Guide to the $\mathbf{pbdPROF}$ Package

Wei-Chen Chen 1, Drew Schmidt 2, Gaurav Sehrawat 3, Pragneshkumar Patel 2, and George Ostrouchov 1,2

¹Computer Science and Mathematics Division, Oak Ridge National Laboratory, Oak Ridge, TN, USA

²Remote Data Analysis and Visualization Center University of Tennessee, Knoxville, TN, USA

³Jaypee Institute of Information Technology Uttar Pradesh, India

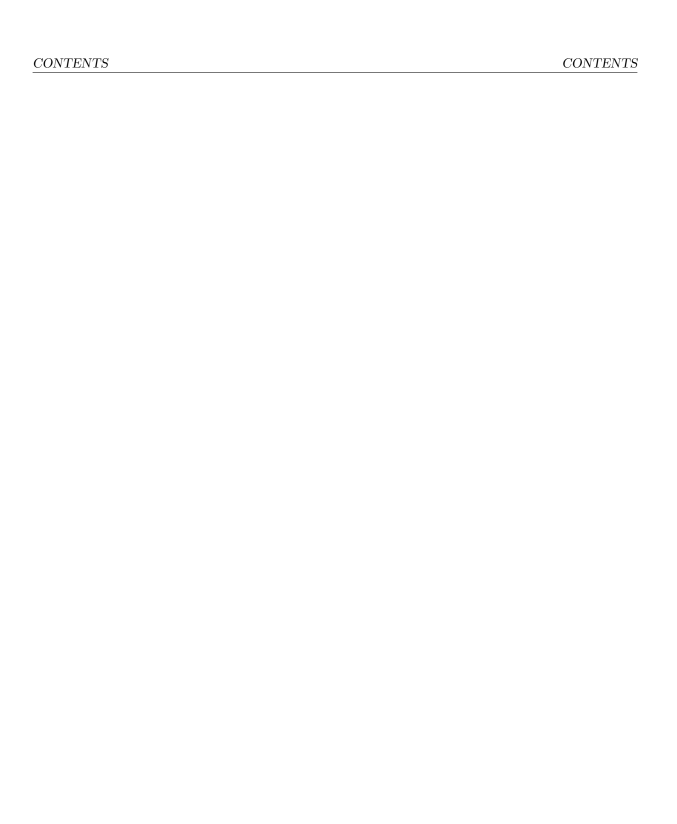
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Warning: The findings and conclusions in this article have not been formally disseminated by the U.S. Department of Energy and should not be construed to represent any determination or policy of University, Agency, and National Laboratory.

This document is written to explain the main functions of **pbdPROF** (Chen *et al.*, 2013), version 0.1-0. Every effort will be made to ensure future versions are consistent with these instructions, but features in later versions may not be explained in this document.

Information about the functionality of this package, and any changes in future versions can be found on website: "Programming with Big Data in R" at http://r-pbd.org/.

1 Introduction

The goal of **pbdPROF** is to utilize external MPI profiling libraries, such as **fpmpi** (Gropp, 2000), **mpiP** (Vetter and McCracken, 2001), or **TAU** (Shende and Malony, 2006), to profile parallel R code and understand hidden MPI communications between processors. The number of communications, sizes of messages, times, and types of functions calls all affect program performance, and so having these measurements can greatly aid in debugging and algorithm design. These MPI profiling libraries are able to hijack calls to MPI functions and then capture the profiling information (as described above), all without disturbing the execution of the original program.

The current main features of **pbdPROF** include:

- 1. providing linking information to pbdR (Ostrouchov et al., 2012) and other MPI-using R packages
- 2. output profiling information associated with MPI calls,
- 3. parsing and summarizing profiling information, and
- 4. support several MPI profiling libraries.

2 Installation

2.1 System Requirements

The **pbdPROF** package requires an MPI installation, such as OpenMPI or MPICH2. Additionally, the package is basically useless without some kind of MPI-using R package, such as **pbdMPI** (Chen *et al.*, 2012a) or **Rmpi** (Yu, 2002). For information regarding how to install MPI or **pbdMPI**, please see the **pbdMPI** vignette (Chen *et al.*, 2012b) or the pbdR website http://r-pbd.org/install.

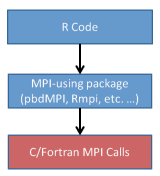
2.2 The Big Picture

Before pressing on, let us stop to take a moment and understand the "big picture" here. The following sections will contain *more than sufficient* detail, to the point where it would be easy to lose sight of the proverbial forest for the trees.

For the remainder of this document, we will be providing information for two fairly distinct groups of people: R-level MPI package developers, and C/Fortran-level MPI package developers. If you are in the former category, then the use of this package is a bit simpler for you. All you need to do is get **pbdPROF**

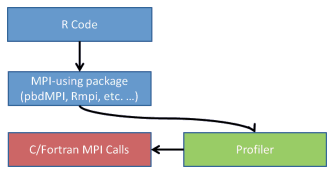
installed and reinstall your MPI-using package of choice (**pbdMPI**, **Rmpi**, etc. ...). Each package that directly uses MPI (packages produced by developers in the latter category) will have to explicitly support **pbdPROF** (or the reader will have to get his/her hands dirty in another developer's makefiles — an unpleasant business). It is worth nothing here that there are instructions in this document for how a developer of the second kind could explicitly add **pbdPROF** support to his/her package.

So why the need to reinstall things? It boils down to how the profilers actually work. Under normal circumstances, a user writes some R code from an MPI-using package (e.g., allreduce(x) from pbdMPI, mpi.allreduce(x, type=2) from Rmpi, etc. ...). This then makes a call to some C or Fortran code



Without a Profiler

which directly interfaces with MPI. You can see this pictures in Figure 1. When you use a profiler, you instead hijack the calls to MPI from the C/Fortran code so that some metadata can be stored about MPI usage. This process is represented in Figure 2. Hopefully it should be clear what, and when, something



With the Profiler

should be reinstalled. For the sake of completion, we summarize the possibilities below:

To enable MPI profiling:

- 1. install **pbdPROF**
- 2. reinstall an MPI-using package and link it with **pbdPROF**
- 3. write and execute your MPI-using R code as normal
- 4. use the pbdPROF utilities read.prof(), plot(), etc. for interpreting profiling results

To disable MPI profiling:

1. reinstall any MPI-using package that was linked it with **pbdPROF**, and this time *do not* link with **pbdPROF**

2.3 Choice of Profiler

The pbdPROF package currently uses the fpmpi library by default. More explicitly, a source copy of fpmpi is located at pbdPROF/src/fpmpi of the pbdPROF source. If this profiler is used, static library will be built and placed in pbdPROF/lib/libfpmpi.a of the pbdPROF install directory. However, external profiling libraries such as mpiP, TAU, or even fpmpi can be also linked with pbdPROF by passing a suitable --configure-args argument during an installation via R CMD INSTALL. We will explain this procedure in depth in Section 2.4 using an external fpmpi and mpiP as an example, TAU will be added in next release. for future Sections.

While it is possible to link with other profiling libraries, at the time of writing (for version 0.2-0), we currently support **fpmpi** and **mpiP**. We anticipate full of **TAU** for the next version of this package.

Regardless of whether **fpmpi**, **mpiP**, or **TAU** is used, we strongly recommend adding **CPPFLAGS="-fPIC"** at the **configure** step.

2.4 fpmpi

We can install **pbdPROF** using the internal **fpmpi** library via

Shell Command

```
R CMD INSTALL pbdPROF_0.1-0.tar.gz
```

By default, this compiles pbdPROF/src/fpmpi/* of the pbdPROF source, generates a static library libfpmpi.a, and installs the library to pbdPROF/lib/ of the pbdPROF install. No shared library is generated or needed, so the directory pbdPROF/libs/ is empty, i.e., there is no need to build pbdPROF.so. The linking argument is saved in Makeconf and installed to pbdPROF/etc/ for later use by other packages, such as pbdMPI or Rmpi.

However, if we choose, we can link with an external **fpmpi** library, via

```
Shell Command
```

```
R CMD INSTALL pbdPROF_0.1-0.tar.gz \
--configure-args="--with-fpmpi='/path_to_fpmpi/lib/libfpmpi.a'"
```

or

Shell Command

```
R CMD INSTALL pbdPR0F_0.1-0.tar.gz \
--configure-args="--with-fpmpi='-L/path_to_fpmpi/lib -lfpmpi'"
```

Or the conventional method in R console

Shell Command

Or

Since fpmpi only builds a static library libfpmpi.a, there is no difference between these two installations of pbdPROF. This only provides the linking arguments, either /path_to_fpmpi/lib/libfpmpi.a or -L/path_to_fpmpi/lib -lfpmpi, which is saved in Makeconf and installed to pbdPROF/etc/ for later use by other packages, such as pbdMPI or Rmpi.

2.4.1 Reinstall pbdMPI

Reinstall **pbdMPI** via

Shell Command

```
R CMD INSTALL pbdMPI_1.0-0.tar.gz --configure-args="--enable-pbdPROF"
```

Package developers who are directly interfacing with MPI (via C or Fortran) should note that pbdMPI/R/get_conf.r and pbdMPI/R/get_lib.r are used in pbdMPI/configure.ac or pbdMPI/configure to determine an appropriate linking flag PROF_LDFLAGS based on preset flags in pbdPROF/etc/Makeconf.

If the internal library is used in **pbdPROF**, then the path to **pbdPROF**/lib/libfpmpi.a is set in the flag PKG_LIBS of **pbdMPI/src/Makevars.in**. If the external library is used in **pbdPROF**, then the linking arguments either <code>/path_to_fpmpi/lib/libfpmpi.a</code> or <code>-L/path_to_fpmpi/lib</code> -lfpmpi is set in the flag PKG_LIBS of <code>pbdMPI/src/Makevars.in</code>. Therefore, the **pbdMPI** can be intercepted by the **fpmpi** library when MPI function calls are evoked.

No mater which library is used, internal or external, the PROF_LDFLAGS in pbdMPI/etc/Makefile provides the linking information to the profiling library. It is also used in PKG_LIBS, which will be exported to other pbdR packages at installation via the flag SPMD_LDFLAGS. Therefore there is no need for additional flags in R CMD INSTALL when reinstalling packages for profiling.

2.4.2 Reinstall pbdBASE

For further profiling, such as **pbdBASE** (Schmidt et al., 2012), one may reinstall the package, via

Shell Command

```
R CMD INSTALL pbdBASE_0.2-2.tar.gz
```

There is no need to provide any flag since **pbdMPI/etc/Makefile** has the information and installation of **pbdBASE** already considers it. Note that since both packages (**pbdMPI** and **pbdBASE**) have MPI-using C/Fortran functions involved, it is necessary to link with **pbdPROF** in order to profile communications evoked by the package.

2.4.3 Reinstall Rmpi

Reinstall Rmpi via

```
wget https://github.com/snoweye/Rmpi_PROF/archive/master.zip
unzip master.zip
mv Rmpi_PROF-master Rmpi
find ./Rmpi -type f -perm 777 -print -exec chmod 644 {} \;
find ./Rmpi -type d -perm 777 -print -exec chmod 755 {} \;
chmod 755 ./Rmpi/configure
chmod 755 ./Rmpi/cleanup
```

```
chmod 755 ./Rmpi/inst/*.sh
R CMD build --no-resave-data Rmpi
R CMD INSTALL Rmpi_0.6-4.tar.gz --configure-args="--enable-pbdPROF'"
```

Note that 0.6-4 is not an official release of **Rmpi**. It is a modified version of 0.6-3 and it is currently available at https://github.com/snoweye/Rmpi_PROF. The authors of **Rmpi** have plans to eventually incorporate these changes, but this can be used as a temporary measure.

2.5 mpiP

We have to install mpiP externally from its source code to use it in pbdPROF. We can install pbdPROF using the external mpiP library via

Shell Command

```
R CMD INSTALL pbdPROF_0.2-0.tar.gz
--configure-args="--with-mpiP='/path/to/your/mpiP/lib/libmpiP.a' "
```

Or

Shell Command

```
R CMD INSTALL pbdPR0F_0.2-0.tar.gz
--configure-args="--with-mpiP='-L/path/to/your/mpiP/lib lmpiP' "
```

Or the conventional method in R console

Shell Command

```
install.packages("pbdPROF",
    configure.args=c("--with-mpiP=/path/to/your/mpiP/lib/libmpiP.a"))
```

Or

Shell Command

pbdPROF/libs/ is empty, i.e., there is no need to build pbdPROF.so. The linking argument is saved in Makeconf and installed to pbdPROF/etc/ for later use by other packages, such as pbdMPI or Rmpi. Since mpiP has external dependency libfpmpi.a on libunwind so while installing mpiP you are suggested to use the below command while configuring mpiP This only provides the linking arguments, either

R Script

```
./configure --disable-libunwind CPPFLAGS="-fPIC -I/usr/lib/openmpi/include" LDFLAGS="-L/usr/lib/openmpi/lib -lmpi"
```

since one has changed the linking so need to reinstall packages depend on CodepbdPROF

2.5.1 Reinstall pbdMPI

Reinstall **pbdMPI** via

```
R CMD INSTALL pbdMPI_1.0-0.tar.gz --configure-args="--enable-pbdPROF"
```

Package developers who are directly interfacing with MPI (via C or Fortran) should note that pbdMPI/R/get_conf.r and pbdMPI/R/get_lib.r are used in pbdMPI/configure.ac or pbdMPI/configure to determine an appropriate linking flag PROF_LDFLAGS based on preset flags in pbdPROF/etc/Makeconf.

if your pbdMPI is correctly installed with all correct linking you will the screenshot just similar to below output during installation of **pbdMPI** or else you might get error

No mater which library is used, internal or external, the PROF_LDFLAGS in pbdMPI/etc/Makefile provides the linking information to the profiling library. It is also used in PKG_LIBS, which will be exported to other pbdR packages at installation via the flag SPMD_LDFLAGS. Therefore there is no need for additional flags in R CMD INSTALL when reinstalling packages for profiling.

2.5.2 Reinstall pbdBASE

For further profiling, such as **pbdBASE** (Schmidt et al., 2012), one may reinstall the package, via

Shell Command

```
R CMD INSTALL pbdBASE_0.2-2.tar.gz
```

There is no need to provide any flag since **pbdMPI/etc/Makefile** has the information and installation of **pbdBASE** already considers it. Note that since both packages (**pbdMPI** and **pbdBASE**) have MPI-using C/Fortran functions involved, it is necessary to link with **pbdPROF** in order to profile communications evoked by the package.

2.5.3 Reinstall Rmpi

Reinstall Rmpi via

```
wget https://github.com/snoweye/Rmpi_PROF/archive/master.zip
unzip master.zip
mv Rmpi_PROF-master Rmpi
```

```
find ./Rmpi -type f -perm 777 -print -exec chmod 644 {} \;
find ./Rmpi -type d -perm 777 -print -exec chmod 755 {} \;
chmod 755 ./Rmpi/configure
chmod 755 ./Rmpi/cleanup
chmod 755 ./Rmpi/inst/*.sh
R CMD build --no-resave-data Rmpi
R CMD INSTALL Rmpi_0.6-4.tar.gz --configure-args="--enable-pbdPROF'"
```

Note that 0.6-4 is not an official release of **Rmpi**. It is a modified version of 0.6-3 and it is currently available at https://github.com/snoweye/Rmpi_PROF. The authors of **Rmpi** have plans to eventually incorporate these changes, but this can be used as a temporary measure.

3 Test Scripts

We provide two short R scripts, one for **pbdMPI** and one for **Rmpi**, to test the installation and profiling capabilities of **pbdPROF**. If the installation is correct, then executing these examples codes should produce profiler output.

3.1 Test with pbdMPI

Below we provide sample scripts to test that the installation of **pbdPROF** was successful. For **pbdMPI**, use:

Test script for pbdMPI

```
### Save this in a file: prof_pbdMPI.r
library(pbdMPI, quiet = TRUE)
init()

set.seed(comm.rank())
x <- allreduce(rnorm(100), op = "sum")

finalize()</pre>
```

and run this code by

R Script

```
mpiexec -np 2 Rscript prof_pbdMPI.r
```

A successful output of fpmpi in the profiling file fpmpi_profile.txt may contain

```
Details for each MPI routine
                Average of sums over all processes
                                              % by message length
                             (max over
                                              processes [rank])
MPI_Allreduce:
                                           0] 050000005000000000000000000
       Calls
                          2
                                      2 [
                :
                                            0] 07000000300000000000000000
       Time
                    3.61e-05
                                3.72e-05 [
                       804
       Data Sent :
                                    804 [
                                            01
                                            0] 0*000000.00000000000000000
       SyncTime :
                    0.00149
                                0.00287 [
       By bin : 1-4 [1,1] [
                                2.5e-05, 2.72e-05] [ 4.1e-05,
```

```
: 513-1024 [1,1] [ 1e-05, 1e-05] [ 1.1e-05, 7.61e-05]
```

In this R script, one MPI C function MPI_Allreduce is called twice and 804 bytes are sent that a hundred of double precision (8 bytes) for 100 normal random variables, and one integer (4 bytes) for checking data type to call the corresponding S4 method.

3.2 Test with Rmpi

For **Rmpi**, use:

Test script for pbdMPI

```
### Save this in a file: prof_Rmpi.r
library(Rmpi, quiet = TRUE)
mpi.comm.dup(0, 1)

set.seed(mpi.comm.rank())
x <- mpi.allreduce(rnorm(100), type = 2, op = "sum")

mpi.quit()</pre>
```

and run this code by

R Script

```
mpiexec -np 2 Rscript prof_Rmpi.r
```

A successful output of fpmpi in the profiling file fpmpi_profile.txt could be

```
Details for each MPI routine
              Average of sums over all processes
                                        % by message length
                         (max over
                                        0.....1
                                                K
                                                       М
                          processes [rank])
MPI_Allreduce:
                                     Calls
                                 1 [
                                     Time
                 4.01e-05
                           4.41e-05 [
      Data Sent :
                     800
                               008
                                     0]
                            0.00204 [
                                     0.00103
      SyncTime
             :
                                  3.6e-05, 4.41e-05] [ 2.79e-05,
      By bin
              : 513-1024
                         [1,1]
                              [
.002041
MPI_Comm_dup:
      Calls
              :
      Time
              :
                 5.81e-05
                 0.000211
      SyncTime
```

Two MPI C functions MPI_Allreduce and MPI_Comm_dup are called one time for each.

4 Profiling with fpmpi

4.1 Demo of pbdMPI

The allreduce.r is originally in **pbdMPI/demo/** and can be profiled by

```
R Script
```

```
mpiexec -np 2 Rscript -e "demo(allreduce,'pbdMPI',ask=F,echo=F)"
```

which will provide an output file fpmpi_profile.txt. Part of output is listed in the next as

```
Processes:
Execute time:
              1.176
Timing Stats: [seconds] [min/max]
                                   [min rank/max rank]
wall-clock: 1.176 sec 1.171488 / 1.180277 0 / 1
user: 0.378 sec 0.360000 / 0.396000
                                  0 / 1
sys: 0.07 sec 0.040000 / 0.100000
                                   1 / 0
Average of sums over all processes
                     Calls
                                Time Msg Length
                                                 %Time by message length
                                          MPI_Allreduce
                        10
                            0.000118
MPI_Barrier
                        21
                              0.0054
Details for each MPI routine
Average of sums over all processes
% by message length
(max over
              processes [rank])
                         K
MPI_Allreduce:
                               10 [ 0] 051004000000000000000000000
 Calls
                   10
         : 0.000118
                        0.000119 [ 0] 061003000000000000000000000
Time
Data Sent :
               188
                            188 [
                                   0]
SyncTime : 0.000312
                        0.000453 [
                                   0] 07.002000000000000000000000
By bin : 1-4 [5,5] [ 7.01e-05, 7.01e-05] [ 0.000117,
: 5-8 [1,1] [ 7.87e-06, 9.06e-06] [ 9.06e-06, 9.06e-06]
: 33-64
           [4,4] [ 3.91e-05, 4.03e-05] [ 4.51e-05,
MPI_Barrier:
 Calls
Time
              0.0054
```

Two MPIC functions MPI_Allreduce and MPI_Barrier are evoked inside this R code. The MPI_Allreduce is called 10 times, span 0.000156 seconds, and 188 bytes are sent. The MPI_Barrier is called 21 times and span 0.00608 seconds.

4.2 Demo of pbdDMAT

The svd.r is originally in pbdDMA/demo/ (Schmidt et al., 2012) and can be profiled by

```
R Script
```

```
mpiexec -np 2 Rscript -e "demo(svd,'pbdDMAT',ask=F,echo=F)"
```

which will provide an output file fpmpi_profile.txt. Part of output is listed in the next as

```
Processes: 2
Execute time: 1.774
Timing Stats: [seconds] [min/max]
                                 [min rank/max rank]
wall-clock: 1.774 sec 1.766181 / 1.781962
user: 0.962 sec 0.956000 / 0.968000
                              1 / 0
sys: 0.046 sec 0.044000 / 0.048000
                                 0 / 1
Average of sums over all processes
                             Time Msg Length %Time by message length
Routine
                   Calls
0.000108
                                       72 0640000000000000000000000000
MPI_Allreduce
                     12
               :
MPI_Barrier
                      8
                          0.000784
Details for each MPI routine
Average of sums over all processes
% by message length
(max over
               processes [rank])
                       K
MPI_Allreduce:
                            12 [
                                  Calls
                12
                               0] 0640000000000000000000000000
Time
          0.000108
                     0.000113 [
Data Sent :
               72
                           72 [
                                0]
SyncTime : 0.000143
                      0.00016 [
                               : 1-4 [6,6] [ 5.44e-05, 6.91e-05] [ 6.91e-05, 8.89e-05]
By bin
: 5-8
      [6,6] [ 4.36e-05, 4.79e-05] [ 5.72e-05, 7.08e-05]
MPI_Barrier:
 Calls
Time
           0.000784
```

Two MPIC functions MPI_Allreduce and MPI_Barrier are evoked inside this R code. The MPI_Allreduce is called 12 times, span 0.000108 seconds, and 72 bytes are sent. The MPI_Barrier is called 8 times and span 0.000784 seconds.

4.3 Demo of Rmpi

The masterSlavePI.r is originally in Rmpi/demo/ and can be profiled by

```
R Script
```

```
mpiexec -np 4 Rscript -e "demo(masterslavePI,'Rmpi',ask=F,echo=F)"
```

which will provide an output file fpmpi_profile.txt. Part of output is listed in the next as

```
Processes: 1

Execute time: 0.05362

Timing Stats: [seconds] [min/max] [min rank/max rank]

wall-clock: 0.05362 sec 0.053622 / 0.053622 0 / 0

user: 0.236 sec 0.236000 / 0.236000 0 / 0

sys: 0.052 sec 0.052000 / 0.052000 0 / 0

Average of sums over all processes

Routine Calls Time Msg Length %Time by message length
```

```
0....................................
MPI_Reduce
                  6.51e-05
                           Details for each MPI routine
Average of sums over all processes
% by message length
(max over
       0.....1.....1.....
processes [rank])
               K
MPI_Reduce:
         1
               Calls
By bin : 5-8 [1,1] [ 6.51e-05, 6.51e-05]
```

One MPI C function MPI_Reduce is evoked inside this R code. The MPI_Reduce is called only 1 time, span 6.51e-05 seconds, and 8 bytes are sent. Note that there is only one processor (master in comm=0) profiled by **fpmpi**, and the other three processors (slaves in comm=1) are not.

5 Profiling with mpiP

5.1 Demo of pbdMPI

The allreduce.r is originally in pbMPI/demo and can be profiled by

```
R Script
```

```
mpiexec -np 2 Rscript -e "demo(allreduce,'pbdMPI',ask=F,echo=F)"
```

which will produce an output file allreduce.r.mpiP part of file is listed below

```
@ Collector Rank
                : 0
                    : 24033
@ Collector PID
@ Final Output Dir
@ Report generation
                    : Single collector task
@ MPI Task Assignment
                    : 0 wolf-vb9
                    : 1 wolf-vb9
@ MPI Task Assignment
 @--- MPI Time (seconds) -----
 Task AppTime MPITime MPI%
    0.153 0.00207 1.35
                    18.35
1
     0.155
            0.0284
     0.308
            0.0305
                   9.90
 @--- Callsites: 6 -----
 ID Lev File/Address
                      Line Parent_Funct
                                              MPI_Call
  0 0x7f335d1108c3
                        [unknown]
                                           Allreduce
  0 0x7f335d110acb
                        [unknown]
                                           Barrier
  0 0x7f335d1107f3
                        [unknown]
3
                                           Allreduce
  0 0x7f2ded6f68c3
                        [unknown]
                                           Allreduce
   0 0x7f2ded6f6acb
                       [unknown]
                                           Barrier
```

6 0 0x7f2ded6f67f3		[unknown]]		Allreduce			
@ Aggregate Tim	e (top tw	enty, desc	ending,	millise	conds)	 		
Call	Site	Time	App%	MPI%	COV			
Barrier	5	28.1	9.13	92.21	0.00			
Barrier	2	1.63	0.53	5.36	0.00			
Allreduce	3	0.322	0.10	1.06	0.00			
Allreduce	6	0.217	0.07	0.71	0.00			
Allreduce	1	0.117	0.04	0.38	0.00			
Allreduce	4	0.083	0.03	0.27	0.00			
@ Aggregate Sen	t Message	Size (top	twenty,	descen	ding, bytes)	 		
Call	Site	Count	Tot	al	Avrg Sent%			
Allreduce	1	4	160		40 42.55			
Allreduce	4	4	160		40 42.55			
Allreduce	3	6	28		4.67 7.45			
Allreduce	6	6	28		4.67 7.45			

The above statistics shows various criteria for the program runned the MPI TIME shows running time per process while executing the allreduce.r.There are four columns Task which is Rank of the processor. In the above sample output there is AppTimewhich is Application level runtime having values 0.153 and 0.155 for first and second ranks respectively ,MPITime which is MPI level runtime of code having value 0.00207 for first rank and 0.0284 for second rank and values 1.35 and 18.35 in MPI% which are percentage of MPITime in AppTime for rank 0 processor and rank 1 respectively. The * shows sum of total ranks in respective column. Furthermore mpiP library provides deeper analysis of each MPI Calls like Aggregate Time and Aggregate Sent Message Size . In Aggregate Time division Call column shows each MPI_Calls used here two are used Barrier and Allreduce. The Barrier calls at Site 5 ran for 28.1 milliseconds of which 9.13 is Application level aggregate time percentage and 92.21 is MPI level aggregate time percentage.

Similarly in Aggregate Sent Message Size division per bytes info of each MPI call is elaborated. For example, for Allreduce at Site 1 has Count value of 4 while Total Message Size is 160 bytes, on average 40 bytes are there. Also Sent percentage is 42.55 for Allreduce at Site 1.

5.2 Demo of pbdDMAT

The svd.r is originally in pbdDMA/demo/ (Schmidt et al., 2012) and can be profiled by

```
R Script

mpiexec -np 2 Rscript -e "demo(svd,'pbdDMAT',ask=F,echo=F)"
```

which will provide an output file svd.r.mpiP. Part of output is listed in the next as

@--- MPI Time (seconds) -----

```
@ Collector Rank : 0
@ Collector PID : 25363
@ Final Output Dir : .
@ Report generation : Single collector task
@ MPI Task Assignment : 0 wolf-vb9
@ MPI Task Assignment : 1 wolf-vb9
```

1	ask	Appl	Time	MPITime	MPI%					
0		0.768	0.0005	27 0	.07					
1		0.784	0.001	95 0	. 25					
*		1.55	0.002		. 16					
() – – –	Callsit	es: 6 -							
]	 [D Le	ev File,	'Address		 Line Paren [.]	 t_Funct		N	MPI_Call	
1	0 (0x7f676e	ef298c3		[unknown]]		Allr	reduce	
2	0 (0x7f676	ef29acb		[unknown]		Barr	rier	
3	0 (0x7f676	ef297f3		[unknown]		Allr	reduce	
4	0 (0x7fa461	caf8c3		[unknown]		Allr	reduce	
5	0 (0x7fa461	lcafacb		[unknown]		Barr	rier	
6	0 (0x7fa461	caf7f3		[unknown]		Allr	reduce	
(66- 06	ice iime	(cop cw	enty, desc	enaing,	millised	onas)	,	
-	 Call			Site	Time	 App%	MPI%		 cov	
- (Bai	Call rrier	 r		Site 5	Time 1.55	App% 0.10	MPI% 62.40	0.00	 COV)	
Baı All	Call rrien Iredu	r uce		Site 5 6	Time 1.55 0.295	App% 0.10 0.02	MPI% 62.40 11.90	0.00	 COV)	
Bai Bai All Bai	Call rien Iredu	r uce r		Site 5 6 2	Time 1.55 0.295 0.256	App% 0.10 0.02 0.02	MPI% 62.40 11.90 10.33	0.00	 COV))	
Bai All Bai	Call rrien Iredu rrien	r uce r uce		Site 5 6 2	Time 1.55 0.295 0.256 0.177	App% 0.10 0.02 0.02 0.01	MPI% 62.40 11.90 10.33 7.14	0.00	COV))))	
Ban All Ban All	Call rien Iredu	r uce r uce uce		Site 5 6 2 3	Time 1.55 0.295 0.256	App% 0.10 0.02 0.02 0.01 0.01	MPI% 62.40 11.90 10.33 7.14	0.00	COV))))	
Ban All Ban All All	Call rrien rredu rredu rredu	r uce r uce uce		Site 5 6 2 3 4	Time 1.55 0.295 0.256 0.177 0.11	App% 0.10 0.02 0.02 0.01 0.01	MPI% 62.40 11.90 10.33 7.14 4.44 3.79	0.00 0.00 0.00 0.00 0.00	COV () () () () () ()	
Ban All Ban All All 	Call rrien rredu rredu rredu	r uce r uce uce		Site 5 6 2 3 4	Time 1.55 0.295 0.256 0.177 0.11 0.094	App% 0.10 0.02 0.02 0.01 0.01 0.01 twenty	MPI% 62.40 11.90 10.33 7.14 4.44 3.79	0.00 0.00 0.00 0.00 0.00 0.00	COV () () () () () ()	
	Call crien crien crien crien credu	r uce r uce uce Aggrega		Site 5 6 2 3 4 1 Message	Time 1.55 0.295 0.256 0.177 0.11 0.094	App% 0.10 0.02 0.02 0.01 0.01 0.01 twenty	MPI% 62.40 11.90 10.33 7.14 4.44 3.79 , descendental	0.00 0.00 0.00 0.00 0.00 0.00	COV)))))) bytes)	
Ban All Ban All All 	Call Trien Tredu Tredu Tredu Tredu Tredu Tredu	r uce r uce uce Aggrega		Site 5 6 2 3 4 1 Message Site	Time 1.55 0.295 0.256 0.177 0.11 0.094	App% 0.10 0.02 0.02 0.01 0.01 0.01	MPI% 62.40 11.90 10.33 7.14 4.44 3.79 , descendental	0.00 0.00 0.00 0.00 0.00 0.00 0.00	COV))))) bytes) g Sent%	
A13 A13 A13 A13 A13 A13 A13 A13	Call rrien rrien rredu redu redu redu redu redu redu	r uce r uce uce ace Aggrega		Site 5 6 2 3 4 1 Message Site 1	Time 1.55 0.295 0.256 0.177 0.11 0.094	App% 0.10 0.02 0.02 0.01 0.01 0.01 twenty 48	MPI% 62.40 11.90 10.33 7.14 4.44 3.79 , descend	0.00 0.00 0.00 0.00 0.00 0.00 0.00 ing,	COV))))) bytes) Sent% 33.33	

The above statistics shows various criteria the code has been profiled for the program runned the MPI TIME shows running time per process while executing the allreduce.r.There are four columns Task which is Rank of the each processor .In the above sample output there is AppTimewhich is Application level runtime having values 0.768 and 0.784 for first and second ranks respectively ,MPITime which is MPI level runtime of code having value 0.000527 for first rank and 0.00195 for second rank and values 0.7 and 0.25 MPI% which are percentage of MPITime in AppTime for rank 0 processor and rank 1 respectively. The * shows sum of total ranks in respective column. Furthermore mpiP library provides deeper analysis of each MPI Calls like Aggregate Time and Aggregate Sent Message Size . In Aggregate Time division Call column shows each MPI_Calls used here two are used Barrier and Allreduce. The Barrier calls at Site 5 ran for 1.5 milliseconds of which 0.10 is Application level aggregate time percentage and 62.40 is MPI level aggregate time percentage.

Similarly in Aggregate Sent Message Size division per bytes info of each MPI call is elaborated. For example, for Allreduce at Site 1 has Count value of 6 while Total Message Size is 48 bytes, on average 8 bytes are there. Also Sent percentage of total bytes is 33.3 for Allreduce at Site 1.

5.3 Demo of Rmpi

The masterSlavePI.r is originally in Rmpi/demo/ and can be profiled by

```
mpiexec -np 4 Rscript -e "demo(masterslavePI,'Rmpi',ask=F,echo=F)"
```

which will provide an output file masterSlavePI.r.mpiP. Part of output is listed in the next as

@ Collector Rank		: 0						
@ Collector PID		: 2	5839					
@ Final Output Di	ir	: .						
@ Report generati		: S	ingle c	ollector	task			
@ MPI Task Assign								
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1								
@ MPI Time								
Task AppTime	e MPI	Time	MPI%					
0 0.0303 (
* 0.0303 (0.00125	4.	12					
@ Callsites:					·			
ID Lev File/Add	dress	L	ine Par	ent_Funct		MPI_	Call	
1 0 0x7f8cdbc03	3628		[unkno	wn]		Comm_fr	еe	
<pre>0 0x7f8cdbc03</pre>	Ba2e		[unkno	wn]		Interco	mm_mer	ge
3 0 0x7f8cdbc02	2ce6		[unkno	wn]		Reduce		
4 0 0x7f8cdbc03	398ъ		[unkno			Comm_fr	ее	
Aggregate	Time (t	op twe	ntv. de	scending.	millisec	onds)		
Call		ite	Tim	e App%	MPI%	COV		
Intercomm_merge	:	2	1.06	3.52	85.47 8.19	0.00		
Reduce	;	3	0.102	0.34	8.19	0.00		
Comm_free	•		0.053		4.25			
Comm_free		1	0.026	0.09	2.09	0.00		
Aggregate	Sent Me	 ssage	Size (t	op twenty	. descend	ing. byt	 es)	
Call					tal	_		
Reduce		3			8	8 100.		
@ Callsite :								
Name	Site	 Rank	Count	Max	Mean	Min	 Ann	 % мрт%
Comm_free	1	0	1	0 026	0.026	0.026		9 2.09
Comm_free	1				0.026			
	-		-	0.020	0.020	0.020	0.00	2.00
Comm_free	4	0	1	0.053	0.053	0.053	0.18	4.25
Comm_free	4	*	1	0.053	0.053	0.053	0.18	4.25
Tn+oncomm mongo	0	0	1	1 06	1.06	1 06	2 50	OE 17
Intercomm_merge	2				1.06			
Intercomm_merge	2	*	1	1.00	1.00	1.00	3.52	85.47
Reduce	3	0	1	0.102	0.102	0.102	0.34	8.19
Reduce	3	*	1	0.102	0.102	0.102	0.34	8.19
@ Callsite N	Message	 Sent s	tatisti	cs (all,	sent byte	s)		
Name	Site	Rank	Count	Ma	x Me	ean	Min	Sum

Reduce	3	0	1	8	8	8	8	
Reduce	3	*	1	8	8	8	8	

The above statistics shows various criteria the code has been profiled for the program runned the MPI TIME shows running time per process while executing the masterSlaveMPI.r. There are four columns Task which is Rank of the each processor. In the above sample output there is AppTimewhich is Application level runtime having values 0.0303 and 0.0303 for first and second ranks respectively ,MPITime which is MPI level runtime of code having value 0.00125 for first rank and 0.00125 for second rank and 4.12 MPI% and 4.12 which is percentage of MPITime in AppTime for rank 0 processor and rank 1 processor respectively. The * shows sum of total ranks in respective column.

Furthermore mpiP library provides deeper analysis of each MPI Calls like Aggregate Time and Aggregate Sent Message Size. In Aggregate Time division Call column shows each MPI_Calls used here two are used Barrier and Allreduce. The Barrier calls at Site 5 ran for 1.5 milliseconds of which 0.10 is Application level aggregate time percentage and 62.40 is MPI level aggregate time percentage.

Similarly in Aggregate Sent Message Size division per bytes info of each MPI call is elaborated. For example, for Allreduce at Site 1 has Count value of 6 while Total Message Size is 48 bytes, on average 8 bytes are there. Also Sent percentage of total bytes is 33.3 for Allreduce at Site 1.

In Callsite Time statistics division further explanation per MPI_Call has been described by factor of Max,Min and Mean. For example the Comm_free Call at Site 1 of Rank 0 has Count value of 1 while Max of various time values is 0.26 and Mean has value of 0.26 and Min also has value of 0.26 since only one processor Rank is used.

6 Visualizing Profiler Outputs

Several useful plotting methods have been provided in the **pbdPROF** package for visualizing fpmpi and mpiP profiler outputs.

In addition, the data is stored in a fairly simple format, so it should be simple enough to create your own plots if these do not suffice.

6.1 Visualizing fpmpi Profiler Output

6.2 Visualizing mpiP Profiler Output

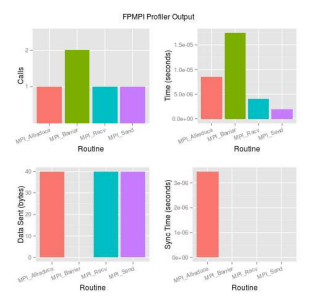
7 Problems with pbdPROF

7.1 Installation

Problem 1: If you have downloaded the package from github and tried to using R CMD INSTALL pbdPROF and you see an error similar to this

```
ERROR: 'configure' exists but is not executable -- see the 'R Installation and Administration Manual'
```

Solution: You have to make the configure executable which means giving it permission , which can done by



fpmpi Plots

R Script

```
chmod +x configure
```

after changing the folder to package's main directory.

Problem 2: If you are using fpmpi (Gropp, 2000) externally and during it's installation you get an error similar to this

```
error : checking for library containing MPI_Init... (cached) no configure: error: Could not find MPI library
```

Solution: You probably need to specify the path to MPI library using this in command line in the fpmpi main directory

R Script

```
./configure CPPFLAGS="-fPIC -I/usr/lib/openmpi/include"
LDFLAGS="-L/usr/lib/openmpi/lib -lmpi"
```

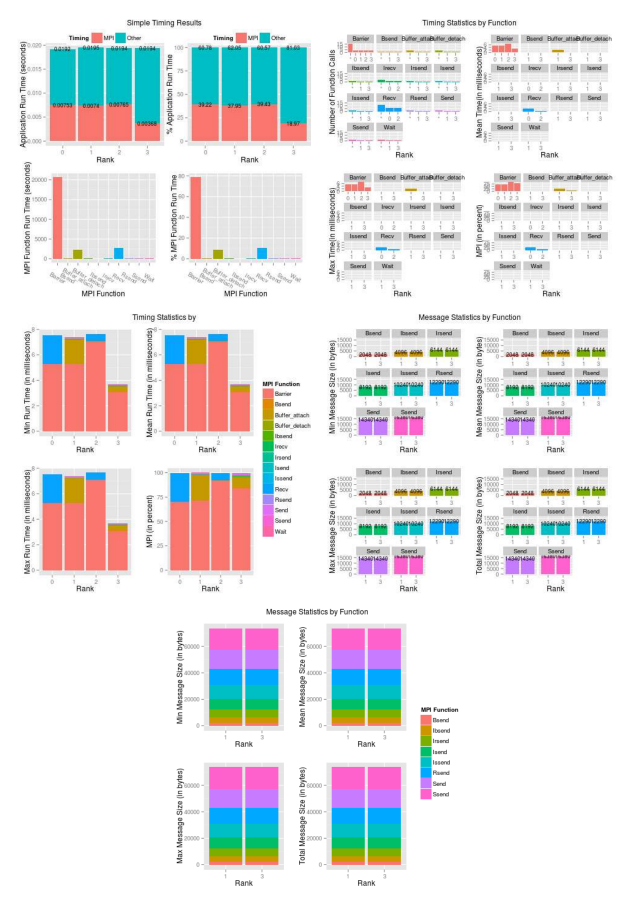
Problem 3: If you are using mpiP (Vetter and McCracken, 2001) externally and during it's installation you get an error similar to this

```
libmpiP.a(wrappers.o): relocation R_X86_64_32 against '.rodata.str1.1' can not
  be used when making a shared object; recompile with -fPIC
libmpiP.a: could not read symbols: Bad value collect2: error: ld returned 1
  exit status
```

Solution: You probably need to specify the path to MPI library using this in command line when installing \mathbf{mpiP}

```
R Script
```

./configure CPPFLAGS="-fPIC -I/usr/lib/openmpi/include"



mpiP Plots

```
LDFLAGS="-L/usr/lib/openmpi/lib -lmpi"
```

Problem 4: If you are using mpiP (Vetter and McCracken, 2001) externally and during pbdMPI (Chen et al., 2012a) installation you get an error similar to this

```
Error : .onLoad failed in loadNamespace() for 'pbdMPI', details:
call: dyn.load(file, DLLpath = DLLpath, ...)
error: unable to load shared object 'pbdMPI.so':
pbdMPI/libs/pbdMPI.so: undefined symbol: _Ux86_64_getcontext
```

Solution: You probably need to disable some external library prerequisite by **mpiP**, using this in command line when installing **mpiP**

R Script

```
./configure --disable-libunwind CPPFLAGS="-fPIC -I/usr/lib/openmpi/include" LDFLAGS="-L/usr/lib/openmpi/lib -lmpi"
```

7.2 Running

Problem 1: While running Rmpi code for profiling, if you encounter the error below:

```
error: mpiexec was unable to launch the specified application as it could not access or execute an executable:

Executable: /path/to/R/package_installation_directory/2.15/Rmpi/Rslaves.sh

Node: "Your_node"

while attempting to start process rank 0.
```

Solution: You need to make executable of the shell scripts in the "inst" directory of "Rmpi" main directory using the following command from command line in "inst" directory:

R Script

```
chmod +x *.sh
```

Problem 2: While running Rmpi code for profiling, if you encounter the error below:

```
[G:12221] [[39704,0],0] ORTE_ERROR_LOG: Not found in file
../../../../orte/mca/plm/base/plm_base_launch_support.c at line 758

mpiexec was unable to start the specified application as it encountered an error.

More information may be available above.
```

Solution:

- 1. You need to check whether your **Rmpi** is working without the **pbdPROF**. If yes try running your **Rmpi** code on single process only.
- 2. If above does not help, then you may need .Rprofile in Rmpi/inst/ to run your code from "inst" directory.
- 3. If still your code does not run ,you need to update your OPENMPI version to the latest one. You

can check your openmpi versionhttp://www.open-mpi.org/software/ompi/through

ompi_info

4. If further you came to this far and luck is not with you somehow(pun intended), there might some configuration problem in your machine.

8 References

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