

## Intel® MPI Benchmarks

**User Guide and Methodology Description** 

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# Contents

•	nformation	
Getting	g Help and Support	8
Si	ubmitting Issues	8
Introdu	uction	9
Ir	ntended Audience	9
Ir	ntroducing Intel(R) MPI Benchmarks	9
W	Vhat's New	10
	Changes in Intel® MPI Benchmarks 4.0	10
	Changes in Intel® MPI Benchmarks 3.2.4	10
	Changes in Intel® MPI Benchmarks 3.2.3	10
	Changes in Intel® MPI Benchmarks 3.2.2	10
	Changes in Intel® MPI Benchmarks 3.2.1	11
	Changes in Intel® MPI Benchmarks 3.2	11
	Changes in Intel® MPI Benchmarks 3.1	12
	Changes in Intel® MPI Benchmarks 3.0	13
N	lotational Conventions	13
D	Occument Version Information	14
R	elated Information	16
Installa	ation and Quick Start	17
М	lemory and Disk Space Requirements	17
	oftware Requirements	
Ir	nstalling Intel® MPI Benchmarks	18
В	uilding Intel(R) MPI Benchmarks	19
	Building Intel® MPI Benchmarks on Linux* OS	19
	Building Intel® MPI Benchmarks on Windows* OS	
R	unning Intel® MPI Benchmarks	22
R	unning Benchmarks in Multiple Mode	22
MPI-1 I	Benchmarks	24
C	lassification of MPI-1 Benchmarks	25
	Single Transfer Benchmarks	25
	Parallel Transfer Benchmarks	25
	Collective Benchmarks	26
Si	ingle Transfer Benchmarks	27
	PingPong, PingPongSpecificSource	27
	PingPing, PingPingSpecificSource	28
Pa	arallel Transfer Benchmarks	29
	Sendrecv	29
	Exchange	30
C	ollective Benchmarks	31
	Reduce	32
	Reduce_scatter	32
	Allreduce	33
	Allgather	33
	Allgatherv	34
	Scatter	34
	Scatterv	35
	Gather	35

	Gatherv	35
	Alltoall	36
	Bcast	36
	Barrier	36
MPI-	-2 Benchmarks	38
	Naming Conventions	40
	IMB-MPI-2 Benchmark Classification	41
	Single Transfer Benchmarks	41
	Parallel Transfer Benchmarks	41
	Collective Benchmarks	42
	MPI-2 Benchmarks Classification	42
	MPI-2 Benchmark Modes	43
	Assured Completion of Transfers	44
	IMB-EXT Benchmarks	44
	Unidir_Put	44
	Unidir_Get	45
	Bidir_Put	46
	Bidir_Get	47
	Accumulate	48
	Window	49
	IMB-IO Blocking Benchmarks	50
	S_[ACTION]_indv	
	S_[ACTION]_expl	52
	P_[ACTION]_indv	
	P_ACTION_expl	55
	P_[ACTION]_shared	56
	P_[ACTION]_priv	58
	C_[ACTION]_indv	59
	C_[ACTION]_expl	59
	C_[ACTION]_shared	60
	Open_Close	60
	IMB-IO Non-blocking Benchmarks	61
	Exploiting CPU	62
	Displaying Results	62
MPI-	-3 Benchmarks	63
	IMB-NBC Benchmarks	63
	Measuring Communication and Computation Overlap	64
	Measuring Pure Communication Time	65
	lallgather	65
	lallgather_pure	66
	lallgatherv	66
	lallgatherv_pure	66
	Iallreduce	
	Iallreduce_pure	67
	Ialltoall	
	Ialltoall_pure	
	Ialitoally	68

Ialltoallv_pure	69
Ibarrier	69
Ibarrier_pure	70
Ibcast	70
Ibcast_pure	70
Igather	<b>7</b> 1
Igather_pure	<b>7</b> 1
Igatherv	72
Igatherv_pure	72
Ireduce	73
Ireduce_pure	73
Ireduce_scatter	74
Ireduce_scatter_pure	74
Iscatter	75
Iscatter_pure	7 <i>6</i>
Iscatterv	7 <i>6</i>
Iscatterv_pure	77
IMB-RMA Benchmarks	77
IMB-RMA Benchmark Modes	77
Classification of IMB-RMA Benchmarks	77
Accumulate	79
All_get_all	
All_put_all	80
Bidir_get	80
Bidir_put	81
Compare_and_swap	81
Exchange_Get	81
Exchange_Put	82
Fetch_and_op	82
Get_accumulate	83
Get_all_local	83
Get_local	84
One_put_all	84
One_get_all	85
Put_all_local	85
Put_local	85
Truly_passive_put	86
Unidir_get	8 <i>6</i>
Unidir_put	87
Benchmark Methodology	88
Control Flow	88
Command-line Control	89
Benchmark Selection Arguments	90
-npmin Option	90
-multi outflag Option	
-off_cache cache_size[,cache_line_size] Option	
-iter Option	92

-iter_policy Option	92
-time Option	93
-mem Option	93
-input <file> Option</file>	93
-msglen <file> Option</file>	94
-map PxQ Option	94
-include [[benchmark1] benchmark2]	95
-exclude [[benchmark1] benchmark2]	95
-msglog [ <minlog>:]<maxlog></maxlog></minlog>	95
-thread_level Option	96
Parameters Controlling Intel® MPI Benchmarks	96
Hard-Coded Settings	99
Communicators, Active Processes	99
Other Preparations for Benchmarking	99
Message/I-O Buffer Lengths	101
Buffer Initialization	101
Warm-Up Phase (IMB-MPI1, IMB-EXT, IMB-NBC, and IMB-RMA)	101
Synchronization	101
Actual Benchmarking	102
Checking Results	104
Output	104
Sample 1 - IMB-MPI1 PingPong Allreduce	105
Sample 2 - IMB-MPI1 PingPing Allreduce	108
Sample 3 - IMB-IO p_write_indv	112
Sample 4 - IMB-EXT.exe	115

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Your feedback is very important to us. To receive technical support for the tools provided in this product and technical information including FAQ's and product updates, you need to register for an Intel(R) Premier Support account at the Registration Center.

This package is supported by Intel(R) Premier Support. Direct customer support requests at:

#### https://premier.intel.com

General information on Intel(R) product-support offerings may be obtained at:

#### http://www.intel.com/software/products/support

The Intel(R) MPI Benchmarks home page can be found at:

### http://www.intel.com/go/imb

When submitting a support issue to Intel(R) Premier Support, please provide specific details of your problem, including:

- The Intel(R) MPI Benchmarks package name and version information
- Host architecture (for example, Intel(R) 64 architecture)
- Compiler(s) and versions
- Operating system(s) and versions
- Specifics on how to reproduce the problem. Include makefiles, command lines, small test cases, and build instructions.

## **Submitting Issues**

- 1. Go to <a href="https://premier.intel.com">https://premier.intel.com</a>
- 2. Log in to the site. Note that your username and password are case-sensitive.
- 3. Click on the **Submit Issue** link in the left navigation bar.
- 4. Choose Development Environment (tools, SDV, EAP) from the Product Type drop-down list. If this is a software or license-related issue, choose the Intel(R) Cluster Studio [XE], Linux\* or the Intel(R) Cluster Studio [XE], Windows\* from the Product Name drop-down list.
- 5. Enter your question and complete the required fields to successfully submit the issue.

#### NOTE:

Notify your support representative prior to submitting source code where access needs to be restricted to certain countries to determine if this request can be accommodated.

## Introduction

This guide presents the Intel® MPI Benchmarks 4.0. The objectives of the Intel® MPI Benchmarks are:

- Provide a concise set of benchmarks targeted at measuring the most important MPI functions.
- Set forth a precise benchmark methodology.
- Report bare timings rather than provide interpretation of the measured results. Show throughput values if and only if these values are well-defined.

Intel® MPI Benchmarks is developed using ANSI C plus standard MPI.

Intel® MPI Benchmarks is distributed as an open source project to enable use of benchmarks across various cluster architectures and MPI implementations.

## **Intended Audience**

This guide is intended for users who want to measure performance of MPI implementations.

## Introducing Intel(R) MPI Benchmarks

Intel® MPI Benchmarks performs a set of MPI performance measurements for point-to-point and global communication operations for a range of message sizes. The generated benchmark data fully characterizes:

- performance of a cluster system, including node performance, network latency, and throughput
- efficiency of the MPI implementation used

The Intel® MPI Benchmarks package consists of the following components:

- IMB-MPI1 benchmarks for MPI-1 functions.
- Two components for MPI-2 functionality:
  - IMB-EXT one-sided communications benchmarks.
  - IMB-IO input/output (I/O) benchmarks.
- Two components for MPI-3 functionality:
  - IMB-NBC benchmarks for nonblocking collective (NBC) operations.
  - *IMB-RMA* one-sided communications benchmarks. These benchmarks measure the Remote Memory Access (RMA) functionality introduced in the MPI-3 standard.

Each component constitutes a separate executable file. You can run all of the supported benchmarks, or specify a single executable file in the command line to get results for a specific subset of benchmarks.

If you do not have the MPI-2 or MPI-3 extensions available, you can install and use IMB-MPI1 that uses only standard MPI-1 functions.

## What's New

This section provides changes for the Intel® MPI Benchmarks as compared to the previous versions of this product.

## Changes in Intel® MPI Benchmarks 4.0

This release includes the following updates as compared to the Intel® MPI Benchmarks 3.2.4:

- Introduced new components IMB-NBC and IMB-RMA that conform to the MPI-3.0 standard.
- Introduced a new feature to set the appropriate policy for automatic calculation of iterations. You can set the policy using the -iter and -iter\_policy options.
- Added new targets to the Linux\* OS Makefiles:
  - NBC for building IMB-NBC
  - RMA for building IMB-RMA
- Updated Microsoft\* Visual Studio\* solutions to include the IMB-NBC and IMB-RMA targets.
- Support for the Microsoft\* Visual Studio\* 2013. Microsoft\* Visual Studio\* 2008 support is removed.

### Changes in Intel® MPI Benchmarks 3.2.4

This release includes the following updates as compared to the Intel® MPI Benchmarks 3.2.3:

Changes of document layout.

### Changes in Intel® MPI Benchmarks 3.2.3

This release includes the following updates as compared to the Intel® MPI Benchmarks 3.2.2:

- Option -msglog to control the message length. Use this option to control the maximum and the second largest minimum of the message transfer sizes. The minimum message transfer size is always 0.
- Thread safety support in the MPI initialization phase. Use MPI\_Init() by default because it is supported for all MPI implementations. You can choose MPI\_Init\_thread() by defining the appropriate macro.
- Option -thread\_level to specify the desired thread level support for MPI\_Init\_thread.
- Support for the Microsoft\* Visual Studio\* 2010 project folder.

### Changes in Intel® MPI Benchmarks 3.2.2

This release includes the following updates as compared to the Intel® MPI Benchmarks 3.2.1:

- Support for large buffers greater than 2 GB for some MPI collective benchmarks (Allgather, Alltoall, Scatter, Gather) to support large core counts.
- New benchmarks: PingPongSpecificSource and PingPingSpecificSource. The exact destination rank is used for these tests instead of MPI\_ANY\_SOURCE as in the PingPong and

PingPing benchmarks. These are not executed by default. Use the -include option to enable the new benchmarks. For example,

```
\ mpirun n 2 IMB_MPI -include PingPongSpecificSource \ PingPingSpecificSource
```

• New options -include/-exclude for better control over the benchmarks list. Use these options to include or exclude benchmarks from the default execution list.

## Changes in Intel® MPI Benchmarks 3.2.1

This release includes the following updates as compared to the Intel® MPI Benchmarks 3.2:

- Fix of the memory corruption issue when the command-line option -msglen is used with the Intel® MPI Benchmarks executable files.
- Fix in the accumulated benchmark related to using the CHECK conditional compilation macro.
- Fix for the integer overflow in dynamic calculations on the number of iterations.
- Recipes for building IA-32 executable files within Microsoft\* Visual Studio\* 2005 and Microsoft\* Visual Studio\* 2008 project folders associated with the Intel® MPI Benchmarks.

## Changes in Intel® MPI Benchmarks 3.2

Intel® MPI Benchmarks 3.2 has the following changes as compared to the previous version:

- The default settings are different.
- Microsoft\* Visual Studio\* project folders are added and can be used on the Microsoft\* Windows\* platforms.
- Makefiles for the Microsoft\* Windows\* nmake utility provided with the Intel® MPI Benchmarks 3.1 are removed.

### Run Time Control by Default

The improved run time control that is associated with the -time flag. This is the default value for the Intel® MPI Benchmarks executable files (with a maximum run time per sample set to 10 seconds by the SECS\_PER\_SAMPLE parameter in the include file IMB\_settings.h).

#### **Makefiles**

The nmake files for Windows\* OS were removed and replaced by Microsoft\* Visual Studio\* solutions.

The Linux\* OS Makefiles received new targets:

- Target MPI1 (default) for building IMB-MPI1
- Target EXT for building IMB-EXT
- Target IO for building IMB-IO
- Target all for building all three of the above

#### Microsoft\* Visual Studio\* Project Folders

Intel® MPI Benchmarks 3.2 contains Microsoft\* Visual Studio\* solutions based on an installation of the Intel® MPI Library. A dedicated folder is created for the Microsoft\* Windows\* OS without duplicating source files. The solutions refer to the source files that are located at their standard location within the Intel® MPI Benchmarks directory structure.

As such solutions are highly version-dependent, see the information in the corresponding ReadMe.txt files that unpack with the folder. You are recommended to learn about the Microsoft\* Visual Studio\* philosophy and the run time environment of your Windows cluster.

## Changes in Intel® MPI Benchmarks 3.1

This release includes the following updates as compared to the Intel® MPI Benchmarks 3.0:

- New control flags
- Better control of the overall repetition counts, run time, and memory exploitation
- A facility to avoid cache re-usage of message buffers as far as possible
- A fix of IMB-IO semantics
- New benchmarks
  - Gather
  - Gatherv
  - Scatter
  - Scatterv
- New command-line flags for better control
  - -off\_cache

Use this flag when measuring performance on high speed interconnects or, in particular, across the shared memory within a node. Traditional Intel® MPI Benchmarks results included a very beneficial cache re-usage of message buffers which led to idealistic results. The flag -off\_cache allows avoiding cache effects and lets the Intel® MPI Benchmarks use message buffers which are very likely not resident in cache.

-iter, -time

Use these flags for enhanced control of the overall run time, which is crucial for large clusters, where collectives tend to run extremely long in the traditional Intel® MPI Benchmarks settings.

#### CAUTION

In the Intel® MPI Benchmarks, the -time flag has been implemented as default.

-mem

Use this flag to determine an a priori maximum (per process) memory usage of the Intel® MPI Benchmarks for the overall message buffers.

### Miscellaneous Changes

In the Exchange benchmark, the two buffers sent by MPI\_Isend are separate. The command line is repeated in the output. Memory management is completely encapsulated in the functions IMB\_v\_alloc/IMB\_v\_free.

## Changes in Intel® MPI Benchmarks 3.0

This release includes the following updates as compared to the Intel® MPI Benchmarks 2.3:

- A call to the MPI\_Init\_thread function to determine the MPI threading environment. The
  MPI threading environment is reported each time an Intel® MPI Benchmark application is
  executed.
- A call to the function MPI\_Get\_version to report the version of the Intel MPI library implementation that the three benchmark applications are linking to.
- New Alltoally benchmark.
- New command-line flag -h[elp] to display the calling sequence for each benchmark application.
- Removal of the outdated Makefile templates. There are three complete makefiles called Makefile, make\_ict, and make\_mpich. The make\_ict option uses the Intel® Composer XE compilers. This option is available for both Intel and non-Intel microprocessors but it may result in additional optimizations for Intel microprocessors.
- Better command-line argument checking, clean message and break on most invalid arguments.

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## **Notational Conventions**

The following conventions are used in this document:

Style	Description
This type style	Commands, arguments, options, file names
THIS_TYPE_STYLE	Environment variables
<this style="" type=""></this>	Placeholders for actual values
[items]	Optional items
{ item   item }	Selectable items separated by vertical bar(s)

# **Document Version Information**

Document Number	Revision Number	Description	Revision Date
320714-001	2.3	Initial version	/10/2004
320714-002	3.0	<ul> <li>The following topics were added:</li> <li>Descriptions of environment amendments</li> <li>The Alltoallv</li> </ul>	/06/2006
320714-003	3.1	The following updates were added:  • Description of Windows version  • Four new benchmarks (Scatter(v), Gather(v))  • IMB-IO functional fix	/07/2007
320714-004	3.2	The following topics were added:  Run time control as default  Microsoft* Visual Studio* solution templates	/08/2008
320714-005	3.2.1	The following updates were added:	/04/2010

		<ul> <li>Fix of the memory corruption</li> <li>Fix in accumulate benchmark related to using the CHECK conditional compilation macro</li> <li>Fix for integer overflow in dynamic calculations on the number of iterations</li> <li>Recipes for building IA-32 executable files within Microsoft* Visual Studio* 2005 and Microsoft* Visual Studio* 2008 project folders associated with the Intel® MPI Benchmarks</li> </ul>	
320714-006	3.2.2	<ul> <li>The following updates were added:</li> <li>Support for large buffers greater than 2 GB for some MPI benchmark</li> <li>New benchmarks         PingPongSpecificSource and PingPingSpecificSource     </li> <li>New options -include/-exclude</li> </ul>	/09/2010
320714-007	3.2.3	The following topics were updated and added:  Changes in the Intel® MPI Benchmarks 3.2.3  Command-line Control  Parameters Controlling IMB  Microsoft* Visual Studio* 2010 project folder support	/08/2011
320714-008	3.2.4	The following updates were added:  Changes of document layout	/06/2012
320714-009	3.2.4	The following updates were added:  • Merged What's new section	/06/2013
320714-010	4.0 Beta	Documented new benchmarks that conform to the MPI-3.0 standard.	/10/2013

320714-011	4.0 Beta Update 1	<ul><li>The following updates were added:</li><li>New option -iter_policy</li><li>Changes in -iter option</li></ul>	/03/2014
320714-012	4.0	Document enhancements	/05/2014

# **Related Information**

For more information, you can see the following related resources:

Intel® MPI Benchmarks Download

Intel® MPI Library Product

# Installation and Quick Start

This section explains how to install and start using the Intel® MPI Benchmarks.

# Memory and Disk Space Requirements

The table below lists memory requirements for benchmarks run with the default settings (standard mode) and with the user-defined settings (optional mode). In this table:

- Q is the number of active processes.
- X is the maximal size of the passing message.

Benchmarks	Standard Mode	Optional Mode
Alltoall	Q*8 MB	Q*2X bytes
Allgather, Allgatherv	(Q+1)*4 MB	(Q+1)*X bytes
Exchange	12 MB	3X bytes
All other MPI-1 benchmarks	8 MB	2X bytes
IMB-EXT	80 MB	2 max(X,OVERALL_VOL) bytes
IMB-IO	32 MB	3X bytes
Ialltoall, Ialltoall_pure	Q*8 MB	Q*2X bytes
Iallgather, Iallgatherv, Iallgather_pure, Iallgatherv_pure	(Q+1)*4 MB	(Q+1)*X bytes
All other IMB-NBC benchmarks	8 MB	2X bytes
Compare_and_swap	12 B	12 B
Exchange_put, Exchange_get	16 MB	4X bytes
All other IMB-RMA benchmarks	8 MB	2X bytes

#### NOTE:

If you do not select the -cache flag, add 2X cache size to all of the above.

For IMB-IO benchmarks, make sure you have enough disk space available:

- 16MB in the standard mode
- max(X,OVERALL\_VOL) bytes in the optional mode

For instructions on enabling the optional mode, see <u>Parameters Controlling Intel® MPI</u> Benchmarks.

## Software Requirements

To run the Intel® MPI Benchmarks, you need:

- cpp, ANSI C compiler, gmake on Linux\* OS or Unix\* OS.
- Enclosed Microsoft Visual\* C++ solutions as the basis for Microsoft Windows\* OS.
- MPI installation, including a startup mechanism for parallel MPI programs.

## Installing Intel® MPI Benchmarks

To install the Intel® MPI Benchmarks, unpack the installation file. The installation directory structure is as follows:

- ReadMe\_IMB.txt
- ./doc documentation directory that contains the User's guide, in PDF and HTML Uncompressed Help formats:
  - IMB\_Users\_Guide.pdf
  - IMB\_Users\_Guide.htm
- ./license license agreement directory that contains the following files:
  - license.txt specifies the source code license granted to you.
  - use-of-trademark-license.txt specifies the license for using the name and/or trademark of the Intel® MPI Benchmarks.
- ./src program source- and Make-files.
- ./WINDOWS Microsoft\* Visual Studio\* solution files.

For basic instructions on how to use the Intel® MPI Benchmarks, see ReadMe\_IMB.txt.

#### See Also

Building Intel® MPI Benchmarks

# Building Intel(R) MPI Benchmarks

This section describes how to build the Intel® MPI Benchmarks on different operating systems.

## Building Intel® MPI Benchmarks on Linux\* OS

To build the benchmarks for Linux\* OS, do the following:

- 1. Set the CC variable to point to the appropriate compiler wrapper, mpicc or mpicc.
- 2. Run one or more makefile commands listed below.

Command	Description
make clean	Remove legacy binary object files and executable files
make MPI1	Build the executable file for the IMB-MPI1 component.
make EXT	Build the executable file for one-sided communications benchmarks.
make IO	Build the executable file for I/O benchmarks.
make NBC	Build the executable file for IMB-NBC benchmarks.
make RMA	Build the executable file for IMB-RMA benchmarks.
make all	Build all executable files available.

To build the benchmarks for Intel® Many Integrated Core Architecture (Intel® MIC Architecture), follow these steps:

1. Build the Intel MPI Benchmarks for the host system:

```
host$ source <install-dir>/composer_xe/bin/compilervars.sh intel64
host$ source <install-dir>/intel64/bin/mpivars.sh
host$ cd <path to IMB directory>/src
host$ make -f make_ict
where
<install-dir>/composer_xe refers to the Intel® Composer XE installation directory
and
```

<install-dir> refers to the Intel® MPI Library installation directory

#### NOTE:

If you have already sourced ictvars.sh for the Bourne command-line shell, you can skip the sourcing of the environment variables controlled by compilervars.sh and mpivars.sh.

2. Build the Intel MPI Benchmarks for the target system based on the Intel® MIC Architecture.

```
host$ cd <path to IMB directory>/src
host$ make -f make ict mic
```

For details on running the resulting executable files on the Intel® MIC Architecture, see the Intel® MPI Library documentation.

#### **See Also**

Running Intel® MPI Benchmarks

### Building Intel® MPI Benchmarks on Windows\* OS

To build the benchmarks for IMB-MPI1, IMB-IO, IMB-EXT, IMB-NBC, or IMB-RMA, follow these steps:

1. Check the environment variable settings for Include, Lib, and Path. Make sure they are set in accordance with this table:

Intel® 64 Architecture Settings	IA-32 Architecture Settings
%I_MPI_ROOT%\intel64\include	%I_MPI_ROOT%\ia32\include
%I_MPI_ROOT%\intel64\lib	%I_MPI_ROOT%\ia32\lib
%I_MPI_ROOT%\intel64\bin	%I_MPI_ROOT%\ia32\bin

#### NOTE:

Intel® MPI Library 5.0 does not support the IA-32 architecture. Use an earlier version of Intel MPI Library to build IA-32 architecture benchmarks.

- 2. Go to the subfolder that corresponds to the Intel® MPI Benchmarks component you would like to build and the Microsoft\* Visual Studio\* version installed on your system. For example, to build IMB-EXT.exe with the Visual\* Studio\* 2010, go to IMB-EXT\_VS\_2010.
- 3. Open the .vcproj or .vcxproj file in Visual Studio\*. The executable file for one of the Intel MPI Benchmarks components is created:
  - IMB-EXT.exe
  - IMB-IO.exe
  - IMB-MPI1.exe
  - IMB-NBC.exe

- IMB-RMA.exe
- 4. From the **Solution Platforms** drop-down list, choose the required architecture (**x64** or **Win32**).
- 5. From the **Solution Configurations** drop-down list, choose **Release**.
- 6. Highlight the project folder in the **Solution Explorer**.
- 7. Go to **Project > Properties** to open **Configuration Properties** dialog box. Make sure you have something like the following settings:

Setting	Value	Notes		
General > Project Defaults				
Character Set	Use Multi-Byte Character Set			
	Debugging			
Debugger to launch	Local Windows Debugger	Depending on your system configuration, you may select other debuggers.		
Command	<ul> <li>x64: \$(I_MPI_ROOT)\intel64\bin\mpiexec.exe</li> <li>IA-32:\$(I_MPI_ROOT)\ia32\bin\mpiexec.exe</li> </ul>			
Command Arguments	-n 2 \$(TargetPath)	\$(TargetPath) should be quoted as in: - n 2 \$(TargetPath)		
	C/C++ > General			
Additional Include Directories	<ul><li>x64: \$(I_MPI_ROOT)\intel64\include</li><li>IA-32:\$(I_MPI_ROOT)\ia32\include</li></ul>			
Warning Level	to Level 1 (/W1)			
	C/C++ > Preprocessor			
Preprocessor Definition	<ul> <li>IMB-EXT: WIN_IMB,         _CRT_SECURE_NO_DEPRECATE, EXT</li> <li>IMB-IO: WIN_IMB,         _CRT_SECURE_NO_DEPRECATE, MPIIO</li> <li>IMB-MPI1: WIN_IMB,         _CRT_SECURE_NO_DEPRECATE, MPII</li> <li>IMB-NBC: WIN_IMB,         _CRT_SECURE_NO_DEPRECATE, NBC</li> <li>IMB-RMA: WIN_IMB,</li> </ul>			

	_CRT_SECURE_NO_DEPRECATE, RMA	
	Linker > Input	
Additional Dependencies	<ul><li>x64: \$(I_MPI_ROOT)\intel64\lib\impi.lib</li><li>IA-32: \$(I_MPI_ROOT)\ia32\lib\impi.lib</li></ul>	

- 8. Go to **Build > Build Solution** to create an executable file.
- 9. Run the executable file using **Debug > Start Without Debugging** command.

## Running Intel® MPI Benchmarks

To run the Intel® MPI Benchmarks, use the following command-line syntax:

```
mpirun -np <P> IMB-<component> [arguments]
```

#### where

- <P> is the number of processes. P=1 is recommended for all I/O and message passing benchmarks except the single transfer ones.
- <component> is the component-specific suffix that can take MPI1, EXT, IO, NBC, and RMA
  values.

By default, all benchmarks run on Q active processes defined as follows:

```
Q=[1,] 2, 4, 8, ..., largest 2^{x}
```

For example, if P=11, the benchmarks run on Q=[1,]2,4,8,11 active processes. Single transfer IMB-IO benchmarks run with Q=1. Single transfer IMB-EXT and IMB-RMA benchmarks run with Q=2.

To pass control arguments other than P, you can use (argc,argv). Process 0 in MPI\_COMM\_WORLD reads all command-line arguments and broadcasts them to all other processes. Control arguments can define various features, such as time measurement, message length, and selection of communicators. For details, see <a href="Command-Line Control">Command-Line Control</a>.

#### See Also

<u>Command-Line Control</u>

<u>Parameters Controlling Intel® MPI Benchmarks</u>

## Running Benchmarks in Multiple Mode

Intel® MPI Benchmarks provides a set of elementary MPI benchmarks.

You can run all benchmarks in the following modes:

- standard (default) the benchmarks run in a single process group.
- multiple the benchmarks run in several process groups.

To run the benchmarks in the multiple mode, add the multi- prefix to the benchmark name.

In the multiple mode, the number of groups may differ depending on the benchmark. For example, if PingPong is running on N≥4 processes, N/2 separate groups of two processes are formed. These process groups are running PingPong simultaneously. Thus, the benchmarks of the single transfer class behave as parallel transfer benchmarks when run in the multiple mode.

#### **See Also**

Classification of MPI-1 Benchmarks Classification of MPI-2 Benchmarks MPI-3 Benchmarks

# MPI-1 Benchmarks

IMB-MPI1 component of the Intel® MPI Benchmarks provides benchmarks for MPI-1 functions. IMB-MPI1 contains the following benchmarks:

Standard Mode	Multiple Mode
PingPong	Multi-PingPong
PingPongSpecificSource (excluded by default)	Multi-PingPongSpecificSource (excluded by default)
PingPing	Multi-PingPing
PingPingSpecificSource (excluded by default)	Multi-PingPingSpecificSource (excluded by default)
Sendrecv	Multi-Sendrecv
Exchange	Multi-Exchange
Bcast	Multi-Bcast
Allgather	Multi-Allgather
Allgatherv	Multi-Allgatherv
Scatter	Multi-Scatter
Scatterv	Multi-Scatterv
Gather	Multi-Gather
Gatherv	Multi-Gatherv
Alltoall	Multi-Alltoall
Alltoallv	Multi-Alltoallv
Reduce	Multi-Reduce
Reduce_scatter	Multi-Reduce_scatter
Allreduce	Multi-Allreduce

Barrier	Multi-Barrier

## Classification of MPI-1 Benchmarks

Intel® MPI Benchmarks introduces the following classes of benchmarks:

- Single Transfer
- Parallel Transfer
- Collective benchmarks

Each class interprets results in a different way.

## Single Transfer Benchmarks

Single transfer benchmarks involve two active processes into communication. Other processes wait for the communication completion. Each benchmark is run with varying message lengths. The timing is averaged between two processes. The basic MPI data type for all messages is MPI\_BYTE.

Throughput values are measured in MBps and can be calculated as follows:

```
throughput = X/2^{20} * 10^6/time = X/1.048576/time,
```

#### where

- time is measured in μ sec.
- x is the length of a message, in bytes.

#### Parallel Transfer Benchmarks

Parallel transfer benchmarks involve more than two active processes into communication. Each benchmark runs with varying message lengths. The timing is averaged over multiple samples. The basic MPI data type for all messages is MPI\_BYTE. The throughput calculations of the benchmarks take into account the multiplicity nmsg of messages outgoing from or incoming to a particular process. For the Sendrecv benchmark, a particular process sends and receives X bytes, the turnover is 2X bytes, nmsg=2. For the Exchange benchmark, the turnover is 4X bytes, nmsg=4.

Throughput values are measured in MBps and can be calculated as follows:

```
throughput = nmsg*X/2^{20} * 10^6/time = nmsg*X/1.048576/time,
```

#### where

- time is measured in µsec.
- x is the length of a message, in bytes.

## **Collective Benchmarks**

Collective benchmarks measure MPI collective operations. Each benchmark is run with varying message lengths. The timing is averaged over multiple samples. The basic MPI data type for all messages is MPI\_BYTE for pure data movement functions and MPI\_FLOAT for reductions.

Collective benchmarks show bare timings.

The following table lists the MPI-1 benchmarks in each class:

Single Transfer	Parallel Transfer	Collective
PingPong	Sendrecv	Bcast Multi-Bcast
PingPongSpecificSource	Exchange	Allgather Multi-Allgather
PingPing	Multi-PingPong	Allgatherv Multi-Allgatherv
PingPingSpecificSource	Multi-PingPing	Alltoall Multi-Alltoall
	Multi-Sendrecv	Alltoallv Multi-Alltoallv
	Multi-Exchange	Scatter Multi-Scatter
		Scatterv Multi-Scatterv
		Gather Multi-Gather
		Gatherv Multi-Gatherv
		Reduce Multi-Reduce

	Reduce_scatter Multi-Reduce_scatter
	Allreduce Multi-Allreduce
	Barrier Multi-Barrier

# Single Transfer Benchmarks

The following benchmarks belong to the single transfer class:

- PingPong
- PingPongSpecificSource
- PingPing
- PingPingSpecificSources

See sections below for definitions of these benchmarks.

## PingPong, PingPongSpecificSource

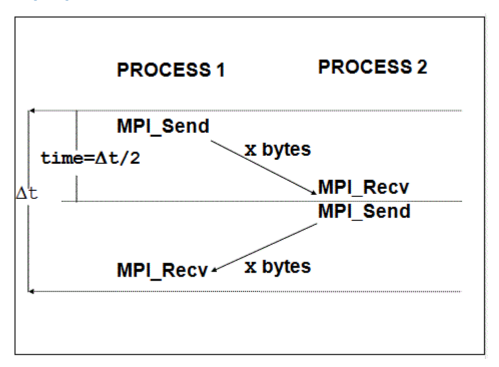
Use PingPong and PingPongSpecificSource for measuring startup and throughput of a single message sent between two processes. PingPong uses the MPI\_ANY\_SOURCE value for destination rank, while PingPongSpecificSource uses an explicit value.

## **PingPong Definition**

Property	Description
Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).
MPI routines	MPI_Send, MPI_Recv
MPI data type	MPI_BYTE
Reported timings	time= $\Delta t/2$ (in $\mu sec$ ) as indicated in the figure below.

Reported throughput	X/(1.048576*time)	

## PingPong Pattern



## PingPing, PingPingSpecificSource

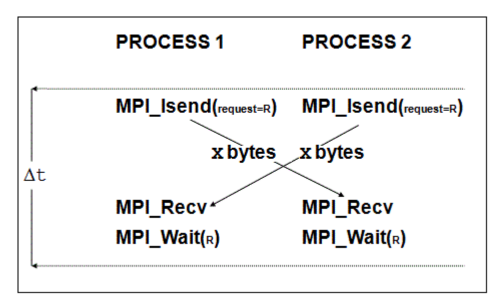
PingPing and PingPingSpecificSource measure startup and throughput of single messages that are obstructed by oncoming messages. To achieve this, two processes communicate with each other using MPI\_Isend/MPI\_Recv/MPI\_Wait calls. The MPI\_Isend calls are issued simultaneously by both processes. For destination rank, PingPing uses the MPI\_ANY\_SOURCE value, while PingPingSpecificSource uses an explicit value.

### **PingPing Definition**

Property	Description
Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).
MPI routines	MPI_Isend/MPI_Wait, MPI_Recv
MPI data type	MPI_BYTE
Reported timings	time=Δt (in μsec)

Reported throughput	X/(1.048576*time)

## **PingPing Pattern**



## Parallel Transfer Benchmarks

The following benchmarks belong to the parallel transfer class:

- Sendrecv
- Exchange
- Multi-PingPong
- Multi-PingPing
- Multi-Sendrecv
- Multi-Exchange

See sections below for definitions of these benchmarks.

#### NOTE:

The definitions of the multiple mode benchmarks are analogous to their standard mode counterparts in the single transfer class.

### **Sendrecy**

The <u>Sendrecv</u> benchmark is based on <u>MPI\_Sendrecv</u>. In this benchmark, the processes form a periodic communication chain. Each process sends a message to the right neighbor and receives

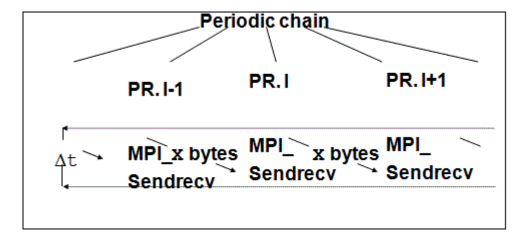
a message from the left neighbor in the chain. The turnover count is two messages per sample (one in, one out) for each process.

In the case of two processes, <u>Sendrecv</u> is equivalent to the <u>PingPing</u> benchmark of <u>IMB1.x</u>. For two processes, it reports the bidirectional bandwidth of the system, as obtained by the optimized <u>MPI\_Sendrecv</u> function.

### **Sendrecy Definition**

Property	Description
Measured pattern	As symbolized between in the figure below.
MPI routines	MPI_Sendrecv
MPI data type	MPI_BYTE
Reported timings	time=∆t (in µsec) as indicated in the figure below.
Reported throughput	2X/(1.048576*time)

#### Sendrecv Pattern



## Exchange

*Exchange* is a communication pattern that often occurs in grid splitting algorithms (boundary exchanges). The group of processes is similar to a periodic chain, and each process exchanges data with both left and right neighbor in the chain.

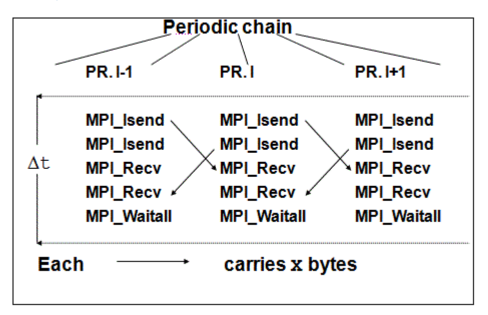
The turnover count is four messages per sample (two in, two out) for each process.

For two Isend messages, separate buffers are used.

## **Exchange Definition**

Property	Description
Measured pattern	As symbolized between in the figure below.
MPI routines	MPI_Isend/MPI_Waitall, MPI_Recv
MPI data type	MPI_BYTE
Reported timings	time=At (in µsec)
Reported throughput	4X/(1.048576*time)

## **Exchange Pattern**



## **Collective Benchmarks**

The following benchmarks belong to the collective class:

- Bcast/multi-Bcast
- Allgather/multi-Allgather
- Allgatherv/multi-Allgatherv
- Alltoall/multi-Alltoall

- Alltoallv/multi-Alltoallv
- Scatter/multi-Scatter
- Scatterv/multi-Scatterv
- Gather/multi-Gather
- Gatherv/multi-Gatherv
- Reduce/multi-Reduce
- Reduce\_scatter/multi-Reduce\_scatter
- Allreduce/multi-Allreduce
- Barrier/multi-Barrier

See sections below for definitions of these benchmarks.

### Reduce

The benchmark for the MPI\_Reduce function. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Reduce
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Root	i%num_procs in iteration i
Reported timings	Bare time
Reported throughput	None

## Reduce\_scatter

The benchmark for the MPI\_Reduce\_scatter function. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM. In the scatter phase, the L items are split as evenly as possible. To be exact, for np number of processes:

```
L = r*np+s
```

where

- r = [L/np]
- $s = L \mod np$

In this case, the process with rank i gets:

- r+1 items when i<s
- r items when i≥s

Property	Description
Measured pattern	MPI_Reduce_scatter
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Reported timings	Bare time
, 3	
Reported throughput	None

## **Allreduce**

The benchmark for the MPI\_Allreduce function. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM.

Property	Description
Measured pattern	MPI_Allreduce
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Reported timings	Bare time
Reported throughput	None

## **Allgather**

The benchmark for the MPI\_Allgather function. Every process inputs X bytes and receives the gathered X\*np bytes, where np is the number of processes.

Property	Description
Measured pattern	MPI_Allgather
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

## **Allgatherv**

The benchmark for the  $\mathtt{MPI\_Allgatherv}$  function. Every process inputs  $\mathtt{X}$  bytes and receives the gathered  $\mathtt{X*np}$  bytes, where  $\mathtt{np}$  is the number of processes. Unlike  $\mathtt{Allgather}$ , this benchmark shows whether MPI produces overhead.

Property	Description
Measured pattern	MPI_Allgatherv
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

## Scatter

The benchmark for the  $\mathtt{MPI\_Scatter}$  function. The root process inputs  $\mathtt{X*np}$  bytes ( $\mathtt{X}$  for each process). All processes receive  $\mathtt{X}$  bytes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Scatter
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	Bare time
Reported throughput	None

## Scatterv

The benchmark for the MPI\_Scatterv function. The root process inputs X\*np bytes (X for each process). All processes receive X bytes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Scatterv
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	Bare time
Reported throughput	None

## Gather

The benchmark for the MPI\_Gather function. The root process inputs X\*np bytes (X from each process). All processes receive X bytes. The root of the operation is changed round-robin.

Description
MPI_Gather
MPI_BYTE
i%num_procs in iteration i
Bare time
None

## **Gatherv**

The benchmark for the MPI\_Gatherv function. All processes input X bytes. The root process receives X\*np bytes, where np is the number of processes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Gatherv

MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	Bare time
Reported throughput	None

## Alltoall

The benchmark for the MPI\_Alltoall function. In the case of np number of processes, every process inputs X\*np bytes (X for each process) and receives X\*np bytes (X from each process).

Property	Description
Measured pattern	MPI_Alltoall
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

## **Bcast**

The benchmark for  $\mathtt{MPI\_Bcast}$ . The root process broadcasts  $\mathtt{X}$  bytes to all other processes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Bcast
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

## Barrier

The benchmark for the MPI\_Barrier function.

Property	Description
Measured pattern	MPI_Barrier
Reported timings	Bare time
Reported throughput	None

# MPI-2 Benchmarks

Intel® MPI Benchmarks provides benchmarks for MPI-2 functions in two components: IMB-EXT and IMB-IO. The table below lists all MPI-2 benchmarks available and specifies whether they support the aggregate mode. For I/O benchmarks, the table also lists nonblocking flavors.

Benchmark	Aggregate Mode	Non-blocking Mode	
IMB-EXT			
Window Multi-Window			
Unidir_Put Multi-Unidir_Put	Supported		
Unidir_Get Multi-Unidir_Get	Supported		
Bidir_Get Multi-Bidir_Get	Supported		
Bidir_Put Multi-Bidir_Put	Supported		
Accumulate Multi-Accumulate	Supported		
Benchmark	Aggregate Mode	Non-blocking Mode	
	IMB-IO		
Open_Close Multi-Open_Close			
S_Write_indv Multi-S_Write_indv	Supported	S_IWrite_indv Multi-S_IWrite_indv	
S_Read_indv Multi-S_Read_indv		S_IRead_indv Multi-S_IRead_indv	
S_Write_expl	Supported	S_IWrite_expl	

Multi-S_Write_expl		Multi-IS_Write_expl
S_Read_expl		S_IRead_expl
Multi-S_Read_expl		Multi-IS_Read_expl
P_Write_indv	Supported	P_IWrite_indv
Multi-P_Write_indv	Supported	Multi-P_IWrite_indv
P_Read_indv		P_IRead_indv
Multi-P_Read_indv		Multi-P_IRead_indv
P_Write_expl	Supported	P_IWrite_expl
Multi-P_Write_expl	Supported	Multi-P_IWrite_expl
P_Read_expl		P_IRead_expl
Multi-P_Read_expl		Multi-P_IRead_expl
P_Write_shared	Supported	P_IWrite_shared
Multi-P_Write_shared	Supported	Multi-P_IWrite_shared
P_Read_shared		P_IRead_shared
Multi-P_Read_shared		Multi-P_IRead_shared
P_Write_priv	Supported	P_IWrite_priv
Multi-P_Write_priv	Supported	Multi-P_IWrite_priv
P_Read_priv		P_IRead_priv
Multi-P_Read_priv		Multi-P_IRead_priv
C_Write_indv	Cupported	C_IWrite_indv
Multi-C_Write_indv	Supported	Multi-C_IWrite_indv
C_Read_indv		C_IRead_indv
Multi-C_Read_indv		Multi-C_IRead_indv
C_Write_expl	Supported	C_IWrite_expl
Multi-C_Write_expl	Supported	Multi-C_IWrite_expl

C_Read_expl		C_IRead_expl
Multi-C_Read_expl		Multi-C_IRead_expl
C_Write_shared Multi-C_Write_shared	Supported	C_IWrite_shared Multi-C_IWrite_shared
C_Read_shared Multi-C_Read_shared		C_IRead_shared Multi-C_IRead_shared

See Also

Benchmark Modes IMB-IO Nonblocking Benchmarks

# Naming Conventions

MPI-2 benchmarks have the following naming conventions:

Convention	Description
Unidir/Bidir	Unidirectional/bidirectional one-sided communications. These are the one-sided equivalents of PingPong and PingPing.
S_	Single transfer benchmark.
c_	Collective benchmark.
P_	Parallel transfer benchmark.
expl	I/O with explicit offset.
indv	I/O with an individual file pointer.
shared	I/O with a shared file pointer.
priv	I/O with an individual file pointer to one private file for each process opened for MPI_COMM_SELF.
[ACTION]	A placeholder for Read or Write component of the benchmark name.

I	Non-blocking flavor. For example, S_IWrite_indv is the nonblocking flavor of the S_IWrite_indv benchmark.
Multi-	The benchmark runs in the multiple mode.

# **IMB-MPI-2 Benchmark Classification**

Intel® MPI Benchmarks introduces three classes of benchmarks:

- Single Transfer
- Parallel Transfer
- Collective

Each class interprets results in a different way.

#### NOTE:

The following benchmarks do not belong to any class:

- Window measures overhead of one-sided communications for the MPI\_Win\_create / MPI\_Win\_free functions
- Open\_close measures overhead of input/output operations for the MPI\_File\_open / MPI\_File\_close functions

#### Single Transfer Benchmarks

This class contains benchmarks of functions that operate on a single data element transferred between one source and one target. For MPI-2 benchmarks, the source of the data transfer can be an MPI process or, in the case of Read benchmarks, an MPI file. The target can be an MPI process or an MPI file.

For I/O benchmarks, the single transfer is defined as an operation between an MPI process and an individual window or a file.

- Single transfer IMB-EXT benchmarks only run with two active processes.
- Single transfer <a>IMB-IO</a> benchmarks only run with one active process.

#### **Parallel Transfer Benchmarks**

This class contains benchmarks of functions that operate on several processes in parallel. The benchmark timings are produced under a global load. The number of participating processes is arbitrary.

In the Parallel Transfer, more than one process participates in the overall pattern.

The final time is measured as the maximum of timings for all single processes. The throughput is related to that time and the overall amount of transferred data (sum over all processes).

#### **Collective Benchmarks**

This class contains benchmarks of functions that are collective as provided by the MPI standard. The final time is measured as the maximum of timings for all single processes. The throughput is not calculated.

#### **MPI-2 Benchmarks Classification**

Single Transfer	Parallel Transfer	Collective	Other
Unidir_Get	Multi_Unidir_Get	Accumulate	Window
Unidir_Put	Multi_Unidir_Put	Multi_Accumulate	Multi_Window
Bidir_Get	Multi_Bidir_Get		
Bidir_Put	Multi_Bidir_Put		
S_[I]Write_indv	P_[I]Write_indv	C_[I]Write_indv	Multi- C_[I]Write_indv
S_[I]Write_indv	P_[I]Write_indv	C_[I]Write_indv Multi- C_[I]Write_indv	Open_close Multi-Open_close
S_[I]Read_indv	P_[I]Read_indv	C_[I]Read_indv  Multi- C_[I]Read_indv	
S_[I]Write_expl	P_[I]Write_expl	C_[I]Write_expl Multi- C_[I]Write_expl	
S_[I]Read_expl	P_[I]Read_expl	C_[I]Read_expl Multi- C_[I]Read_expl	
	P_[I]Write_shared	C_[I]Write_shared Multi- C_[I]Write_shared	

P_[I]Read_shared	C_[I]Read_shared  Multi- C_[I]Write_shared	
P_[I]Write_priv		
P_[I]Read_priv		

# MPI-2 Benchmark Modes

MPI-2 benchmarks can run in the following modes:

- Blocking/nonblocking mode. These modes apply to the IMB-IO benchmarks only. For details, see sections IMB-IO Blocking Benchmarks and IMB-IO Nonblocking Benchmarks.
- Aggregate/non-aggregate mode. Non-aggregate mode is not available for nonblocking flavors of IMB-IO benchmarks.

The following example illustrates aggregation of M transfers for IMB-EXT and blocking Write benchmarks:

```
Select a repetition count M

time = MPI Wtime();
issue M disjoint transfers
assure completion of all transfers
time = (MPI_Wtime() - time) / M
```

In this example:

- M is a repetition count:
  - M = 1 in the non-aggregate mode
  - M = n\_sample in the aggregate mode. For the exact definition of n\_sample see the <u>Actual Benchmarking</u> section.
- A transfer is issued by the corresponding one-sided communication call (for IMB-EXT) and by an MPI-IO write call (for IMB-IO).
- Disjoint means that multiple transfers (if M>1) are to/from disjoint sections of the window or file. This permits to avoid misleading optimizations when using the same locations for multiple transfers.

The variation of  $\underline{M}$  provides important information about the system and the MPI implementation, crucial for application code optimizations. For example, the following possible internal strategies of an implementation could influence the timing outcome of the above pattern.

- Accumulative strategy. Several successive transfers (up to M in the example above) are accumulated without an immediate completion. At certain stages, the accumulated transfers are completed as a whole. This approach may save time of expensive synchronizations. This strategy is expected to produce better results in the aggregate case as compared to the non-aggregate one.
- Non-accumulative strategy. Every Transfer is completed before the return from the corresponding function. The time of expensive synchronizations is taken into account. This strategy is expected to produce equal results for aggregate and non-aggregate cases.

#### **Assured Completion of Transfers**

Following the MPI standard, assured completion of transfers is the minimum sequence of operations after which all processes of the file communicator have a consistent view after a write.

The aggregate and non-aggregate modes differ in when the assured completion of data transfers takes place:

- after each transfer (non-aggregate mode)
- after a bunch of multiple transfers (aggregate mode)

For Intel® MPI Benchmarks, assured completion means the following:

- For IMB-EXT benchmarks, MPI Win fence
- For IMB-IO Write benchmarks, a triplet
   MPI\_File\_sync/MPI\_Barrier(file\_communicator)/MPI\_File\_sync. This fixes the non-sufficient definition in the Intel® MPI Benchmarks 3.0.

# **IMB-EXT Benchmarks**

This section provides definitions of IMB-EXT benchmarks. The benchmarks can run with varying transfer sizes x, in bytes. The timings are averaged over multiple samples. See the <u>Benchmark Methodology</u> section for details. In the definitions below, a single sample with a fixed transfer size x is used.

The <u>Unidir</u> and <u>Bidir</u> benchmarks are exact equivalents of the message passing <u>PingPong</u> and <u>PingPing</u>, respectively. Their interpretation and output are analogous to their message passing equivalents.

#### Unidir\_Put

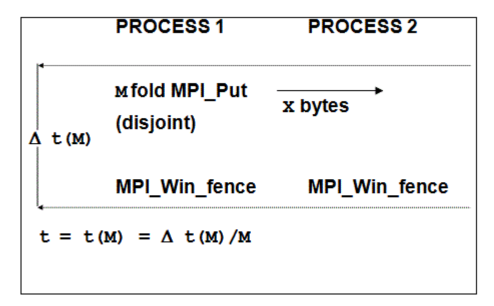
This is the benchmark for the MPI\_Put function. The following table and figure provide the basic definitions and a schematic view of the pattern.

#### Unidir\_Put Definition

Property	Description

Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).
MPI routine	MPI_Put
MPI data type	MPI_BYTE (origin and target)
Reported timings	t=t(M) (in µsec) as indicated in the figure below, non-aggregate (M=1) and aggregate (M=n_sample). For details, see Actual Benchmarking.
Reported throughput	X/t, aggregate and non-aggregate

# Unidir\_Put Pattern



# Unidir\_Get

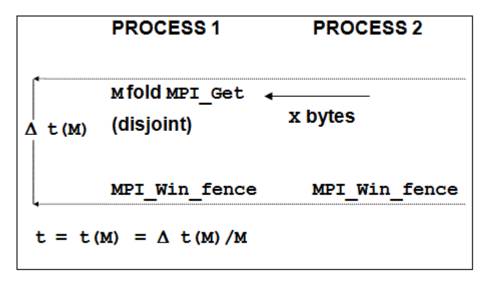
This is the benchmark for the MPI\_Get

# Unidir\_Get Definition

Property	Description
Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).

MPI routine	MPI_Get
MPI data type	MPI_BYTE, for both origin and target
Reported timings	t=t(M) (in µsec) as indicated in the figure below, non-aggregate (M=1)and aggregate (M=n_sample). For details, see Actual Benchmarking.
Reported throughput	X/t, aggregate and non-aggregate

#### Unidir\_Get Pattern



# Bidir\_Put

This is the benchmark for the MPI\_Put function with bidirectional transfers. See the basic definitions below.

#### Bidir\_Put Definition

Property	Description
Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).
MPI routine	MPI_Put
MPI data type	MPI_BYTE, for both origin and target

Reported timings	t=t(M) (in µsec)as indicated in the figure below, non-aggregate (M=1)and aggregate (M=n_sample). For details, see Actual Benchmarking.
Reported throughput	X/t, aggregate and non-aggregate

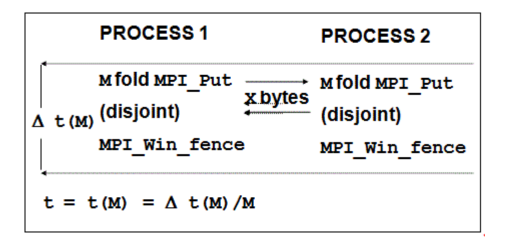
# Bidir\_Get

This is the benchmark for the  $\mathtt{MPI\_Get}$  function, with bidirectional transfers. Below see the basic definitions and a schematic view of the pattern.

# **Bidir\_Get Definition**

Property	Description
Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).
MPI routine	MPI_Get
MPI data type	MPI_BYTE, for both origin and target
Reported timings	t=t(M) (in µsec) as indicated in the figure below, non-aggregate (M=1)and aggregate (M=n_sample). For details, see Actual Benchmarking.
Reported throughput	X/t, aggregate and non-aggregate

# Bidir\_Get Pattern



#### Accumulate

This is the benchmark for the MPI\_Accumulate function. It reduces a vector of length L = x/sizeof(float) of float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM. See the basic definitions and a schematic view of the pattern below.

#### **Accumulate Definition**

Property	Description
Measured pattern	As symbolized between in the figure below. This benchmark runs on two active processes (Q=2).
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Root	О
Reported timings	t=t(M) (in µsec) as indicated in the figure below, non-aggregate (M=1)and aggregate (M=n_sample). For details, see Actual Benchmarking.
Reported throughput	None

#### Accumulate Pattern

```
all active processes

Mfold MPI_Accumulate (x bytes → rank 0)
(disjoint)

Δ t(M)

MPI_Win_fence

t = t(M) = Δ t(M)/M
```

#### Window

This is the benchmark for measuring the overhead of an <a href="MPI\_Win\_create/MPI\_Win\_free">MPI\_Win\_free</a> combination. In the case of an unused window, a negligible non-trivial action is performed inside the window. It minimizes optimization effects of the MPI implementation.

The MPI\_Win\_fence function is called to properly initialize an access epoch. This is a correction as compared to earlier releases of the Intel® MPI Benchmarks.

See the basic definitions and a schematic view of the pattern below.

#### **Window Definition**

Property	Description
Measured pattern	MPI_Win_create/MPI_Win_fence/MPI_Win_free
Reported timings	$t=\Delta t$ (M) (in $\mu sec$ ) as indicated in the figure below.
Reported throughput	None

#### Window Pattern

```
MPI_Win_create (size = X)

Λ t MPI_Win_fence

MPI_Put (1 byte → Window)

MPI_Win_free
```

# **IMB-IO Blocking Benchmarks**

This section describes blocking I/O benchmarks. The benchmarks can run with varying transfer sizes X, in bytes. The timings are averaged over multiple samples. The basic MPI data type for all data buffers is MPI\_BYTE. In the definitions below, a single sample with a fixed I/O size X is used.

Every benchmark contains an elementary I/O action, denoting a pure read or write. Thus, all benchmark flavors have a Write and a Read component. The [ACTION] placeholder denotes a Read or a Write alternatively.

The Write flavors of benchmarks include a file synchronization with different placements for aggregate and non-aggregate modes.

Figure: I/O Benchmarks, Aggregation for Output

# Output: M fold aggregation

м fold elementary I/O action (output), disjoint file sections

**∆** t(M)

# non-aggregate mode:

$$t = \Delta t(M = 1)$$

## aggregate mode:

$$t = \Delta t(M = n \text{ sample}) / M$$

$$(choice of M = n sample)$$

# Input: No aggregation

 $t = \Delta t$  single elementary I/O action (input)

# S\_[ACTION]\_indv

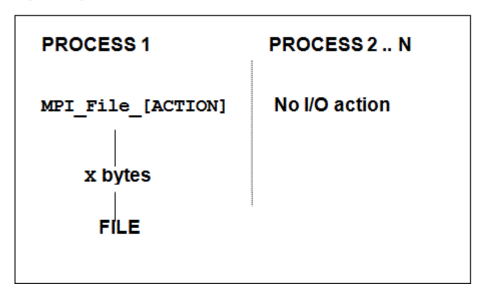
File I/O performed by a single process. This pattern mimics the typical case when a particular master process performs all of the I/O. See the basic definitions and a schematic view of the pattern below.

#### S\_[ACTION]\_indv Definition

Property	Description
Measured pattern	As symbolized in figure I/O benchmarks, aggregation for output
Elementary I/O action	As symbolized in the figure below.
MPI routines for the blocking mode	MPI_File_write/MPI_File_read

MPI routines for the nonblocking mode	MPI_File_iwrite/MPI_File_iread
etype	MPI_BYTE
File type	MPI_BYTE
MPI data type	MPI_BYTE
Reported timings	t (in µsec) as indicated in the figure I/O benchmarks, aggregation for output, aggregate and non-aggregate for the Write flavor.
Reported throughput	x/t, aggregate and non-aggregate for the Write flavor

## S\_[ACTION]\_indv Pattern



# S\_[ACTION]\_expl

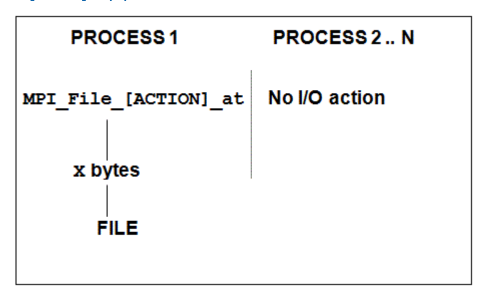
This benchmark mimics the same situation as S\_[ACTION]\_indv, with a different strategy to access files. See the basic definitions and a schematic view of the pattern below.

# S\_[ACTION]\_expl Definition

Property	Description
Measured pattern	As symbolized in figure I/O benchmarks, aggregation for output

Elementary I/O action	As symbolized in the figure below.
MPI routines for the blocking mode	MPI_File_write_at/MPI_File_read_at
MPI routines for the nonblocking mode	MPI_File_iwrite_at/MPI_File_iread_at
etype	MPI_BYTE
File type	MPI_BYTE
MPI data type	MPI_BYTE
Reported timings	t (in µsec) as indicated in the figure <a href="I/O">I/O</a> <a href="Denchmarks">benchmarks</a> , aggregation for output, <a href="aggregate">aggregate</a> and non-aggregate for the Write flavor.
Reported throughput	x/t, aggregate and non-aggregate for the Write flavor

#### S\_[ACTION]\_expl pattern



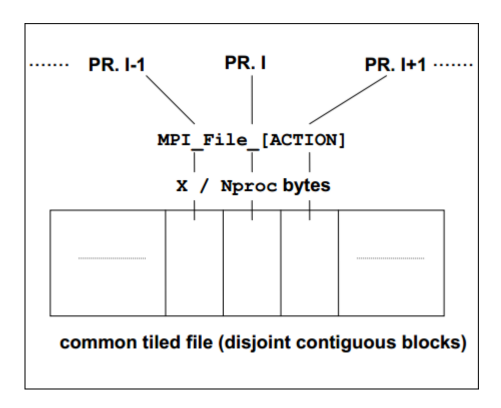
# P\_[ACTION]\_indv

This pattern accesses the file in a concurrent manner. All participating processes access a common file. See the basic definitions and a schematic view of the pattern below.

## P\_[ACTION]\_indv Definition

Property	Description
Measured pattern	As symbolized in figure <u>I/O benchmarks</u> , aggregation for output
Elementary I/O action	As symbolized in the figure below. In this figure, Nproc is the number of processes.
MPI routines for the blocking mode	MPI_File_write/MPI_File_read
MPI routines for the nonblocking mode	MPI_File_iwrite/MPI_File_iread
etype	MPI_BYTE
File type	Tiled view, disjoint contiguous blocks
MPI data type	MPI_BYTE
Reported timings	t (in µsec) as indicated in the figure <a href="I/O">I/O</a> <a href="Denchmarks">benchmarks</a> , aggregation for output, <a href="aggregate">aggregate</a> and non-aggregate for the Write flavor.
Reported throughput	x/t, aggregate and non-aggregate for the Write flavor

# P\_[ACTION]\_indv Pattern



# P\_ACTION\_expl

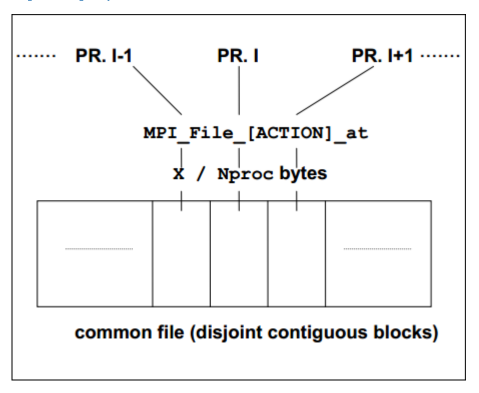
P\_[ACTION]\_expl follows the same access pattern as P\_[ACTION]\_indv with an explicit file pointer type. See the basic definitions and a schematic view of the pattern below.

#### P\_[ACTION]\_expl Definition

Property	Description
Measured pattern	As symbolized in figure I/O benchmarks, aggregation for output
Elementary I/O action	As symbolized in the figure below. In this figure, Nproc is the number of processes.
MPI routines for the blocking mode	MPI_File_write_at/MPI_File_read_at
MPI routines for the nonblocking mode	MPI_File_iwrite_at/MPI_File_iread_at
etype	MPI_BYTE
File type	MPI_BYTE
MPI data type	MPI_BYTE

Reported timings	t (in µsec) as indicated in the figure <a href="I/O">I/O</a> <a href="Denchmarks">benchmarks</a> , aggregation for output, <a href="aggregate">aggregate</a> and non-aggregate for the Write flavor.
Reported throughput	x/t, aggregate and non-aggregate for the Write flavor

#### P\_[ACTION]\_expl Pattern



# P\_[ACTION]\_shared

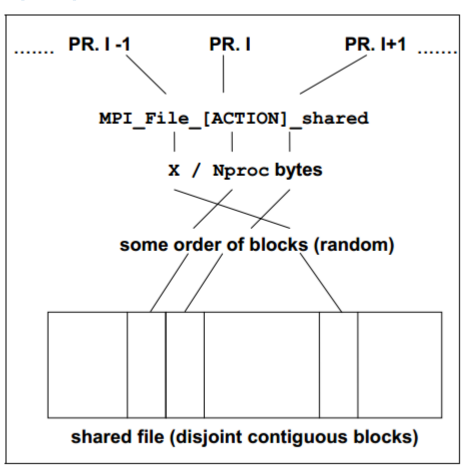
Concurrent access to a common file by all participating processes, with a shared file pointer. See the basic definitions and a schematic view of the pattern below.

#### P\_[ACTION]\_shared Definition

Property	Description
Measured pattern	As symbolized in figure I/O benchmarks, aggregation for output
Elementary I/O action	As symbolized in the figure below. In this figure, Nproc is the number of processes.

MPI routines for the blocking mode	MPI_File_write_at/MPI_File_read_at
MPI routines for the nonblocking mode	MPI_File_iwrite_at/MPI_File_iread_at
etype	MPI_BYTE
File type	MPI_BYTE
MPI data type	MPI BYTE
I data type	_
Reported timings	t (in µsec) as indicated in the figure I/O benchmarks, aggregation for output, aggregate and non-aggregate for the Write flavor.
Reported throughput	x/t, aggregate and non-aggregate for the Write flavor

# P\_[ACTION]\_shared Pattern



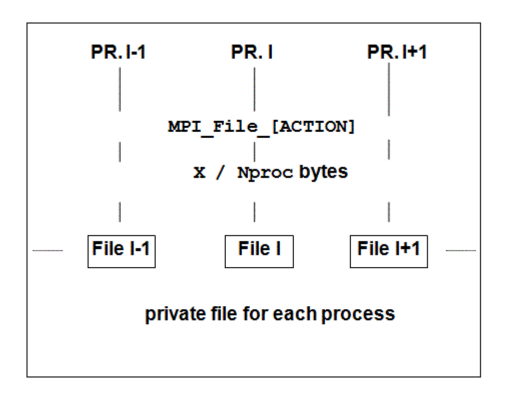
## P\_[ACTION]\_priv

This pattern tests the case when all participating processes perform concurrent I/O to different private files. This benchmark is particularly useful for the systems that allow completely independent I/O operations from different processes. The benchmark pattern is expected to show parallel scaling and obtain optimum results. See the basic definitions and a schematic view of the pattern below.

#### P\_[ACTION]\_priv Definition

Property	Description
Measured pattern	As symbolized in figure I/O benchmarks, aggregation for output
Elementary I/O action	As symbolized in the figure below. In this figure, Nproc is the number of processes.
MPI routines for the blocking mode	MPI_File_write/MPI_File_read
MPI routines for the nonblocking mode	MPI_File_iwrite/MPI_File_iread
etype	MPI_BYTE
File type	MPI_BYTE
MPI data type	MPI_BYTE
Reported timings	$\Delta t$ (in µsec), aggregate and non-aggregate for the Write flavor.
Reported throughput	x/Δt, aggregate and non-aggregate for the Write flavor

# P\_[ACTION]\_priv Pattern



## C\_[ACTION]\_indv

C\_[ACTION]\_indv tests collective access from all processes to a common file, with an individual file pointer. Below see the basic definitions and a schematic view of the pattern.

This benchmark is based on the following MPI routines:

- MPI\_File\_read\_all/MPI\_File\_write\_all for the blocking mode
- MPI\_File\_..\_all\_begin/MPI\_File\_..\_all\_end for the nonblocking mode

All other parameters and the measuring method are the same as for the  $P_[ACTION]_{indv}$  benchmark.

#### See Also

P\_[ACTION]\_indv

#### C\_[ACTION]\_expl

This pattern performs collective access from all processes to a common file, with an explicit file pointer.

This benchmark is based on the following MPI routines:

- MPI File read at all/MPI File write at all for the blocking mode
- MPI\_File\_..\_at\_all\_begin/MPI\_File\_..\_at\_all\_end for the nonblocking mode

All other parameters and the measuring method are the same as for the  $P_[ACTION]_{expl}$  benchmark.

#### **See Also**

P\_[ACTION]\_expl

#### C\_[ACTION]\_shared

The benchmark of a collective access from all processes to a common file, with a shared file pointer.

This benchmark is based on the following MPI routines:

- MPI\_File\_read\_ordered/MPI\_File\_write\_ordered for the blocking mode
- MPI\_File\_..\_ordered\_begin/MPI\_File\_..\_ordered\_end for the nonblocking mode

All other parameters and the measuring method are the same as for the P\_[ACTION]\_shared benchmark.

#### See Also

P\_[ACTION]\_shared

#### Open\_Close

The benchmark for the MPI\_File\_open/MPI\_File\_close functions. All processes open the same file. To avoid MPI implementation optimizations for an unused file, a negligible non-trivial action is performed with the file. See the basic definitions of the benchmark below.

#### Open\_Close Definition

Property	Description
Measured pattern	MPI_File_open/MPI_File_close
etype	MPI_BYTE
File type	MPI_BYTE
Reported timings	$t=\Delta t$ (in $\mu sec$ ), as indicated in the figure below.
Reported throughput	None

#### Open\_Close Pattern

# all active processes MPI\_File open Δ t MPI\_File\_write (1 byte → File) MPI\_File\_close

# **IMB-IO Non-blocking Benchmarks**

Intel® MPI Benchmarks implements blocking and nonblocking modes of the IMB-IO benchmarks as different benchmark flavors. The Read and Write components of the blocking benchmark name are replaced for nonblocking flavors by IRead and IWrite, respectively.

The definitions of blocking and nonblocking flavors are identical, except for their behavior in regard to:

- Aggregation. The nonblocking versions only run in the non-aggregate mode.
- Synchronism. Only the meaning of an elementary transfer differs from the equivalent blocking benchmark.

Basically, an elementary transfer looks as follows:

```
time = MPI_Wtime()
for ( i=0; i<n_sample; i++ )
{
    Initiate transfer
    Exploit CPU
Wait for the end of transfer
}
time = (MPI_Wtime()-time)/n_sample</pre>
```

The Exploit CPU section in the above example is arbitrary. Intel® MPI Benchmarks exploits CPU as described below.

#### **Exploiting CPU**

Intel® MPI Benchmarks uses the following method to exploit the CPU. A kernel loop is executed repeatedly. The kernel is a fully vectorizable multiplication of a 100x100 matrix with a vector. The function is scalable in the following way:

```
IMB_cpu_exploit(float desired_time, int initialize);
```

The input value of <code>desired\_time</code> determines the time for the function to execute the kernel loop, with a slight variance. At the very beginning, the function is called with <code>initialize=1</code> and an input value for <code>desired\_time</code>. This determines an Mflop/s rate and a timing <code>t\_CPU</code>, as close as possible to <code>desired\_time</code>, obtained by running without any obstruction. During the actual benchmarking, <code>IMB\_cpu\_exploit</code> is called with <code>initialize=0</code>, concurrently with the particular I/O action, and always performs the same type and number of operations as in the initialization step.

#### **Displaying Results**

Three timings are crucial to interpret the behavior of nonblocking I/O, overlapped with CPU exploitation:

- t\_pure is the time for the corresponding pure blocking I/O action, non-overlapping with CPU activity
- t\_CPU is the time the IMB\_cpu\_exploit periods (running concurrently with nonblocking I/O) would use when running dedicated
- t\_ovrl is the time for the analogous nonblocking I/O action, concurrent with CPU activity (exploiting t\_CPU when running dedicated)

```
A perfect overlap means: t_ovrl = max(t_pure,t_CPU)

No overlap means: t_ovrl = t_pure+t_CPU.
```

The actual amount of overlap is:

```
overlap=(t_pure+t_CPU-t_ovrl)/min(t_pure,t_CPU)(*)
```

The Intel® MPI Benchmarks result tables report the timings t\_ovrl, t\_pure, t\_CPU and the estimated overlap obtained by the (\*) formula above. At the beginning of a run, the Mflop/s rate is corresponding to the t\_CPU displayed.

# MPI-3 Benchmarks

Intel® MPI Benchmarks provides two sets of benchmarks conforming to the MPI-3 standard:

- <u>IMB-NBC</u> benchmarks for nonblocking collective (NBC) operations
- <u>IMB-RMA</u> one-sided communications benchmarks that measure the Remote Memory Access (RMA) functionality introduced in the MPI-3 standard.

#### See Also

IMB-NBC Benchmarks
IMB-RMA Benchmarks

# **IMB-NBC** Benchmarks

Intel® MPI Benchmarks provides two types of benchmarks for nonblocking collective (NBC) routines that conform to the MPI-3 standard:

- benchmarks for measuring the overlap of communication and computation
- benchmarks for measuring pure communication time

#### TIP

When you run the IMB-NBC component, only the overlap benchmarks are enabled by default. To measure pure communication time, specify the particular benchmark name or use the - include command-line parameter to run the \_pure flavor of the benchmarks.

The following table lists all IMB-NBC benchmarks:

Benchmarks Measuring Communication and Computation Overlap (Enabled by Default)	Benchmarks Measuring Pure Communication Time (Disabled by Default)
Ibcast	Ibcast_pure
Iallgather	
Iallgatherv	
Igather	Igather_pure
Igatherv	Igatherv_pure
Iscatter	Iscatter_pure
Iscatterv	Iscatterv_pure
Ialltoall	Ialltoall_pure
Ialltoallv	Ialltoallv_pure
Ireduce	Ireduce_pure

Ireduce_scatter	Ireduce_scatter_pure
Iallreduce	Iallreduce_pure
Ibarrier	Ibarrier_pure

#### See Also

<u>Measuring Communication and Computation Overlap</u> <u>Measuring Pure Communication Time</u>

#### Measuring Communication and Computation Overlap

Semantics of nonblocking collective operations enables you to run inter-process communication in the background while performing computations. However, the actual overlap depends on the particular MPI library implementation. You can measure a potential overlap of communication and computation using IMB-NBC benchmarks. The general benchmark flow is as follows:

- 1. Measure the time needed for a pure communication call.
- 2. Start a nonblocking collective operation.
- 3. Start computation using the IMB\_cpu\_exploit function, as described in the IMB-IO Nonblocking Benchmarks chapter. To ensure correct measurement conditions, the computation time used by the benchmark is close to the pure communication time measured at step 1.
- 4. Wait for communication to finish using the MPI\_Wait function.

#### Displaying Results

The timing values to interpret the overlap potential are as follows:

- t\_pure is the time of a pure communication operation, non-overlapping with CPU activity.
- t\_CPU is the time the IMB\_cpu\_exploit function takes to complete when run concurrently with the nonblocking communication operation.
- t\_ovrl is the time of the nonblocking communication operation takes to complete when run concurrently with a CPU activity.
  - If t\_ovrl = max(t\_pure,t\_CPU), the processes are running with a perfect overlap.
  - If t\_ovrl = t\_pure+t\_CPU, the processes are running with no overlap.

Since different processes in a collective operation may have different execution times, the timing values are taken for the process with the biggest  $t_{ovrl}$  execution time. The IMB-NBC result tables report the timings  $t_{ovrl}$ ,  $t_{pure}$ ,  $t_{CPU}$  and the estimated overlap in percent calculated by the following formula:

```
overlap = 100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))
```

#### See Also

<u>IMB-NBC Benchmarks</u> <u>Measuring Pure Communication Time</u>

#### **Measuring Pure Communication Time**

To measure pure execution time of nonblocking collective operations, use the <u>pure</u> flavor of the <u>IMB-NBC</u> benchmarks. The benchmark methodology is consistent with the one used for regular <u>collective operations</u>:

- Each rank performs the predefined amount of iterations and calculates the mean value.
- The basic MPI data type for all messages is MPI\_BYTE for pure data movement functions and MPI\_FLOAT for reductions.
- If the operation requires the root process to be specified, the root process is selected round-robin through iterations.

These benchmarks are not included into the default list of IMB-NBC benchmarks. To run a benchmark, specify the particular benchmark name or use the -include command-line parameter. For example:

```
mpirun -np 2 IMB-NBC Ialltoall_pure
mpirun -np 2 IMB-NBC -include Iallgather_pure Ialltoall_pure
```

#### Displaying Results

Pure nonblocking collective benchmarks show bare timing values. Since execution time may vary for different ranks, three timing values are shown: maximum, minimum, and the average time among all the ranks participating in the benchmark measurements.

#### See Also

IMB-NBC Benchmarks
Measuring Communication and Computation Overlap
Command-Line Control

#### lallgather

The benchmark for MPI\_Iallgather that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Iallgather/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see <a href="Measuring Communication and Computation Overlap">Measuring Communication and Computation Overlap</a>.</li> </ul>
Reported throughput	None

#### lallgather\_pure

The benchmark for the MPI\_Iallgather function that measures pure communication time. Every process inputs X bytes and receives the gathered X\*np bytes, where np is the number of processes.

Property	Description
Measured pattern	MPI_Iallgather/MPI_Wait
·	
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

## **lallgatherv**

The benchmark for MPI\_Iallgatherv that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Iallgatherv/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>
Reported throughput	None

# lallgatherv\_pure

The benchmark for the MPI\_Iallgatherv function that measures pure communication time. Every process inputs X bytes and receives the gathered X\*np bytes, where np is the number of processes. Unlike Iallgather\_pure, this benchmark shows whether MPI produces overhead.

Property	Description
Measured pattern	MPI_Iallgatherv/MPI_Wait

MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

#### lallreduce

The benchmark for MPI\_Iallreduce that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Iallreduce/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>
Reported throughput	None

# lallreduce\_pure

The benchmark for the MPI\_Iallreduce function that measures pure communication time. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM.

Property	Description
Measured pattern	MPI_Iallreduce/MPI_Wait
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Reported timings	Bare time

Reported throughput	None

#### lalltoall

The benchmark for MPI\_Ialltoall that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Ialltoall/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>
Reported throughput	None

# lalltoall\_pure

The benchmark for the MPI\_Ialltoall function that measures pure communication time. In the case of np number of processes, every process inputs X\*np bytes (X for each process) and receives X\*np bytes (X from each process).

Property	Description
. ,	·
Measured pattern	MPI_Ialltoall/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

#### **lalltoally**

The benchmark for  $\mathtt{MPI\_Ialltoallv}$  that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Ialltoallv/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>
Reported throughput	None

# lalltoallv\_pure

The benchmark for the  $MPI_Ialltoallv$  function that measures pure communication time. In the case of np number of processes, every process inputs X\*np bytes (X for each process) and receives X\*np bytes (X from each process).

Property	Description
Measured pattern	MPI_Ialltoallv/MPI_Wait
·	
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

#### **Ibarrier**

The benchmark for  ${\tt MPI\_Ibarrier}$  that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Ibarrier/IMB_cpu_exploit/MPI_Wait
Reported timings	<ul><li>t_ovrl</li><li>t_pure</li><li>t_CPU</li></ul>

	<pre>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</pre>
	For details, see Measuring Communication and Computation Overlap.
Reported throughput	None

# lbarrier\_pure

The benchmark for the  ${\tt MPI\_Ibarrier}$  function that measures pure communication time.

Property	Description
Measured pattern	MPI_Ibarrier/MPI_Wait
Reported timings	Bare time
Reported throughput	None

#### **Ibcast**

The benchmark for MPI\_Ibcast that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Ibcast/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>
Reported throughput	None

# lbcast\_pure

The benchmark for  $\mathtt{MPI\_Ibcast}$  that measures pure communication time. The root process broadcasts  $\mathtt{X}$  bytes to all other processes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Ibcast/MPI_Wait
MPI data type	MPI_BYTE
Reported timings	Bare time
Reported throughput	None

# **Igather**

The benchmark for MPI\_Igather that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Igather/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>
Reported throughput	None

# lgather\_pure

The benchmark for the  $\mathtt{MPI\_Igather}$  function that measures pure communication time. The root process inputs  $\mathtt{X*np}$  bytes ( $\mathtt{X}$  from each process). All processes receive  $\mathtt{X}$  bytes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Igather/MPI_Wait
MPI data type	MPI_BYTE

Root	i%num_procs in iteration i
Reported timings	Bare time
Reported throughput	None

# **Igatherv**

The benchmark for  ${\tt MPI\_Igatherv}$  that measures communication and computation overlap.

Property	Description
Measured pattern	MPI_Igatherv/IMB_cpu_exploit/MPI_Wait
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see <a href="Measuring Communication and Computation Overlap">Measuring Communication and Computation Overlap</a>.</li> </ul>
Reported throughput	None

# Igatherv\_pure

The benchmark for the  $\mathtt{MPI\_Igatherv}$  function that measures pure communication time. All processes input  $\mathtt{X}$  bytes. The root process receives  $\mathtt{X*np}$  bytes, where  $\mathtt{np}$  is the number of processes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Igatherv/MPI_Wait
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i

Reported timings	Bare time
Reported throughput	None

### **Ireduce**

The benchmark for  ${\tt MPI\_Ireduce}$  that measures communication and computation overlap.

Property	Description	
Measured pattern	MPI_Ireduce/IMB_cpu_exploit/MPI_Wait	
MPI data type	MPI_FLOAT	
MPI operation	MPI_SUM	
Root	i%num_procs in iteration i	
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>	
Reported throughput	None	

# Ireduce\_pure

The benchmark for the MPI\_Ireduce function that measures pure communication time. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Ireduce/MPI_Wait
MPI data type	MPI_FLOAT
MPI operation	MPI_SUM
Root	i%num_procs in iteration i

Reported timings	Bare time
Reported throughput	None

## lreduce\_scatter

The benchmark for MPI\_Ireduce\_scatter that measures communication and computation overlap. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is MPI\_FLOAT. The MPI operation is MPI\_SUM. In the scatter phase, the L items are split as evenly as possible. To be exact, for np number of processes:

```
L = r*np+s
```

#### where

- r = [L/np]
- s = L mod np

In this case, the process with rank i gets:

- r+1 items when i<s
- r items when i≥s

Property	Description	
Measured pattern	MPI_Ireduce_scatter/IMB_cpu_exploit/MPI_Wait	
MPI data type	MPI_FLOAT	
MPI operation	MPI_SUM	
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see <a href="Measuring Communication and Computation Overlap">Measuring Communication and Computation Overlap</a>.</li> </ul>	
Reported throughput	None	

## Ireduce\_scatter\_pure

The benchmark for the MPI\_Ireduce\_scatter function that measures pure communication time. It reduces a vector of length L = X/sizeof(float) float items. The MPI data type is

MPI\_FLOAT. The MPI operation is MPI\_SUM. In the scatter phase, the  $\bf L$  items are split as evenly as possible. To be exact, for np number of processes:

L = r\*np+s

where

- r = [L/np]
- s = L mod np

In this case, the process with rank i gets:

- r+1 items when i<s
- r items when i≥s

Property	Description	
Measured pattern	MPI_Ireduce_scatter/MPI_Wait	
MPI data type	MPI FLOAT	
wii i data type		
MPI operation	MPI_SUM	
Reported timings	Bare time	
Reported throughput	None	

### **Iscatter**

The benchmark for MPI\_Iscatter that measures communication and computation overlap.

Property	Description	
Measured pattern	MPI_Iscatter/IMB_cpu_exploit/MPI_Wait	
MPI data type	MPI_BYTE	
Root	i%num_procs in iteration i	
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>	

Reported throughput	None

### lscatter\_pure

The benchmark for the MPI\_Iscatter function that measures pure communication time. The root process inputs X\*np bytes (X for each process). All processes receive X bytes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Iscatter/MPI_Wait
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	Bare time
Reported tirrings	Date time
Reported throughput	None

## **Iscatterv**

The benchmark for  ${\tt MPI\_Iscatterv}$  that measures communication and computation overlap.

Property	Description	
Measured pattern	MPI_Iscatterv/IMB_cpu_exploit/MPI_Wait	
MPI data type	MPI_BYTE	
Root	i%num_procs in iteration i	
Reported timings	<ul> <li>t_ovrl</li> <li>t_pure</li> <li>t_CPU</li> <li>overlap=100.*max(0,min(1, (t_pure+t_CPU-t_ovrl) / min(t_pure, t_CPU))</li> <li>For details, see Measuring Communication and Computation Overlap.</li> </ul>	
Reported throughput	None	

#### Iscatterv\_pure

The benchmark for the  $\mathtt{MPI\_Iscatterv}$  function that measures pure communication time. The root process inputs  $\mathtt{X*np}$  bytes ( $\mathtt{X}$  for each process). All processes receive  $\mathtt{X}$  bytes. The root of the operation is changed round-robin.

Property	Description
Measured pattern	MPI_Iscatterv/MPI_Wait
MPI data type	MPI_BYTE
Root	i%num_procs in iteration i
Reported timings	Bare time
Reported throughput	None

# **IMB-RMA Benchmarks**

Intel® MPI Benchmarks provides a set of remote memory access (RMA) benchmarks that use the passive target communication mode to measure one-sided operations compliant with the MPI-3 standard.

#### **IMB-RMA Benchmark Modes**

When running the IMB-RMA benchmarks, you can choose between the following modes:

- Standard (default) or multiple mode. You can enable the multiple mode for all IMB-RMA benchmarks using the -multi command-line parameter. For details, see <u>Running</u> Benchmarks in Multiple Mode.
- Aggregate or non-aggregate mode. For details on these modes, see the MPI-2
  Benchmark Modes chapter. Some IMB-RMA benchmarks support the non-aggregate mode only. To determine which benchmarks can run in the aggregate mode, see Classification of IMB-RMA Benchmarks.

#### Classification of IMB-RMA Benchmarks

All the IMB-RMA benchmarks fall into the following categories:

#### Single Transfer

In these benchmarks, one process accesses the memory of another process, in unidirectional or bidirectional manner. Single Transfer IMB-RMA benchmarks only run on two active processes. Throughput values are measured in MBps and can be calculated as follows:

```
throughput = X/2^{20} * 10^{6}/time = X/1.048576/time,
```

#### where

- time is measured in µsec.
- x is the length of a message, in bytes.

#### **Multiple Transfer**

In these benchmarks, one process accesses the memory of several other processes.

Throughput values are measured in MBps and can be calculated as follows:

```
throughput = (X/2^{20} * 10^6/time) * N = X/1.048576/time * N, where
```

- time is measured in µsec.
- x is the length of a message, in bytes.
- N is the number of target processes.

#### NOTE

The final throughput value is multiplied by the amount of target processes since the transfer is performed to every process except the origin process itself.

#### **Parallel Transfer**

This class contains benchmarks that operate on several processes in parallel. These benchmarks show bare timing values: maximum, minimum, and the average time among all the ranks participating in the benchmark measurements.

The table below lists all IMB-RMA benchmarks and specifies their properties:

Benchmark	Туре	Aggregated Mode
Unidir_put	Single Transfer	Supported
Unidir_get	Single Transfer	Supported
Bidir_put	Single Transfer	Supported
Bidir_get	Single Transfer	Supported
One_put_all	Multiple Transfer	N/A
One_get_all	Multiple Transfer	N/A
All_put_all	Parallel Transfer	N/A
All_get_all	Parallel Transfer	N/A
Put_local	Single Transfer	Supported
Put_all_local	Multiple Transfer	N/A
Exchange_put	Parallel Transfer	N/A

Exchange_get	Parallel Transfer	N/A	
Accumulate	Single Transfer	Supported	
Get_accumulate	Single Transfer	Supported	
Fetch_and_op	Single Transfer	Supported	
Compare_and_swap	Single Transfer	Supported	
Truly_passive_put	Single Transfer*	N/A	
Get_local	Single Transfer	Supported	
Get_all_local	Multiple Transfer	N/A	

<sup>\*</sup> The output format differs from the regular Single Transfer output. For details, see <a href="mailto:Truly\_passive\_put">Truly\_passive\_put</a>.

# Accumulate

This benchmark measures the MPI\_Accumulate operation in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call.

Property	Description
Measured pattern	MPI_Accumulate/MPI_Win_flush
MPI data type	MPI_FLOAT (origin and target)
MPI operation	MPI_SUM
Reported timings	Bare time
Reported throughput	MBps

## All\_get\_all

The benchmark tests the scenario when all processes communicate with each other using the MPI\_Get operation. To avoid congestion due to simultaneous access to the memory of a process by all other processes, different ranks choose different targets at each particular step. For example, while looping through all the possible target ranks, the next target is chosen as follows: (target\_rank + current\_rank)\*num\_ranks.

Property	Description
Measured pattern	(N*MPI_Get)/MPI_Win_flush_all, where N is the number of target processes
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	None

## All\_put\_all

The benchmark tests the scenario when all processes communicate with each other using MPI\_Put operation. To avoid congestion due to simultaneous access to the memory of a process by all other processes, different ranks choose different targets at each particular step. For example, while looping through all the possible target ranks, the next target is chosen as follows: (target\_rank + current\_rank)\*num\_ranks.

Property	Description
Measured pattern	(N*MPI_Put)/MPI_Win_flush_all, where N is the number of target processes
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	None

## Bidir\_get

This benchmark measures the bidirectional MPI\_Get operation in passive target communication mode. The benchmark runs on two active processes. These processes initiate an access epoch to each other using the MPI\_Lock function, get data from the target, close the access epoch, and call the MPI\_Barrier function.

Property	Description
Measured pattern	MPI_Get/MPI_Win_flush
MPI data type	MPI_BYTE (origin and target)

Reported timings	Bare time
Reported throughput	MBps

# Bidir\_put

This benchmark measures the bidirectional MPI\_Put operation in passive target communication mode. The benchmark runs on two active processes. These processes initiate an access epoch to each other using the MPI\_Lock function, transfer data, close the access epoch, and call the MPI\_Barrier function.

Property	Description
Measured pattern	MPI_Put/MPI_Win_flush
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

# Compare\_and\_swap

This benchmark measures the MPI\_Compare\_and\_swap operation in passive target communication mode. The target process is waiting in the MPI\_Barrier call.

Property	Description
Measured pattern	MPI_Compare_and_swap/MPI_Win_flush
MPI data type	MPI_INT (origin and target)
Reported timings	Bare time
Reported throughput	MBps

# Exchange\_Get

This benchmark tests the scenario when each process exchanges data with its left and right neighbor processes using the MPI\_Get operation.

Property	Description

Measured pattern	(2*MPI_Get)/(2*MPI_Win_flush)
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	None

# Exchange\_Put

This benchmark tests the scenario when each process exchanges data with its left and right neighbor processes using the MPI\_Put operation.

Property	Description
Measured pattern	(2*MPI_Put)/(2*MPI_Win_flush)
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	None

# Fetch\_and\_op

This benchmark measures the MPI\_Fetch\_and\_op operation in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call.

Property	Description
Measured pattern	MPI_Fetch_and_op/MPI_Win_flush
MPI data type	MPI_FLOAT (origin and target)
MPI operation	MPI_SUM
Reported timings	Bare time

Domonto di the considerant	MDrs. or
Reported throughput	MBps

#### Get\_accumulate

This benchmark measures the MPI\_Get\_Accumulate operation in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call.

Property	Description
Measured pattern	MPI_Get_Accumulate/MPI_Win_flush
MPI data type	MPI_FLOAT (origin and target)
MPI operation	MPI_SUM
Reported timings	Bare time
Reported throughput	MBps

## Get\_all\_local

This benchmark tests the MPI\_Get operation where one active process obtains data from all other processes. All target processes are waiting in the MPI\_Barrier call, while the active process performs the transfers. The completion of the origin process is ensured by the MPI\_Win\_flush\_local\_all operation. Since local completion of the MPI\_Get operation is semantically equivalent to a regular completion, the benchmark flow is very similar to the One\_get\_all benchmark.

#### NOTE

This benchmark is not enabled in IMB-RMA by default. Specify the benchmark name in the command line or use the -include command-line parameter to run this benchmark.

Property	Description
Measured pattern	(N*MPI_Get)/MPI_Win_flush_local_all, where N is the number of target processes
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time

		1
Reported throughput	MBps	

### Get\_local

This benchmark measures the combination of MPI\_Get and MPI\_Win\_flush\_all operations in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call. Since local completion of the MPI\_Get operation at the origin side is semantically equivalent to a regular completion, performance results are expected to be very close to the Unidir\_Get benchmark results.

#### NOTE

This benchmark is not enabled in IMB-RMA by default. Specify the benchmark name in the command line or use the -include command-line parameter to run this benchmark.

Property	Description
Measured pattern	MPI_Get/MPI_Win_flush_local
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

### One\_put\_all

This benchmark tests the MPI\_Put operation using one active process that transfers data to all other processes. All target processes are waiting in the MPI\_Barrier call while the origin process performs the transfers.

Property	Description
Measured pattern	(N*MPI_Put)/MPI_Win_flush_all, where N is the number of target processes
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

## One\_get\_all

This benchmark tests the MPI\_Get operation using one active process that gets data from all other processes. All target processes are waiting in the MPI\_Barrier call while the origin process accesses their memory.

Property	Description
Measured pattern	(N*MPI_Get)/MPI_Win_flush_all , where N is the number of target processes.
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

# Put\_all\_local

This benchmark tests the MPI\_Put operation where one active process transfers data to all other processes. All target processes are waiting in the MPI\_Barrier call, while the origin process performs the transfers. The completion of the origin process is ensured by the MPI\_Win\_flush\_local\_all operation.

Property	Description
Measured pattern	(N*MPI_Put)/MPI_Win_flush_local_all, where N is the number of target processes
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

## Put\_local

This benchmark measures the combination of MPI\_Put and MPI\_Win\_flush\_all operations in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call.

Property	Description
Measured pattern	MPI_Put/MPI_Win_flush_local

MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

# Truly\_passive\_put

This benchmark verifies whether the MPI implementation supports the truly one-sided communication mode. In this mode, the origin process can complete its access epoch even if the target process is outside the MPI stack.

The Truly\_passive\_put benchmark returns two timing values:

- The time needed for the origin process to complete the MPI\_Put operation while the target process is waiting in the MPI stack in the MPI\_Barrier call.
- The time needed for the origin process to complete the MPI\_Put operation while the target process performs computations outside the MPI stack before the MPI\_Barrier call.

To ensure measurement correctness, the time spent by the target process in the computation function should be comparable to the time needed for successful completion of the MPI\_Put operation by the origin process.

Property	Description
Measured pattern	MPI_Put/MPI_Win_flush, while the target process performs computations before the MPI_Barrier call
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	None

## Unidir\_get

This benchmark measures the MPI\_Get operation in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call.

Property	Description

Measured pattern	MPI_Get/MPI_Win_flush
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

# Unidir\_put

This benchmark measures the MPI\_Put operation in passive target communication mode. The benchmark runs on two active processes. The target process is waiting in the MPI\_Barrier call.

Property	Description
Measured pattern	MPI_Put/MPI_Win_flush
MPI data type	MPI_BYTE (origin and target)
Reported timings	Bare time
Reported throughput	MBps

# Benchmark Methodology

This section describes:

- Different ways to manage Intel® MPI Benchmarks control flow
- Command-line syntax for running the benchmarks
- Sample output data you can receive

# **Control Flow**

Intel® MPI Benchmarks provides different ways to manage its control flow:

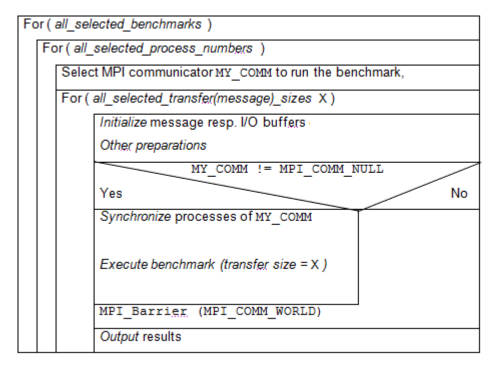
- Hard-coded control mechanisms. For example, the selection of process numbers for running the central benchmarks. See the <u>Hard-coded Settings</u> section for details.
- Preprocessor parameters. The required control parameters are either command-line arguments or parameter selections in the settings.h / setting\_io.h include files. See Parameters Controlling Intel® MPI Benchmarks for details.

Intel® MPI Benchmarks also offers different modes of control:

- Standard mode. In this mode, all configurable sizes are predefined and should not be changed. This ensures comparability for result tables.
- Optional mode. In this mode, you can set these parameters at your choice. You can use this mode to extend the result tables to larger transfer sizes.

The following graph shows the control flow inside the Intel® MPI Benchmarks.

Control flow of Intel® MPI Benchmarks



# **Command-line Control**

You can control all the aspects of the Intel® MPI Benchmarks through the command-line. The general command-line syntax is the following:

```
[-h{elp}]
IMB-MPI1
            [-npmin
                        <NPmin>]
                        <MultiMode>]
            [-multi
            [-off_cache <cache_size[,cache_line_size]>
            [-iter
     <msgspersample[,overall_vol[,msgs_nonaggr[,iter_policy]]]>]
            [-iter_policy
                              <iter_policy>]
            [-time
                       <max_runtime per sample>]
            [-mem
                        <max. mem usage per process>]
            [-msglen
                        <Lengths_file>]
            [-map
                        <PxQ>]
            [-input
                        <filename>]
                        [benchmark1 [,benchmark2 [,...]]]
            [-include]
            [-exclude] [benchmark1 [,benchmark2 [,...]]]
            [-msglog [<minlog>:]<maxlog>]
            [benchmark1 [,benchmark2 [,...]]]
```

The command line is repeated in the output. The options may appear in any order.

#### Examples:

Use the following command line to get out-of-cache data for PingPong:

```
mpirun -np 2 IMB-MPI1 pingpong -off_cache -1
```

Use the following command line to run a very large configuration: restrict iterations to 20, max. 1.5 seconds run time per message size, max. 2 GBytes for message buffers:

```
mpirun -np 512 IMB-MPI1 -npmin 512
    alltoallv -iter 20 -time 1.5 -mem 2
```

Other examples:

### **Benchmark Selection Arguments**

Benchmark selection arguments are a sequence of blank-separated strings. Each argument is the name of a benchmark in exact spelling, case insensitive.

For example, the string IMB-MPI1 PingPong Allreduce specifies that you want to run PingPong and Allreduce benchmarks only.

Default: no benchmark selection. All benchmarks of the selected component are run.

#### -npmin Option

Specifies the minimum number of processes  $P_{\min}$  to run all selected benchmarks on. The  $P_{\min}$  value after -npmin must be an integer.

Given P\_min, the benchmarks run on the processes with the numbers selected as follows:

```
P_min, 2P_min, 4P_min, ..., largest 2*P_min <P, P
```

#### NOTE:

You may set  $P_{\min}$  to 1. If you set  $P_{\min} > P$ , Intel MPI Benchmarks interprets this value as  $P_{\min} = P$ .

Default: no -npmin selection. Active processes are selected as described in the Running Intel® MPI Benchmarks section.

## -multi outflag Option

Defines whether the benchmark runs in the multiple mode. The argument after -multi is a meta-symbol <outflag> that can take an integer value of 0 or 1. This flag controls the way of displaying results:

- Outflag = 0 only display maximum timings (minimum throughputs) over all active groups
- Outflag = 1 report on all groups separately. The report may be long in this case.

When the number of processes running the benchmark is more than half of the overall number MPI\_COMM\_WORLD, the multiple benchmark coincides with the non-multiple one, as not more than one process group can be created.

Default: no -multi selection. Intel® MPI Benchmarks run non-multiple benchmark flavors.

#### -off\_cache cache\_size[,cache\_line\_size] Option

Use the <code>-off\_cache</code> flag to avoid cache re-usage. If you do not use this flag (default), the communications buffer is the same within all repetitions of one message size sample. In this case, <code>Intel® MPI</code> Benchmarks reuses the cache, so throughput results might be non-realistic.

The argument after off\_cache can be a single number (cache\_size), two comma-separated numbers (cache\_size,cache\_line\_size), or -1:

- cache\_size is a float for an upper bound of the size of the last level cache, in MB.
- cache\_line\_size is assumed to be the size of a last level cache line (can be an upper estimate).
- -1 indicates that the default values from IMB\_mem\_info.h should be used. The cache\_size and cache\_line\_size values are assumed to be statically defined in IMB\_mem\_info.h.

The sent/received data is stored in buffers of size ~2x MAX(cache\_size, message\_size). When repetitively using messages of a particular size, their addresses are advanced within those buffers so that a single message is at least 2 cache lines after the end of the previous message. When these buffers are filled up, they are reused from the beginning.

-off\_cache is effective for IMB-MPI1 and IMB-EXT. You are not recommended to use this option for IMB-IO.

#### Examples

Use the default values defined in IMB\_mem\_info.h:

```
-off_cache -1
```

2.5 MB last level cache, default line size:

```
-off_cache 2.5
```

16 MB last level cache, line size 128:

```
-off_cache 16,128
```

The off\_cache mode might also be influenced by eventual internal caching with the Intel® MPI Library. This could make results interpretation complicated.

Default: no cache control. Data may come out of cache.

#### -iter Option

Use this option to control the number of iterations.

By default, the number of iterations is controlled through parameters MSGSPERSAMPLE, OVERALL VOL, MSGS NONAGGR, and ITER POLICY defined in IMB settings.h.

You can optionally add one or more arguments after the -iter flag, to override the default values defined in IMB\_settings.h. Use the following guidelines for the optional arguments:

- To override the MSGSPERSAMPLE value, use a single integer.
- To override the OVERALL\_VOL use two comma-separated integers. The first integer defines the MSGSPERSAMPLE value. The second integer overrides the OVERALL\_VOL value.
- To override the MSGS\_NONAGGR value, use three comma-separated integer numbers. The first integer defines the MSGSPERSAMPLE value. The second integer overrides the OVERALL\_VOL value. The third overrides the MSGS\_NONAGGR value.
- To override the -iter\_policy argument, enter it after the integer arguments, or right after the -iter flag if you do not use any other arguments.

#### Examples

To define MSGSPERSAMPLE as 2000, and OVERALL\_VOL as 100, use the following command line:

```
-iter 2000,100
```

To define MSGS\_NONAGGR as 150, you need to define values for MSGSPERSAMPLE and OVERALL\_VOL as shown in the following command line:

```
-iter 1000,40,150
```

To define MSGSPERSAMPLE as 2000 and set the multiple\_np policy, use the following command line (see -iter\_policy):

```
-iter 2000, multiple_np
```

### -iter\_policy Option

Use this option to set a policy for automatic calculation of the number of iterations. Use one of the following arguments to override the default ITER\_POLICY value defined in IMB\_settings.h:

Policy	Description
dynamic	Reduces the number of iterations when the maximum run time per sample (see - time) is expected to be reached. Using this policy ensures faster execution, but may lead to inaccuracy of the results
multiple_np	Reduces the number of iterations when the message size is growing. Using this policy ensures the accuracy of the results, but may lead to longer execution time. You can control the execution time through the -time option

auto	Automatically chooses which policy to use:  • applies multiple_np to collective operations where one of the ranks acts as the root of the operation (for example, MPI_Bcast)  • applies dynamic to all other types of operations
off	The number of iterations does not change during the execution

You can also set the policy through the -iter option. See -iter.

Default: ITER\_POLICY value defined in IMB\_settings.h. The default policy is dynamic.

### -time Option

Specifies the number of seconds for the benchmark to run per message size. The argument after -time is a floating-point number.

The combination of this flag with the -iter flag or its default alternative ensures that the Intel® MPI Benchmarks always chooses the maximum number of repetitions that conform to all restrictions.

A rough number of repetitions per sample to fulfill the -time request is estimated in preparatory runs that use ~1 second overhead.

Default: -time is activated. The floating-point value specifying the run-time seconds per sample is set in the SECS\_PER\_SAMPLE variable defined in IMB\_settings.h/IMB\_settings\_io.h. The current value is 10.

### -mem Option

Specifies the number of GB to be allocated per process for the message buffers benchmarks/message. If the size is exceeded, a warning is returned, stating how much memory is required for the overall run not to be interrupted.

The argument after <code>-mem</code> is a floating-point number.

Default: the memory is restricted by MAX\_MEM\_USAGE defined in IMB\_mem\_info.h.

### -input <File> Option

Use the ASCII input file to select the benchmarks. For example, the IMB\_SELECT\_EXT file looks as follows:

```
#
# IMB benchmark selection file
#
```

```
# Every line must be a comment (beginning with #), or it
# must contain exactly one IMB benchmark name
#
#Window
Unidir_Get
#Unidir_Put
#Bidir_Get
#Bidir_Put
Accumulate
```

With the help of this file, the following command runs only <code>Unidir\_Get</code> and <code>Accumulate</code> benchmarks of the <code>IMB-EXT</code> component:

```
mpirun .... IMB-EXT -input IMB_SELECT_EXT
```

## -msglen <File> Option

Enter any set of non-negative message lengths to an ASCII file, line by line, and call the Intel\$ MPI Benchmarks with arguments:

```
-msglen Lengths
```

The Lengths value overrides the default message lengths. For IMB-IO, the file defines the I/O portion lengths.

### -map PxQ Option

Numbers processes along rows of the matrix:

0	P	 (Q-2)P	(Q-1)P
1			
P-1	2P-1	(Q-1)P-1	QP-1

For example, to run Multi-PingPongbetween two nodes of size P, with each process on one node communicating with its counterpart on the other, call:

```
mpirun -np <2P> IMB-MPI1 -map <P>x2 PingPong
```

## -include [[benchmark1] benchmark2 ...]

Specifies the list of additional benchmarks to run. For example, to add PingPongSpecificSource and PingPingSpecificSource benchmarks, call:

mpirun -np 2 IMB-MPI1 -include PingPongSpecificSource PingPingSpecificSource

### -exclude [[benchmark1] benchmark2 ...]

Specifies the list of benchmarks to be exclude from the run. For example, to exclude Alltoall and Allgather, call:

mpirun -np 2 IMB-MPI1 -exclude Alltoall Allgather

## -msglog [<minlog>:]<maxlog>

This option allows you to control the lengths of the transfer messages. This setting overrides the MINMSGLOG and MAXMSGLOG values. The new message sizes are 0, 2^minlog, ..., 2^maxlog.

0.94

1.06

65.14

115.16

For example, try running the following command line:

```
mpirun -np 2 IMB-MPI1 -msglog 3:7 PingPong
```

Intel® MPI Benchmarks selects the lengths 0,8,16,32,64,128, as shown below:

```
#-----
# Benchmarking PingPong
# #processes = 2
#-----
    #bytes #repetitions t[µsec]
                         Mbytes/sec
       0
              1000
                      0.70
                               0.00
              1000
                      0.73
                              10.46
       8
                      0.74
       16
             1000
                              20.65
       32
              1000
                      0.94
                              32.61
```

Alternatively, you can specify only the maxlog value:

1000

1000

#-----

64

128

<sup>#</sup> Benchmarking PingPong

#### # #processes = 2

			#
Mbytes/sec	t[µsec]	#repetitions	#bytes
0.00	0.69	1000	0
1.33	0.72	1000	1
2.69	0.71	1000	2
5.28	0.72	1000	4
10.47	0.73	1000	8

# -thread\_level Option

This option specifies the desired thread level for MPI\_Init\_thread(). See description of MPI\_Init\_thread() for details. The option is available only if the Intel® MPI Benchmarks is built with the USE\_MPI\_INIT\_THREAD macro defined. Possible values for <level> are single, funneled, serialized, and multiple.

# Parameters Controlling Intel® MPI Benchmarks

Parameters controlling the default settings of the Intel® MPI Benchmarks are set by preprocessor definition in files <a href="IMB\_settings.h">IMB\_settings.h</a> (for IMB-MPI1 and IMB-EXT benchmarks) and <a href="IMB\_settings\_io.h">IMB\_settings\_io.h</a> (for IMB-IO benchmarks). Both include files have identical structure, but differ in the predefined parameter values.

To enable the optional mode, define the IMB\_OPTIONAL parameter in the IMB\_settings.h/IMB\_settings\_io.h. After you change the settings in the optional section, you need to recompile the Intel® MPI Benchmarks.

The following table describes the Intel MPI Benchmarks parameters and lists their values for the standard mode.

Parameter	Values in IMB_settings.h	Values in IMB_settings_io.h	Description
USE_MPI_INIT_THREAD	Not set	Not set	Set to initialize Intel® MPI Benchmarks by MPI_Init_thread() instead of MPI_Init()
IMB_OPTIONAL	Not set	Not set	Set to activate optional settings
MINMSGLOG	0	0	The second smallest data transfer size is max(unit, 2 <sup>MINMSGLOG</sup> (the smallest size is always 0),

			where unit=sizeof(float) for reductions, unit=1 for all other cases.  You can override this parameter value using the -msglog flag.
MAXMSGLOG	22	24	The largest message size used is 2 <sup>MAXMSGLOG</sup> You can override this parameter value using the -msglog flag.
ITER_POLICY	imode_dynamic		The policy used for calculating the number of iterations.  You can override this parameter value using the -iter_policy or -iter flag.
MSGSPERSAMPLE	1000	50	The maximum repetition count for all IMB-MPI1 benchmarks. You can override this parameter value using the -iter flag.
MSGS_NONAGGR	100	10	The maximum repetition count for non-aggregate benchmarks (relevant only for IMB-EXT). You can override this parameter value using the -time flag.
OVERALL_VOL	40 Mbytes	16*1048576	For all sizes smaller than OVERALL_VOL, the repetition count is reduced so that not more than OVERALL_VOL bytes are processed all in all. This permits you to avoid unnecessary repetitions for large message sizes. Finally, the real repetition count for message size X is MSGSPERSAMPLE (X=0), min(MSGSPERSAMPLE,

			max(1,OVERALL_VOL/X)) (X>0)  Note that OVERALL_VOL does not restrict the size of the maximum data transfer. 2 <sup>MAXMSGLOG</sup> OVERALL_VOL.  You can override this parameter value using the -mem flag.
SECS_PER_SAMPLE	10		Number of iterations is dynamically set so that this number of run time seconds is not exceeded per message length.
N_BARR	2	2	Number of MPI_Barrier for synchronization.
TARGET_CPU_SECS	0.01 seconds	0.1 seconds	CPU seconds (as float) to run concurrently with nonblocking benchmarks (currently irrelevant for IMB-MPI1)

In the example below, the IMB\_settings\_io.h. file has the IMB\_OPTIONAL parameter enabled, so that user-defined parameters are used. I/O sizes of 32 and 64 MB, and a smaller repetition count are selected, extending the standard mode tables. You can modify the optional values as required.

```
#define FILENAME IMB_out
#define IMB_OPTIONAL
#ifdef IMB_OPTIONAL
#define MINMSGLOG 25
#define MAXMSGLOG 26
#define MSGSPERSAMPLE 10
#define MSGS_NONAGGR 10
#define OVERALL_VOL 16*1048576
#define SECS_PER_SAMPLE 10
#define TARGET_CPU_SECS 0.1 /* unit seconds */
#define N_BARR 2
#else
```

```
/*Do not change anything below this line*/
#define MINMSGLOG 0

#define MAXMSGLOG 24

#define MSGSPERSAMPLE 50

#define MSGS_NONAGGR 10

#define OVERALL_VOL 16*1048576

#define TARGET_CPU_SECS 0.1 /* unit seconds */
#define N_BARR 2

#endif
```

If IMB\_OPTIONAL is deactivated, Intel MPI Benchmarks uses the default standard mode values.

# **Hard-Coded Settings**

The sections below describe Intel® MPI Benchmarks hard-coded settings:

- Communicators, Active Processes
- Other Preparations for Benchmarking
- Message /I-O Buffer Lengths
- Buffer Initialization
- Warm-Up Phase (MPI-1, EXT)
- Synchronization
- Actual Benchmarking

### Communicators, Active Processes

Communicator management is repeated in every "select MY\_COMM" step. If it exists, the previous communicator is freed. When running  $Q \le P$  processes, the first Q ranks of MPI\_COMM\_WORLD are put into one group, and the remaining P = Q get MPI\_COMM\_NULL.

The group of MY\_COMM calls the active processes group.

### Other Preparations for Benchmarking

#### Window (IMB-EXT and IMB-RMA)

- 1. An Info is set and MPI\_Win\_create is called, creating a window of size X for MY\_COMM.
- 2. For IMB-EXT, MPI\_Win\_fence is called to start an access epoch.

#### NOTE

IMB-RMA benchmarks do not require MPI\_Win\_fence since they use the passive target communication mode.

#### File (IMB-IO)

To initialize the IMB-IO file, follow these steps:

- 1. Select a file name. This parameter is located in the IMB\_settings\_io.h include file. In the case of a multi-<MPI command>, a suffix \_g<groupid> is appended to the name. If the file name is per process, a second event suffix \_<rank> is appended.
- 2. Delete the file if it exists: open the file with MPI\_MODE\_DELETE\_ON\_CLOSE and close it.
- 3. Select a communicator to open the file: MPI\_COMM\_SELF for S\_benchmarks and P\_[ACTION]\_priv.
- 4. Select a mode: MPI\_MODE\_CREATE | MPI\_MODE\_RDWR
- 5. Select an info routine as explained below.

#### Info

Intel® MPI Benchmarks uses an external function <code>User\_Set\_Info</code> which you implement for the current system. The default version is:

```
#include mpi.h
void User_Set_Info ( MPI_Info* opt_info)
#ifdef MPIIO

{/* Set info for all MPI_File_open calls */
*opt_info = MPI_INFO_NULL;
}
#endif
#ifdef EXT

{/* Set info for all MPI_Win_create calls */
*opt_info = MPI_INFO_NULL;
}
#endif
#endif
```

The Intel® MPI Benchmarks use no assumptions and imposes no restrictions on how this routine is implemented.

#### View (IMB-IO)

The file view is determined by the following settings:

- disp = 0,
- datarep = native
- etype, filetypeas defined in the benchmark definitions above

• info as defined in the "Info" section above.

# Message/I-O Buffer Lengths

#### IMB-MPI1, IMB-EXT

Set in IMB\_settings.h and used unless the -msglen flag is selected.

#### IMB-IO

Set in IMB\_settings\_io.h and used unless the -msglen flag is selected.

#### **Buffer Initialization**

Communication and I/O buffers are dynamically allocated as void\* and used as MPI\_BYTE
buffers for all benchmarks except Accumulate, see Memory Requirements. To assign the buffer contents, a cast to an assignment type is performed. This facilitates result checking which may become necessary. Besides, a sensible data type is mandatory for Accumulate.

Intel® MPI Benchmarks sets the buffer assignment type assign\_type in IMB\_settings.h/IMB\_settings\_io.h. Currently, int is used for IMB-IO, float for IMB-EXT. The values are set by a CPP macro as follows.

For IMB-EXT benchmarks:

```
#define BUF_VALUE(rank,i) (0.1*((rank)+1)+(float)( i)
For IMB-IO benchmarks:
#define BUF_VALUE(rank,i) 10000000*(1+rank)+i%10000000
```

In every initialization, communication buffers are seen as typed arrays and initialized as follows:

```
((assign_type*)buffer)[i] = BUF_VALUE(rank,i;
```

where rank is the MPI rank of the calling process.

#### Warm-Up Phase (IMB-MPI1, IMB-EXT, IMB-NBC, and IMB-RMA)

Before starting the actual benchmark measurement for IMB-MPI1, IMB-EXT, IMB-NBC, and IMB-RMA, the selected benchmark is executed N\_WARMUP times with a sizeof(assign\_type) message length. The N\_WARMUP value is defined in IMB\_settings.h, see <a href="Parameters Controlling Intel® MPI Benchmarks">Parameters Controlling Intel® MPI Benchmarks</a> for details. The warm-up phase eliminates the initialization overheads from the benchmark measurement.

### **Synchronization**

Before the actual benchmark measurement is performed, the constant  $N_BARR$  is used to regulate calls to:

```
MPI_Barrier(MPI_COMM_WORLD)
```

The N\_BARR constant is defined in IMB\_settings.h and IMB\_settings\_io.h, with the current value of 2.

See figure Control flow of IMB to ensure that all processes are synchronized.

# **Actual Benchmarking**

To reduce measurement errors caused by insufficient clock resolution, every benchmark is run repeatedly. The repetition count is as follows:

For IMB-MPI1, IMB-NBC, and aggregate flavors of IMB-EXT, IMB-IO, and IMB-RMA benchmarks, the repetition count is MSGSPERSAMPLE. This constant is defined in IMB\_settings\_io.h, with 1000 and 50 values, respectively.

To avoid excessive run times for large transfer sizes X, an upper bound is set to OVERALL\_VOL/X. The OVERALL\_VOL value is defined in IMB\_settings.h/IMB\_settings\_io.h, with 4MB and 16MB values, respectively.

Given transfer size X, the repetition count for all aggregate benchmarks is defined as follows:

```
n_sample = MSGSPERSAMPLE (X=0)
n_sample = max(1,min(MSGSPERSAMPLE,OVERALL_VOL/X)) (X>0)
```

The repetition count for non-aggregate benchmarks is defined completely analogously, with MSGSPERSAMPLE replaced by MSGS\_NONAGGR. A reduced count is recommended as non-aggregate run times are usually much longer.

In the following examples, elementary transfer means a pure function (MPI\_[Send, ...], MPI\_Put, MPI\_Get, MPI\_Accumulate, MPI\_File\_write\_XX, MPI\_File\_read\_XX), without any further function call. Assured completion transfer completion is:

- MPI\_Win\_fence for IMB-EXT benchmarks
- a triplet MPI\_File\_sync/MPI\_Barrier(file\_communicator)/MPI\_File\_sync for IMB-IO
   Write benchmarks
- MPI\_Win\_flush, MPI\_Win\_flush\_all, MPI\_Win\_flush\_local, or MPI\_Win\_flush\_local\_all for IMB-RMA benchmarks
- empty for all other benchmarks

#### **MPI-1 Benchmarks**

```
for ( i=0; i<N_BARR; i++ ) MPI_Barrier(MY_COMM)
time = MPI_Wtime()
for ( i=0; i<n_sample; i++ )
    execute MPI pattern
time = (MPI_Wtime()-time)/n_sample</pre>
```

#### IMB-EXT and Blocking I/O Benchmarks

For aggregate benchmarks, the kernel loop looks as follows:

```
for ( i=0; i<N_BARR; i++ )MPI_Barrier(MY_COMM)

/* Negligible integer (offset) calculations ... */
time = MPI_Wtime()

for ( i=0; i<n_sample; i++ )
   execute elementary transfer
   assure completion of all transfers

time = (MPI_Wtime()-time)/n_sample</pre>
```

For non-aggregate benchmarks, every single transfer is safely completed:

```
for ( i=0; i<N_BARR; i++ )MPI_Barrier(MY_COMM)

/* Negligible integer (offset) calculations ... */
time = MPI_Wtime()

for ( i=0; i<n_sample; i++ )

{
   execute elementary transfer
   assure completion of transfer
}

time = (MPI_Wtime()-time)/n_sample</pre>
```

#### Non-blocking I/O Benchmarks

A nonblocking benchmark has to provide three timings:

- t\_pure blocking pure I/O time
- t\_ovrl- nonblocking I/O time concurrent with CPU activity
- t\_CPU pure CPU activity time

The actual benchmark consists of the following stages:

- Calling the equivalent blocking benchmark as defined in <u>Actual Benchmarking</u> and taking benchmark time as t\_pure.
- Closing and re-opening the particular file(s).
- Re-synchronizing the processes.
- Running the nonblocking case, concurrent with CPU activity (exploiting t\_CPU when running undisturbed), taking the effective time as t\_ovrl.

The desired CPU time to be matched approximately by t\_CPU is set in IMB\_settings\_io.h:
#define TARGET\_CPU\_SECS 0.1 /\* unit seconds \*/

# **Checking Results**

To check whether your MPI implementation is working correctly, you can use the CPP flag - DCHECK.

Activate the CPP flag -DCHECK through the CPPFLAGS variable and recompile the Intel® MPI Benchmarks executable files. Every message passing result from the Intel® MPI Benchmarks are checked against the expected outcome. Output tables contain an additional column called Defects that displays the difference as floating-point numbers.

#### NOTE:

The -DCHECK results are not valid as real benchmark data. Deactivate -DCHECK and recompile to get the proper results.

# **Output**

The benchmark output includes the following information:

- General information:
  - machine, system, release, and version are obtained by IMB\_g\_info.c.
- The calling sequence (command-line flags) are repeated in the output chart
- Results for the non-multiple mode

After a benchmark completes, three time values are available, extended over the group of active processes:

- Tmax the maximum time
- Tmin the minimum time
- Tavg the average time

The time unit is  $\mu$ .

Single Transfer Benchmarks:

```
Display X = message size [bytes], T=Tmax[\musec], bandwidth = X / 1.048576 / T
```

Parallel Transfer Benchmarks:

```
Display X = message; size, Tmax, Tmin and Tavg, bandwidth based on time = Tmax
```

#### NOTE

IMB-RMA benchmarks show only bare timings for Parallel Transfer benchmarks.

Collective Benchmarks:

```
Display X = message size;(except for Barrier), Tmax, Tmin; and Tavg
Results for the multiple mode
```

- -multi 0: the same as above, with min, avg over all groups.
- -multi 1: the same for all groups, max, min, avg over single groups.

### Sample 1 - IMB-MPI1 PingPong Allreduce

# or through the flag => -time

The following example shows the results of the PingPong and Allreduce benchmark:

```
<...> np 2 IMB-MPI1 PingPong Allreduce
#-----
# Intel (R) MPI Benchmark Suite V3.2, MPI1 part
#-----
# Date
                   : Thu Sep 4 13:20:07 2008
# Machine
                    : x86_64
# System
                    : Linux
# Release
                    : 2.6.9-42.ELsmp
                    : #1 SMP Wed Jul 12 23:32:02 EDT 2006
# Version
# MPI Version
                    : 2.0
# MPI Thread Environment: MPI_THREAD_SINGLE
# New default behavior from Version 3.2 on:
 the number of iterations per message size is cut down
# dynamically when a certain run time (per message size sample)
# is expected to be exceeded. Time limit is defined by variable
# SECS_PER_SAMPLE (=> IMB_settings.h)
```

```
# Calling sequence was:
# ./IMB-MPI1 PingPong Allreduce
# Minimum message length in bytes: 0
# Maximum message length in bytes: 4194304
# MPI_Datatype
                       : MPI_BYTE
# MPI_Datatype for reductions : MPI_FLOAT
# MPI_Op
                    : MPI_SUM
#
# List of Benchmarks to run:
# PingPong
# Allreduce
#-----
# Benchmarking PingPong
# #processes = 2
#-----
#bytes #repetitions t[µsec] Mbytes/sec
    0
          1000
                .. ..
    1
          1000
    2
          1000
    4
          1000
          1000
   8
   16
          1000
   32
       1000
   64 1000
  128 1000
```

```
256
             1000
  512
             1000
  1024
             1000
  2048
             1000
  4096
             1000
  8192
             1000
 16384
             1000
 32768
             1000
 65536
              640
131072
              320
262144
           160
524288
              80
1048576
              40
              20
2097152
4194304
              10
#-----
# Benchmarking Allreduce
# ( #processes = 2 )
#-----
\texttt{\#bytes} \quad \texttt{\#repetitions} \quad \texttt{t\_min[} \texttt{\mu} \texttt{sec} \texttt{]} \quad \texttt{t\_avg[} \texttt{\mu} \texttt{sec} \texttt{]}
    0
             1000
    4
             1000
             1000
    8
    16
          1000
    32
             1000
   64
             1000
   128
              1000
   256
             1000
   512
             1000
  1024
             1000
  2048
             1000
  4096
            1000
```

8192	1000
16384	1000
32768	1000
65536	640
131072	320
262144	160
524288	80
1048576	40
2097152	20
4194304	10

# All processes entering MPI\_Finalize

# Sample 2 - IMB-MPI1 PingPing Allreduce

The following example shows the results of the PingPing

```
<..>
-np 6 IMB-MPI1
 pingping allreduce -map 2x3 -msglen Lengths -multi 0
Lengths file:
100
1000
10000
100000
1000000
#-----
# Intel (R) MPI Benchmark Suite V3.2.2, MPI1 part
# Date
                   : Thu Sep 4 13:26:03 2008
# Machine
                  : x86_64
# System
                  : Linux
```

```
# Release
                     : 2.6.9-42.ELsmp
# Version
                    : #1 SMP Wed Jul 12 23:32:02 EDT 2006
# MPI Version
                     : 2.0
# MPI Thread Environment: MPI_THREAD_SINGLE
# New default behavior from Version 3.2 on:
# the number of iterations per message size is cut down
# dynamically when a certain run time (per message size sample)
# is expected to be exceeded. Time limit is defined by variable
# SECS_PER_SAMPLE (=> IMB_settings.h)
# or through the flag => -time
# Calling sequence was:
# IMB-MPI1 pingping allreduce -map 3x2 -msglen Lengths
         -multi 0
# Message lengths were user-defined
# MPI_Datatype
                                 : MPI_BYTE
# MPI_Datatype for reductions : MPI_FLOAT
# MPI_Op
                                   : MPI_SUM
#
# List of Benchmarks to run:
# (Multi-)PingPing
# (Multi-)Allreduce
#-----
# Benchmarking Multi-PingPing
# ( 3 groups of 2 processes each running simultaneously )
# Group 0: 0 3
# Group 1: 1 4
```

```
# Group 2: 2 5
#
#-----
# bytes #rep.s t min[µsec] t max[µsec] t avg[µsec] Mbytes/sec
  0 1000 .. .. .. ..
 100 1000
 1000 1000
 10000 1000
100000 419
1000000 41
#-----
# Benchmarking Multi-Allreduce
# ( 3 groups of 2 processes each running simultaneously )
# Group 0: 0 3
# Group 1: 1 4
# Group 2: 2 5
#-----
#bytes #repetitions t min[µsec] t max[µsec] t avg[µsec]
  0
        1000
              . .
                   . .
 100 1000
 1000 1000
 10000 1000
100000
         419
1000000
         41
#-----
# Benchmarking Allreduce
#processes = 4; rank order (rowwise):
```

```
0 3
# 1 4
# ( 2 additional processes waiting in MPI_Barrier)
#-----
# bytes #repetitions t_min[µsec] t_max[µsec] t_avg[µsec]
  0
         1000
                . .
 100 1000
 1000 1000
 10000 1000
        419
100000
1000000
          41
#-----
# Benchmarking Allreduce
# processes = 6; rank order (rowwise):
# 0 3
  2 5
# bytes #repetitions t_min[µsec] t_max[µsec] t_avg[µsec]
  0
         1000
                .. .. ..
 100
         1000
 1000
         1000
10000
       1000
100000 419
1000000 41
```

```
# All processes entering MPI_Finalize
```

### Sample 3 - IMB-IO p\_write\_indv

The following example shows the results of the p\_write\_indv benchmark:

```
<...> IMB-IO -np 2 p_write_indv -npmin 2
#-----
# Date
                    : Thu Sep 4 13:43:34 2008
# Machine
                    : x86_64
# System
                   : Linux
# Release
                   : 2.6.9-42.ELsmp
                   : #1 SMP Wed Jul 12 23:32:02 EDT 2006
# Version
# MPI Version
                   : 2.0
# MPI Thread Environment: MPI THREAD SINGLE
# New default behavior from Version 3.2 on:
# the number of iterations per message size is cut down
# dynamically when a certain run time (per message size sample)
# is expected to be exceeded. Time limit is defined by variable
# SECS_PER_SAMPLE (=> IMB_settings.h)
# or through the flag => -time
# Calling sequence was:
# ./IMB-IO p_write_indv -npmin 2
# Minimum io portion in bytes: 0
# Maximum io portion in bytes: 16777216
# List of Benchmarks to run:
# P_Write_Indv
#-----
```

```
# Benchmarking P_Write_Indv
# #processes = 2
#-----
 MODE: AGGREGATE
 0
      50
                 .. .. ..
          . .
   1 50
   2 50
   4 50
   8
      50
   16
      50
   32
      50
  64
      50
  128
     50
  256
     50
     50
  512
  1024
      50
  2048
      50
 4096
      50
 8192
      50
 16384 50
 32768
     50
 65536 50
 131072
      50
 262144
      50
524288
      32
     16
1048576
2097152
     8
4194304
     4
     2
8388608
```

```
16777216 1
#-----
# Benchmarking P_Write_Indv
# #processes = 2
#-----
# MODE: NON-AGGREGATE
 #bytes #rep.s t_min[\musec] t_max t_avg Mb/sec
   0 10 .. .. ..
   1 10
   2
      10
   4
      10
   8 10
  16 10
  32 10
  64 10
  128
      10
  256
      10
  512
      10
  1024 10
 2048 10
 4096 10
 8192 10
 16384
      10
 32768
      10
 65536
      10
 131072 10
```

262144 10 524288 10

1048576 10

```
2097152 8
4194304 4
8388608 2
16777216 1
```

# All processes entering MPI\_Finalize

## Sample 4 - IMB-EXT.exe

The example below shows the results for the Window benchmark received after running IMB-EXT.exe on a Microsoft Windows\* cluster using two processes. The performance diagnostics for Unidir\_Get, Unidir\_Put, Bidir\_Get, Bidir\_Put, and Accumulate are omitted.

```
<..> -n 2 IMB-EXT.exe
#----
   Intel (R) MPI Benchmark Suite V3.2.2, MPI-2 part
#-----
# Date
                   : Fri Sep 05 12:26:52 2008
# Machine : Intel64 Family 6 Model 15 Stepping 6, GenuineIntel
                  : Windows Server 2008
# System
# Release
                  : .0.6001
# Version
                   : Service Pack 1
# MPI Version
                : 2.0
# MPI Thread Environment: MPI_THREAD_SINGLE
# New default behavior from Version 3.2 on:
# the number of iterations per message size is cut down
# dynamically when a certain run time (per message size sample)
# is expected to be exceeded. Time limit is defined by variable
# SECS_PER_SAMPLE (=> IMB_settings.h)
# or through the flag => -time
```

```
# Calling sequence was:
# \\master-node\MPI_Share_Area\IMB_3.1\src\IMB-EXT.exe
# Minimum message length in bytes: 0
# Maximum message length in bytes: 4194304
# MPI_Datatype
                        : MPI_BYTE
# MPI_Datatype for reductions
                        : MPI_FLOAT
qO_IqM #
                        : MPI_SUM
# List of Benchmarks to run:
# Window
# Unidir_Get
# Unidir_Put
# Bidir_Get
# Bidir_Put
# Accumulate
#-----
# Benchmarking Window
# #processes = 2
#-----
     #bytes #repetitions t min[µsec] t max[µsec] t avg[µsec]
                 100 .. ..
         4
                 100
         8
                 100
        16
                100
        32
                100
                100
        64
       128
                100
                100
       256
```

100
100
100
100
100
100
100
100
100
100
80
40
20
10

. . .

#### # All processes entering MPI\_Finalize

The above example listing shows the results of running IMB-EXT.exe on a Microsoft Windows\* cluster using two processes.

The listing only shows the result for the Window benchmark. The performance diagnostics for Unidir\_Get, Unidir\_Put, Bidir\_Get, Bidir\_Put, and Accumulate are omitted.