

The functions call MPI info functions.

## Usage

```
info.create(info = .pbd_env$SPMD.CT$info)
info.set(info = .pbd_env$SPMD.CT$info, key, value)
info.free(info = .pbd_env$SPMD.CT$info)
info.c2f(info = .pbd_env$SPMD.CT$info)
```

## Arguments

info	a info number.
key	a character string to be set.
value	a character string to be set associate with key.

## Details

These functions are for internal functions. Potentially, they set info for initialization of master and workers.

## Value

An invisible state of MPI call is returned.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
info.create(0L)
info.set(0L, \"file\", \"appschema\")
info.free(0L)

### Finish.
finalize()
```

```

"
# execmpi(spmd.code, nnranks = 2L)

## End(Not run)

```

**irecv-method***A Rank Receives (Nonblocking) an Object from the Other Rank***Description**

This method lets a rank receive (nonblocking) an object from the other rank in the same communicator. The default return is the object sent from the other rank.

**Usage**

```
irecv(x.buffer = NULL, rank.source = .pbd_env$SPMD.CT$rank.source,
      tag = .pbd_env$SPMD.CT$tag, comm = .pbd_env$SPMD.CT$comm,
      request = .pbd_env$SPMD.CT$request,
      status = .pbd_env$SPMD.CT$status)
```

**Arguments**

x.buffer	a buffer to store x sent from the other rank.
rank.source	a source rank where x sent from
tag	a tag number.
comm	a communicator number.
request	a request number.
status	a status number.

**Details**

A corresponding send()/isend() should be evoked at the corresponding rank rank.source.

**Warning:** irecv() is not safe for R since R is not a thread safe package that a dynamic returning object requires certain blocking or barrier at some where. Current, the default method is equivalent to the default method of recv().

**Value**

An object is returned by default.

**Methods**

For calling spmd.irecv.\*():

```

signature(x = "ANY")
signature(x = "integer")
signature(x = "numeric")
signature(x = "raw")

```

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[recv\(\)](#), [send\(\)](#), [isend\(\)](#).

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
if(.comm.rank == 0){
  y <- send(matrix(x, nrow = 1))
} else if(.comm.rank == 1){
  y <- irecv()
}
comm.print(y, rank.print = 1)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

*is.comm.null*

*Check if a MPI\_COMM\_NULL*

## Description

The functions check MPI\_COMM\_NULL.

**Usage**

```
is.comm.null(comm = .pbdr_env$SPMD.CT$comm)
```

**Arguments**

comm a comm number.

**Details**

These functions are for internal uses.

**Value**

TRUE if input comm is MPI\_COMM\_NULL, otherwise FALSE.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**Examples**

```
## Not run:  
### Save code in a file "demo.r" and run with 2 processors by  
### SHELL> mpiexec -np 2 Rscript demo.r  
  
spmd.code <- "  
### Initial.  
suppressMessages(library(pbdMPI, quietly = TRUE))  
init()  
.comm.size <- comm.size()  
.comm.rank <- comm.rank()  
  
### Examples.  
is.comm.null(0L)  
is.comm.null(1L)  
  
### Finish.  
finalize()  
"  
# execmpi(spmd.code, nranks = 2L)  
  
## End(Not run)
```

---

**isend-method***A Rank Send (Nonblocking) an Object to the Other Rank*

---

**Description**

This method lets a rank send (nonblocking) a object to the other rank in the same communicator. The default return is NULL.

**Usage**

```
isend(x, rank.dest = .pbd_env$SPMD.CT$rank.dest,
      tag = .pbd_env$SPMD.CT$tag,
      comm = .pbd_env$SPMD.CT$comm,
      request = .pbd_env$SPMD.CT$request,
      check.type = .pbd_env$SPMD.CT$check.type)
```

**Arguments**

x	an object to be sent from a rank.
rank.dest	a rank of destination where x send to.
tag	a tag number.
comm	a communicator number.
request	a request number.
check.type	if checking data type first for handshaking.

**Details**

A corresponding `recv()` or `irecv()` should be evoked at the corresponding rank `rank.dest`. See details of `send()` for the arugments `check.type`.

**Value**

A NULL is returned by default.

**Methods**

For calling `spmd.isend.*()`:

```
signature(x = "ANY")
signature(x = "integer")
signature(x = "numeric")
signature(x = "raw")
```

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[send\(\)](#), [recv\(\)](#), [irecv\(\)](#).

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
if(.comm.rank == 0){
  y <- isend(matrix(x, nrow = 1))
} else if(.comm.rank == 1){
  y <- recv()
}
comm.print(y, rank.print = 1)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

**Description**

The function set/get a point address in R where the point point to a structure containing MPI arrays.

## Usage

```
arrange.mpi.apts()
```

## Details

Since Rmpi/pbdMPI use pre-allocate memory to store comm, status, datatype, info, request, this function provides a variable in R to let different APIs share the same memory address.

If the package loads first, then this sets ‘`.__MPI_APTS__`’ in the `.GlobalEnv` of R. If the package does not load before other MPI APIs, then this points an structure point to the external memory according to ‘`.__MPI_APTS__`’, i.e. pre-set by other MPI APIs.

`pbdMPI/R/arrange.mpi.apts` provides the R code, and `pbdMPI/src/pkg_*.*` provides the details of this call.

## Value

‘`.__MPI_APTS__`’ is set in the `.GlobalEnv` of R.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:  
### See source code for the details.  
  
## End(Not run)
```

## Description

These functions are designed to get or print MPI\_COMM pointer and its address when the SPMD code in R be a foreign application of other applications.

## Usage

```
get.mpi.comm.ptr(comm = .pbd_env$SPMD.CT$comm, show.msg = FALSE)  
addr.mpi.comm.ptr(comm.ptr)
```

### Arguments

comm	a communicator number.
comm.ptr	a communicator pointer.
show.msg	if showing message for debug only.

### Details

`get.mpi.comm.ptr()` returns an R external pointer that points to the address of the comm.  
`addr.mpi.comm.ptr()` takes the R external points, and prints the address of the comm. This function is mainly for debugging.

### Value

`get.mpi.comm.ptr()` returns an R external pointer.  
`addr.mpi.comm.ptr()` prints the comm pointer address and the address of MPI\_COMM\_WORLD.

### Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

### References

Programming with Big Data in R Website: <https://pbdr.org/>

### Examples

```
### Save code in a file "demo.r" and run with 2processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

ptr1 <- get.mpi.comm.ptr(1, show.msg = TRUE)
addr.mpi.comm.ptr(ptr1)

comm.split(color = as.integer(comm.rank()/2), key = comm.rank())

ptr1.new <- get.mpi.comm.ptr(1, show.msg = TRUE)
addr.mpi.comm.ptr(ptr1.new)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code = spmd.code, nranks = 2L)
```

---

probe

*Probe Functions*

---

## Description

The functions call MPI probe functions.

## Usage

```
probe(rank.source = .pbd_env$SPMD.CT$rank.source,
      tag = .pbd_env$SPMD.CT$tag, comm = .pbd_env$SPMD.CT$comm,
      status = .pbd_env$SPMD.CT$status)
iprobe(rank.source = .pbd_env$SPMD.CT$rank.source,
       tag = .pbd_env$SPMD.CT$tag, comm = .pbd_env$SPMD.CT$comm,
       status = .pbd_env$SPMD.CT$status)
```

## Arguments

rank.source	a source rank where an object sent from.
tag	a tag number.
comm	a communicator number.
status	a status number.

## Details

These functions are for internal functions. Potentially, they set/get probe for receiving data.

## Value

An invisible state of MPI call is returned.

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## Examples

```
## Not run:
### See source code of spmd.recv.default() for an example.

## End(Not run)
```

---

recv-method*A Rank Receives (Blocking) an Object from the Other Rank*

---

**Description**

This method lets a rank receive (blocking) an object from the other rank in the same communicator. The default return is the object sent from the other rank.

**Usage**

```
recv(x.buffer = NULL, rank.source = .pbd_env$SPMD.CT$rank.source,
     tag = .pbd_env$SPMD.CT$tag, comm = .pbd_env$SPMD.CT$comm,
     status = .pbd_env$SPMD.CT$status,
     check.type = .pbd_env$SPMD.CT$check.type)
```

**Arguments**

x.buffer	a buffer to store x sent from the other rank.
rank.source	a source rank where x sent from
tag	a tag number.
comm	a communicator number.
status	a status number.
check.type	if checking data type first for handshaking.

**Details**

A corresponding send() should be evoked at the corresponding rank rank.source.

These are high level S4 methods. By default, check.type is TRUE and an additional send()/recv() will make a handshaking call first, then deliver the data next. i.e. an integer vector of length two (type and length) will be delivered first between send() and recv() to ensure a buffer (of right type and right size/length) is properly allocated at the rank.dest side.

Currently, four data types are considered: integer, double, raw/byte, and default/raw.object. The default method will make a serialize() call first to convert the general R object into a raw vector before sending it away. After the raw vector is received at the rank.dest side, the vector will be unserialize() back to the R object format.

check.type set as FALSE will stop the additional handshaking call, but the buffer should be prepared carefully by the user self. This is typically for the advanced users and more specifically calls are needed. i.e. calling those spmd.send.integer with spmd.recv.integer correspondingly.

check.type also needs to be set as FALSE for more efficient calls such as isend()/recv() or send()/irecv(). Currently, no check types are implemented in those mixed calls.

**Value**

An object is returned by default and the buffer will be overwritten implicitly.

## Methods

For calling `spmd.recv.*()`:

```
signature(x = "ANY")
signature(x = "integer")
signature(x = "numeric")
signature(x = "raw")
```

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

`irecv()`, `send()`, `isend()`.

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
if(.comm.rank == 0){
  y <- send(matrix(x, nrow = 1))
} else if(.comm.rank == 1){
  y <- recv()
}
comm.print(y, rank.print = 1)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)
```

---

reduce-method*A Rank Receive a Reduction of Objects from Every Rank*

---

## Description

This method lets a rank receive a reduction of objects from every rank in the same communicator based on a given operation. The default return is an object as the input.

## Usage

```
reduce(x, x.buffer = NULL, op = .pbd_env$SPMD.CT$op,
       rank.dest = .pbd_env$SPMD.CT$rank.source,
       comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

x	an object to be gathered from all ranks.
x.buffer	a buffer to hold the return object which probably has x with the same type of x.
op	a reduction operation applied on combine all x.
rank.dest	a rank of destination where all x reduce to.
comm	a communicator number.

## Details

By default, the object is reduced to .pbd\_env\$SPMD.CT\$rank.source, i.e. *rank 0L*.

All x on all ranks are likely presumed to have the same size and type.

x.buffer can be NULL or unspecified. If specified, the type should be either integer or double specified correctly according to the type of x.

## Value

The reduced object of the same type as x is returned by default.

## Methods

For calling spmd.reduce.\*():

```
signature(x = "ANY", x.buffer = "missing")
signature(x = "integer", x.buffer = "integer")
signature(x = "numeric", x.buffer = "numeric")
signature(x = "logical", x.buffer = "logical")
signature(x = "float32", x.buffer = "float32")
```

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[allgather\(\)](#), [gather\(\)](#), [reduce\(\)](#).

**Examples**

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
y <- reduce(matrix(x, nrow = 1), op = \\"sum\\")
comm.print(y)

y <- reduce(x, double(N), op = \\"prod\\")
comm.print(y)

x <- as.logical(round(runif(N)))
y <- reduce(x, logical(N), op = \\"land\\")
comm.print(y)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code = spmd.code, nranks = 2L)
```

## Description

This method lets a rank scatter objects to every rank in the same communicator. The default input is a list of length equal to ‘comm size’ and the default return is an element of the list.

## Usage

```
scatter(x, x.buffer = NULL, x.count = NULL, displs = NULL,
       rank.source = .pbd_env$SPMD.CT$rank.source,
       comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

x	an object of length ‘comm size’ to be scattered to all ranks.
x.buffer	a buffer to hold the return object which probably has ‘size of element of x’ with the same type of the element of x.
x.count	a vector of length ‘comm size’ containing all object lengths.
displs	c(0L, cumsum(x.count)) by default.
rank.source	a rank of source where elements of x scatter from.
comm	a communicator number.

## Details

All elements of x are likely presumed to have the same size and type.

x.buffer, x.count, and displs can be NULL or unspecified. If specified, the type should be one of integer, double, or raw specified correctly according to the type of x.

If x.count is specified, then the spmd.scatterv.\*() is called.

## Value

An element of x is returned according to the rank id.

## Methods

For calling spmd.scatter.\*():

```
signature(x = "ANY", x.buffer = "missing", x.count = "missing")
signature(x = "integer", x.buffer = "integer", x.count = "missing")
signature(x = "numeric", x.buffer = "numeric", x.count = "missing")
signature(x = "raw", x.buffer = "raw", x.count = "missing")
```

For calling spmd.scatterv.\*():

```
signature(x = "ANY", x.buffer = "missing", x.count = "integer")
signature(x = "ANY", x.buffer = "ANY", x.count = "integer")
signature(x = "integer", x.buffer = "integer", x.count = "integer")
signature(x = "numeric", x.buffer = "numeric", x.count = "integer")
signature(x = "raw", x.buffer = "raw", x.count = "integer")
```

## Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[bcast\(\)](#).

## Examples

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- split(1:(N * .comm.size), rep(1:.comm.size, N))
y <- scatter(lapply(x, matrix, nrow = 1))
comm.print(y)
y <- scatter(x, double(N))
comm.print(y)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)
```

## Description

These functions set/end/reset seeds to all ranks. By default, these functions are wrappers of **rlecuyer** which implements the random number generator with multiple independent streams developed by L'Ecuyer et al (2002).

## Usage

```
comm.set.seed(seed, diff = FALSE, state = NULL,
             comm = .pbd_env$SPMD.CT$comm)
comm.seed.state(comm = .pbd_env$SPMD.CT$comm)
comm.end.seed(comm = .pbd_env$SPMD.CT$comm)
comm.reset.seed(comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

seed	one integer or six integers as in <b>rlecuyer</b> .
diff	if all ranks use the same stream. (default = FALSE)
state	a new state to overwrite seed.
comm	a communicator number.

## Details

`comm.set.seed()` sets the given seed to all ranks. If `diff = FALSE`, then all ranks generate one stream and use that stream. Otherwise, all ranks generate `COMM.SIZE` streams and use the stream named by `COMM.RANK`.

Also, `comm.set.seed()` can assign to arbitrarily `state` obtained from `comm.seed.state()`.

`comm.seed.state()` obtains current state of seed which ends the stream first (update state), gets the state, and continues the stream (pretend as nothing happens).

`comm.end.seed()` ends and deletes seed from all ranks.

`comm.reset.seed()` resets seed to initial start steps which end the current seed and reset everything back to the start stream. Use this function with caution.

## Value

Several hidden objects are set in the `.GlobalEnv`, see **rlecuyer** package for details.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Pierre L'Ecuyer, Simard, R., Chen, E.J., and Kelton, W.D. (2002) An Object-Oriented Random-Number Package with Many Long Streams and Substreams. *Operations Research*, 50(6), 1073-1075.

<https://www.iro.umontreal.ca/~lecuyer/myftp/papers/streams00.pdf>

Sevcikova, H. and Rossini, T. (2012) rlecuyer: R interface to RNG with multiple streams. R Package, URL <https://cran.r-project.org/package=rlecuyer>

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

.lec.SetPackageSeed(), .lec.CreateStream(), .lec.CurrentStream(), .lec.CurrentStreamEnd(), .lec.DeleteStream(), .lec.SetSeed(), and .lec.GetState().

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()

### Examples.
comm.set.seed(123456)
comm.print(runif(5), all.rank = TRUE)
comm.reset.seed()
comm.print(runif(5), all.rank = TRUE)
comm.end.seed()

### Obtain the seed state.
comm.set.seed(123456, diff = TRUE)
comm.print(runif(5), all.rank = TRUE)
saved.seed <- comm.seed.state()    ### save the state.
comm.print(runif(5), all.rank = TRUE)
comm.end.seed()

### Start from a saved state.
comm.set.seed(123456, state = saved.seed) ### rewind to the state.
comm.print(runif(5), all.rank = TRUE)
comm.end.seed()

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

## Description

This method lets a rank send (blocking) an object to the other rank in the same communicator. The default return is NULL.

## Usage

```
send(x, rank.dest = .pbd_env$SPMD.CT$rank.dest,
     tag = .pbd_env$SPMD.CT$tag,
     comm = .pbd_env$SPMD.CT$comm,
     check.type = .pbd_env$SPMD.CT$check.type)
```

## Arguments

x	an object to be sent from a rank.
rank.dest	a rank of destination where x send to.
tag	a tag number.
comm	a communicator number.
check.type	if checking data type first for handshaking.

## Details

A corresponding `recv()` should be evoked at the corresponding rank `rank.dest`.

These are high level S4 methods. By default, `check.type` is TRUE and an additional `send()`/`recv()` will make a handshaking call first, then deliver the data next. i.e. an integer vector of length two (type and length) will be delivered first between `send()` and `recv()` to ensure a buffer (of right type and right size/length) is properly allocated at the `rank.dest` side.

Currently, four data types are considered: `integer`, `double`, `raw/byte`, and `default/raw.object`. The default method will make a `serialize()` call first to convert the general R object into a raw vector before sending it away. After the raw vector is received at the `rank.dest` side, the vector will be  `unserialize()` back to the R object format.

`check.type` set as FALSE will stop the additional handshaking call, but the buffer should be prepared carefully by the user self. This is typically for the advanced users and more specifically calls are needed. i.e. calling those `spmd.send.integer` with `spmd.recv.integer` correspondingly.

`check.type` also needs to be set as FALSE for more efficient calls such as `isend()`/`recv()` or `send()`/`irecv()`. Currently, no check types are implemented in those mixed calls.

## Value

A NULL is returned by default.

## Methods

For calling `spmd.send.*()`:

```
signature(x = "ANY")
signature(x = "integer")
signature(x = "numeric")
signature(x = "raw")
```

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[isend\(\)](#), [recv\(\)](#), [irecv\(\)](#).

**Examples**

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
if(.comm.rank == 0){
  y <- send(matrix(x, nrow = 1))
} else if(.comm.rank == 1){
  y <- recv()
}
comm.print(y, rank.print = 1)

### Finish.
finalize()
"
pbdMPI::execmpi(spmd.code, nranks = 2L)
```

**Description**

This method lets a rank send an object to the other rank and receive an object from another rank in the same communicator. The default return is x.

## Usage

```
sendrecv(x, x.buffer = NULL,
         rank.dest = (comm.rank(.pbd_env$SPMD.CT$comm) + 1) %%  

                      comm.size(.pbd_env$SPMD.CT$comm),
         send.tag = .pbd_env$SPMD.CT$tag,
         rank.source = (comm.rank(.pbd_env$SPMD.CT$comm) - 1) %%  

                      comm.size(.pbd_env$SPMD.CT$comm),
         recv.tag = .pbd_env$SPMD.CT$tag,
         comm = .pbd_env$SPMD.CT$comm, status = .pbd_env$SPMD.CT$status)
```

## Arguments

x	an object to be sent from a rank.
x.buffer	a buffer to store x sent from the other rank.
rank.dest	a rank of destination where x send to.
send.tag	a send tag number.
rank.source	a source rank where x sent from.
recv.tag	a receive tag number.
comm	a communicator number.
status	a status number.

## Details

A corresponding `sendrecv()` should be evoked at the corresponding ranks `rank.dest` and `rank.source`. `rank.dest` and `rank.source` can be `as.integer(NULL)` to create a silent `sendrecv` operation which is more efficient than setting `rank.dest` and `rank.source` to be equal.

## Value

A x is returned by default.

## Methods

For calling `spmd.sendrecv.*()`:

```
signature(x = "ANY", x.buffer = "ANY")
signature(x = "integer", x.buffer = "integer")
signature(x = "numeric", x.buffer = "numeric")
signature(x = "raw", x.buffer = "raw")
```

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[sendrecv.replace\(\)](#).

## Examples

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.size
y <- sendrecv(matrix(x, nrow = 1))
comm.print(y, rank.print = 1)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

## sendrecv.replace-method

*Send and Receive an Object to and from Other Ranks*

## Description

This method lets a rank send an object to the other rank and receive an object from another rank in the same communicator. The default return is x.

## Usage

```
sendrecv.replace(x,
rank.dest = (comm.rank(.pbdd_env$SPMD.CT$comm) + 1) %%
               comm.size(.pbdd_env$SPMD.CT$comm),
send.tag = .pbdd_env$SPMD.CT$tag,
rank.source = (comm.rank(.pbdd_env$SPMD.CT$comm) - 1) %%
```

```

comm.size(.pbd_env$SPMD.CT$comm),
recv.tag = .pbd_env$SPMD.CT$tag,
comm = .pbd_env$SPMD.CT$comm, status = .pbd_env$SPMD.CT$status)

```

## Arguments

x	an object to be sent from a rank.
rank.dest	a rank of destination where x send to.
send.tag	a send tag number.
rank.source	a source rank where x sent from.
recv.tag	a receive tag number.
comm	a communicator number.
status	a status number.

## Details

A corresponding `sendrecv.replace()` should be evoked at the corresponding ranks `rank.dest` and `rank.source`.

`rank.dest` and `rank.source` can be `as.integer(NULL)` to create a silent `sendrecv` operation which is more efficient than setting `rank.dest` and `rank.source` to be equal.

**Warning:** `sendrecv.replace()` is not safe for R since R is not a thread safe package that a dynamic returning object requires certain blocking or barrier at some where. The replaced object or memory address ‘MUST’ return correctly. This is almost equivalent to `sendrecv()`.

## Value

A x is returned by default.

## Methods

For calling `spmd.sendrecv.replace.*()`:

```

signature(x = "ANY")
signature(x = "integer")
signature(x = "numeric")
signature(x = "raw")

```

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

[sendrecv\(\)](#).

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.size
x <- sendrecv.replace(matrix(x, nrow = 1))
comm.print(x, rank.print = 1)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

**Set global pbd options**

*Set Global pbdR Options*

**Description**

This is an advanced function to set pbdR options.

**Usage**

```
pbd_opt(..., bytext = "", envir = .GlobalEnv)
```

**Arguments**

- ... in argument format option = value to set .pbd\_env\$option <- value inside the envir.
- bytext in text format "option = value" to set .pbd\_env\$option <- value inside the envir.
- envir by default the global environment is used.

## Details

. . . allows multiple options in `envir$.pbd_env`, but only in a simple way.

`bytext` allows to assign options by text in `envir$.pbd_env`, but can assign advanced objects. For example, "`option$suboption <- value`" will set `envir$.pbd_env$option$suboption <- value`.

## Value

No value is returned.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)> and Drew Schmidt.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

`.pbd_env`, `SPMD.CT()`, `SPMD.OP()`, `SPMD.IO()`, `SPMD.TP()`, and `.mpiopt_init()`.

## Examples

```
## Not run:  
### Save code in a file "demo.r" and run with 4 processors by  
### SHELL> mpiexec -np 4 Rscript demo.r  
  
### Initial.  
suppressMessages(library(pbdMPI, quietly = TRUE))  
init()  
  
### Examples.  
ls(.pbd_env)  
pbd_opt(ICTXT = c(2, 2))  
pbd_opt(bytext = "grid.new <- list(); grid.new$ICTXT <- c(4, 4)")  
pbd_opt(BLDIM = c(16, 16), bytext = "grid.new$BLDIM = c(8, 8)")  
ls(.pbd_env)  
.pbd_env$ICTXT  
.pbd_env$BLDIM  
.pbd_env$grid.new  
  
### Finish.  
finalize()  
  
## End(Not run)
```

---

**sourcetag***Functions to Obtain source and tag*

---

**Description**

The functions extract MPI\_ANY\_SOURCE, MPI\_ANY\_TAG, MPI\_status.source and MPI\_status.tag.

**Usage**

```
anysource()
anytag()
get.sourcetag(status = .pbd_env$SPMD.CT$status)
```

**Arguments**

status	a status number.
--------	------------------

**Details**

These functions are for internal uses.

**Value**

Corresponding status will be returned.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
.comm.size <- comm.size()
.comm.rank <- comm.rank()
if(.comm.size < 2)
  comm.stop(\"At least two processors are required.\")"
```

```

### Examples.
if(.comm.rank != 0){
  send(as.integer(.comm.rank * 10), rank.dest = 0L,
       tag = as.integer(.comm.rank + 10))
}
if(.comm.rank == 0){
  for(i in 1:(.comm.size - 1)){
    ret <- recv(x.buffer = integer(1),
                 rank.source = anysource(), tag = anytag())
    sourcetag <- get.sourcetag()
    print(c(sourcetag, ret))
  }
}

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)

```

**SPMD Control***Sets of controls in pbdMPI.***Description**

These sets of controls are used to provide default values in this package.

**Format**

Objects contain several parameters for communicators and methods.

**Details**

The elements of `.pbd_env$SPMD.CT` are default values for controls including

Elements	Default	Usage
comm	0L	a communicator index
intercomm	2L	an inter communicator index
info	0L	an info index
newcomm	1L	a new communicator index
op	"sum"	an operation
port.name	"spmdport"	an operation
print.all.rank	FALSE	if all ranks print message
print.quiet	FALSE	if print/cat rank information
rank.root	0L	a rank of root
rank.source	0L	a rank of source
rank.dest	1L	a rank of destination
request	0L	a request index

<code>serv.name</code>	"spmdserv"	a service name
<code>status</code>	0L	a status index
<code>tag</code>	0L	a tag number
<code>unlist</code>	FALSE	if unlist returning
<code>divide.method</code>	"block"	a way to divide jobs or data
<code>mpi.finalize</code>	TRUE	if shutdown MPI
<code>quit</code>	TRUE	if quit when errors occur
<code>msg.flush</code>	TRUE	if flush message for comm.cat/comm.print
<code>msg.barrier</code>	TRUE	if barrier message for comm.cat/comm.print
<code>Rprof.all.rank</code>	FALSE	if call Rprof on all ranks
<code>lazy.check</code>	TRUEE	if use lazy check on all ranks

The elements of `.pbd_env$SPMD.OP` list the implemented operations for `reduce()` and `allreduce()`. Currently, four operations are implemented "sum", "prod", "max", and "min".

The elements of `.SPMD.IO` are default values for input and output including

Elements	Default	Usage
<code>max.read.size</code>	5.2e6	max of reading size (5 MB)
<code>max.test.lines</code>	500	max of testing lines
<code>read.method</code>	"gbd"	default reading method
<code>balance.method</code>	"block"	default load balance method

where `balance.method` is only used for "gbd" reading method when `nrows = -1` and `skip = 0` are set.

The elements of `.pbd_env$SPMD.TP` are default values mainly for task pull including

Elements	Default	Usage
<code>bcast</code>	FALSE	if bcase() objects to all ranks
<code>barrier</code>	TRUE	if call barrier() for all ranks
<code>try</code>	TRUE	if use try() in works
<code>try.silent</code>	FALSE	if silent the try() message

See `task.pull()` for details.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

---

**SPMD Control Functions**

*Sets of controls in pbdMPI.*

---

### Description

These sets of controls are used to provide default values in this package. The values are not supposed to be changed in general.

### Author(s)

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

### References

Programming with Big Data in R Website: <https://pbdr.org/>

### See Also

.pbd\_env.

---

**Task Pull**

*Functions for Task Pull Parallelism*

---

### Description

These functions are designed in SPMD but assuming rank 0 is a master and rests are workers.

### Usage

```
task.pull(jids, FUN, ..., rank.master = .pbd_env$SPMD.CT$rank.root,
          comm = .pbd_env$SPMD.CT$comm, bcast = .pbd_env$SPMD.TP$bcast,
          barrier = .pbd_env$SPMD.TP$barrier,
          try = .pbd_env$SPMD.TP$try,
          try.silent = .pbd_env$SPMD.TP$try.silent)

task.pull.workers(FUN = function(jid, ...){ return(jid) }, ...,
                  rank.master = .pbd_env$SPMD.CT$rank.root,
                  comm = .pbd_env$SPMD.CT$comm,
                  try = .pbd_env$SPMD.TP$try,
                  try.silent = .pbd_env$SPMD.TP$try.silent)
task.pull.master(jids, rank.master = .pbd_env$SPMD.CT$rank.root,
                 comm = .pbd_env$SPMD.CT$comm)
```

## Arguments

jids	all job ids (a vector of positive integers).
FUN	a function to be evaluated in workers.
...	extra variables for FUN.
rank.master	a rank of master where jid sent from.
comm	a communicator number.
bcast	if bcast to all ranks.
barrier	if barrier for all ranks.
try	if use <code>try()</code> to avoid breaks. CAUTION: <code>try = FALSE</code> is not safe and can stop all MPI/R jobs.
try.silent	if turn off the error message from <code>try()</code> .

## Details

All of these functions are for SPMD, NOT for master/workers.

FUN is a user defined function which has `jid` as the first argument and other variables are given in ...

The `jids` will be asked by workers when jobs are available and workers are no job in hand.

## Value

A list with length `comm.size() - 1` will be returned for mater, but `NULL` for workers. Each element of the list contains returns `ret` of the FUN call.

## Author(s)

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

## References

Programming with Big Data in R Website: <https://pbdr.org/>

## See Also

[get.jid\(\)](#).

## Examples

```
## Not run:
### Under command mode, run the demo with 2 processors by
### (Use Rscript.exe for windows system)
# mpiexec -np 2 Rscript -e "demo(task_pull,'pbMPI',ask=F,echo=F)"
### Or
# execmpi("demo(task_pull,'pbMPI',ask=F,echo=F)", nranks = 2L)

## End(Not run)
```

---

Utility execmpi	<i>Execute MPI code in system</i>
-----------------	-----------------------------------

---

## Description

This function basically saves code in a spmd.file and executes MPI via R's system call e.g. `system("mpexec -np 1 Rscript spmd.file")`.

## Usage

```
execmpi(spmd.code = NULL, spmd.file = NULL,
        mpicmd = NULL, nranks = 1L, rscmd = NULL, verbose = TRUE,
        disable.current.mpi = TRUE)
runmpi(spmd.code = NULL, spmd.file = NULL,
       mpicmd = NULL, nranks = 1L, rscmd = NULL, verbose = TRUE,
       disable.current.mpi = TRUE)
```

## Arguments

<code>spmd.code</code>	SPMD code to be run via mpicmd and Rscript.
<code>spmd.file</code>	a file contains SPMD code to be run via mpicmd and Rscript.
<code>mpicmd</code>	MPI executable command. If NULL, system default will be searched.
<code>nranks</code>	number of processes to run the SPMD code envoked by mpicmd.
<code>rscmd</code>	Rscript executable command. If NULL, system default will be searched.
<code>verbose</code>	print SPMD code outputs and MPI messages.
<code>disable.current.mpi</code>	force to finalize the current MPI comm if any, for unix-alike system only.

## Details

When the `spmd.code` is NULL: The code should be already saved in the file named `spmd.file` for using.

When the `spmd.code` is not NULL: The `spmd.code` will be dumped to a temp file (`spmd.file`) via the call `writeLines(spmd.code, conn)` where `conn <- file(spmd.file, open = "wt")`. The file will be closed after the dumping.

When `spmd.file` is ready (either dumped from `spmd.code` or provided by the user), the steps below will be followed: If `spmd.file = NULL`, then a temporary file will be generated and used to dump `spmd.code`.

For Unix-alike systems, the command `cmd <- paste(mpicmd, "-np", nranks, rscmd, spmd.file, ">", log.file, "2>&1 & echo \"PID=$!\" &")` is executed via `system(cmd, intern = TRUE, wait = FALSE, ignore.stdout = TRUE, ignore.stderr = TRUE)`. The `log.file` is a temporary file to save the outputs from the `spmd.code`. The results saved to the `log.file` will be read back in and `cat` and return to R.

For Windows, the `cmd` will be `paste(mpicmd, "-np", nranks, rscmd, spmd.file)` and is executed via `system(cmd, intern = TRUE, wait = FALSE, ignore.stdout = TRUE, ignore.stderr = TRUE)`.

**Value**

Basically, only the PID of the MPI job (in background) will be returned in Linux-alike systems. For Windows, the MPI job is always wait until it is complete.

**Note**

For Unix-alike systems, in new R and MPI, the `pbdMPI::execmpi(...)` may carry the current MPI `comm` into `system(cmd, ...)` calls. Because the `comm` has been established-loaded by the `init()` call because of `::`, the `mpiexec` inside the `system(cmd, ...)` calls will be confused with the exist `comm`.

Consider that `pbdMPI::execmpi(...)` is typically called in interactive mode (or actually only done for CRAN check in most case), an argument `disable.current.mpi = TRUE` is added/needed to finalize the existing `comm` first before `system(cmd, ...)` be executed.

This function is NOT recommended for running SPMD programs. The recommended way is to run under shell command.

**Author(s)**

Wei-Chen Chen <[wccsnow@gmail.com](mailto:wccsnow@gmail.com)> and Drew Schmidt.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**See Also**

`pbdCS::pbdRscript()`.

**Examples**

```
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.file <- tempfile()
cat("
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
allreduce(2)
finalize()
", file = spmd.file)
pbdMPI::execmpi(spmd.file = spmd.file, nranks = 2L)
```

---

**wait***Wait Functions*

---

**Description**

The functions call MPI wait functions.

**Usage**

```
wait(request = .pbd_env$SPMD.CT$request,
      status = .pbd_env$SPMD.CT$status)
waitany(count, status = .pbd_env$SPMD.CT$status)
waiitsome(count)
waitall(count)
```

**Arguments**

request	a request number.
status	a status number.
count	a count number.

**Details**

These functions are for internal uses. Potentially, they wait after some nonblocking MPI calls.

**Value**

An invisible state of MPI call is returned.

**Author(s)**

Wei-Chen Chen <wccsnow@gmail.com>, George Ostrouchov, Drew Schmidt, Pragneshkumar Patel, and Hao Yu.

**References**

Programming with Big Data in R Website: <https://pbdr.org/>

**Examples**

```
## Not run:
### Save code in a file "demo.r" and run with 2 processors by
### SHELL> mpiexec -np 2 Rscript demo.r

spmd.code <- "
### Initial.
suppressMessages(library(pbdMPI, quietly = TRUE))
init()
```

```
.comm.size <- comm.size()
.comm.rank <- comm.rank()

### Examples.
N <- 5
x <- (1:N) + N * .comm.rank
if(.comm.rank == 0){
  isend(list(x))
}
if(.comm.rank == 1){
  y <- irecv(list(x))
}
wait()
comm.print(y, rank.print = 1L)

### Finish.
finalize()
"
# execmpi(spmd.code, nranks = 2L)

## End(Not run)
```

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