

Intel® MPI Library Developer Reference for Linux* OS

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Intel® MPI Library Developer Reference for Linux* OS

Documentation for older versions of the Intel® MPI Library are available for download only. For a list of available documentation downloads by product version, see Download Documentation for Intel Parallel Studio XE. To download the previous versions of Intel MPI Library documentation, refer to this page.

What's New

This Developer Reference provides you with the complete reference for the Intel MPI Library. It is intended to help a user fully utilize the Intel MPI Library functionality. For examples and detailed functionality description, please refer to the Intel MPI Library Developer Guide.

The following are some popular topics in the Intel MPI Library Developer Reference:

Command Reference

Command Reference provides reference information on compilation and runtime commands (mpirun, cpuinfo, impi_info) and describes how to use these commands.

Environment Variable Reference

Environment Variable Reference provides syntax, arguments, and descriptions for Fabrics Control, Tuning, Autotuning, Process Pinning, and I_MPI_ADJUST Family environment variables.

Global Options and Environment Variables for mpiexec.hydra

Describes the Global Options and provides Environment Variables used with the Hydra process manager.

Mpitune Configuration Options

Tune your MPI application with the mpitune utility.

Introduction

This Developer Reference provides you with the complete reference for the Intel® MPI Library. It is intended to help an experienced user fully utilize the Intel MPI Library functionality. You can freely redistribute this document in any desired form.

Document Organization

Section	Description
Section 1.Introduction	Introduces this document and the Intel MPI Library.
Section 2.Command Reference	Describes compilation and job startup commands and their options.
Section 3. Environment Variable Reference	Describes environment variables .
Section 4.Miscellaneous	Contains information not related to the sections above.

Introducing Intel® MPI Library

Intel® MPI Library is a multi-fabric message passing library that implements the Message Passing Interface, v3.1 (MPI-3.1) specification. It provides a standard library across Intel® platforms that enable adoption of MPI-3.1 functions as their needs dictate.

Intel® MPI Library enables developers to change or to upgrade processors and interconnects as new technology becomes available without changes to the software or to the operating environment.

You can get the latest information for the Intel® MPI Library at https://software.intel.com/intel-mpi-library.

What's New

This page lists changes to the product that are reflected in the documentation. For a list of all changes, refer to the Release Notes.

Intel® oneAPI 2021.4

• Changed the I MPI STARTUP MODE default value to pmi shm netmod.

Intel® oneAPI 2021.3

- Changed the default pinning order to bunch.
- Added new value for the I_MPI_SHM variable (icx).

Additionally, minor updates have been made to fix inaccuracies in the document and improve user experience.

Intel® oneAPI 2021.2

No documentation changes.

Intel® oneAPI Gold

- Removed all content specific to Intel® Parallel Studio XE (see notice on title page).
- Added Intel[®] Ethernet 800 Series support.
- Added MPI + OpenMP offload examples.
- Added new algorithm for MPI_Sendrecv_replace (I_MPI_ADJUST_SENDRECV_REPLACE=2).
- Added I MPI OFFLOAD variable to GPU Support topic.
- Reworked directory layout:

- Removed intel64/.
- Mpivars.[c]sh and mpi modulefile moved to env/.
- Mpivars.[c]sh renamed to vars.[c]sh.
- Removed deprecated symbolic links.
- Removed static libraries for debug configurations.

Notational Conventions

The following conventions are used in this document.

This type style	Document names
This type style	Hyperlinks
This type style	Commands, arguments, options, file names
THIS TYPE STYLE	Environment variables
<this style="" type=""></this>	Variables or placeholders for actual values
[items]	Optional items
{ item item }	Selectable items separated by vertical bar(s)

Related Information

Description of some of the Intel® MPI Library functionality is available in man1 pages: mpiexec.hydra, hydra_nameserver, and compiler wrappers.

The following related documents that might be useful to the user:

- Product Web Site
- Intel® MPI Library Support
- Intel® Cluster Tools Products
- Intel® Software Development Products

Command Reference



This section provides information on different command types and how to use these commands:

- Compilation Commands lists the available Intel® MPI Library compiler commands, related options, and environment variables.
- mpirun provides the description and examples for the mpirun command.
- mpiexec.hydra gives full information on the mpiexec.hydra command, its options, environment variables, and related features and utilities.
- cpuinfo provides the syntax, arguments, description and output examples for the cpuinfo utility.
- impi_info provides information on available environment variables.
- mpitune provides information on configuration options for the mpitune utility.

Compiler Commands

The following table lists the available Intel® MPI Library compiler commands with their underlying compilers and programming languages.

Intel MPI Library Compiler Wrappers

Compiler Command	Default Compiler	Supported Languages
Generic Compilers		
mpicc	gcc, cc	С
mpicxx	g++	C/C++
mpifc	gfortran	Fortran77*/Fortran 95*
GNU* Compilers		
mpigcc	gcc	С
mpigxx	g++	C/C++
mpif77	gfortran	Fortran 77
mpif90	gfortran	Fortran 95
Intel® Fortran, C++ Compile	rs	
mpiicc	icc	С
mpiicpc	ісрс	C++
mpiifort	ifort	Fortran77/Fortran 95

Notes on Compiler Commands

- Compiler commands are available only in the Intel MPI Library Software Development Kit (SDK).
- For the supported versions of the listed compilers, refer to the *Release Notes*.
- To display mini-help of a compiler command, execute it without any parameters.
- Compiler wrapper scripts are located in the <install-dir>/bin directory, where <install-dir> is the Intel MPI Library installation directory.
- The environment settings can be established by sourcing the <install-dir>/env/vars.[c]sh script. To use a specific library configuration, pass one of the following arguments to the script to switch to the corresponding configuration: release, debug, release mt, or debug mt.
- Ensure that the corresponding underlying compiler is already in your PATH. If you use Intel® compilers, source the vars.sh script from the installation directory to set up the compiler environment.

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Compilation Command Options

-nostrip

Use this option to turn off the debug information stripping while linking the Intel® MPI Library statically.

-config=<name>

Use this option to source a compiler configuration file. The file should contain the environment settings to be used with the specified compiler.

Use the following naming convention for configuration files:

```
<install-dir>/etc/mpi<compiler>-<name>.conf
```

where:

- <compiler>={cc, cxx, f77, f90}, depending on the language compiled.
- <name> is the name of the underlying compiler with spaces replaced by hyphens; for example, the <name> value for cc -64 is cc--64.

-profile=<profile_name>

Use this option to specify an MPI profiling library. $<profile_name>$ is the name of the configuration file (profile) that loads the corresponding profiling library. The profiles are taken from <install-dir>/etc.

The Intel MPI Library comes with several predefined profiles for the Intel® Trace Collector:

- <install-dir>/etc/vt.conf regular tracing library
- <install-dir>/etc/vtfs.conf fail-safe tracing library
- <install-dir>/etc/vtmc.conf correctness checking tracing library
- <install-dir>/etc/vtim.conf load imbalance tracing library

You can also create your own profile as conf. You can define the following environment variables in a configuration file:

- PROFILE PRELIB libraries (and paths) to load before the Intel MPI Library
- PROFILE POSTLIB libraries to load after the Intel MPI Library
- PROFILE INCPATHS C preprocessor arguments for any include files

For example, create a file myprof.conf with the following lines:

```
PROFILE_PRELIB="-L<path_to_myprof>/lib -lmyprof"
PROFILE_INCPATHS="-I<paths_to_myprof>/include"
```

Use the -profile=myprof option for the relevant compiler wrapper to select this new profile.

-t or -trace

Use the -t or -trace option to link the resulting executable file against the Intel® Trace Collector library. Using this option has the same effect as the -profile=vt option.

You can also use the <code>I_MPI_TRACE_PROFILE</code> environment variable to <code><profile_name></code> to specify another profiling library. For example, set <code>I_MPI_TRACE_PROFILE</code> to <code>vtfs</code> to link against the fail-safe version of the Intel Trace Collector.

To use this option, include the installation path of the Intel® Trace Collector in the VT_ROOT environment variable. Source the vars.[c]sh script provided in the Intel® Trace Analyzer and Collector installation folder.

-trace-imbalance

Use the -trace-imbalance option to link the resulting executable file against the load imbalance tracing library of Intel Trace Collector. Using this option has the same effect as the -profile=vtim option.

To use this option, include the installation path of the Intel Trace Collector in the VT_ROOT environment variable. Source the vars.[c]sh script provided in the Intel® Trace Analyzer and Collector installation folder.

-check mpi

Use this option to link the resulting executable file against the Intel® Trace Collector correctness checking library. The default value is libVTmc.so. Using this option has the same effect as the -profile=vtmc option.

You can also use the <code>I_MPI_CHECK_PROFILE</code> environment variable to <code><profile_name></code> to specify another checking library.

To use this option, include the installation path of the Intel Trace Collector in the VT_ROOT environment variable. Source the vars.[c]sh script provided in the Intel® Trace Analyzer and Collector installation folder.

-ilp64

Use this option to enable partial ILP64 support. All integer arguments of the Intel MPI Library are treated as 64-bit values in this case.

-no ilp64

Use this option to disable the ILP64 support explicitly. This option must be used in conjunction with -i8 option of Intel® Fortran Compiler.

If you specify the -i8 option for the separate compilation with Intel Fortran Compiler, you still have to use the i8 or i1p64 option for linkage.

-dynamic_log

Use this option in combination with the -t option to link the Intel Trace Collector library dynamically. This option does not affect the default linkage method for other libraries.

To run the resulting programs, include \$VT ROOT/slib in the LD LIBRARY PATH environment variable.

-g

Use this option to compile a program in debug mode and link the resulting executable file against the debugging version of the Intel MPI Library. See I_MPI_DEBUG for information on how to use additional debugging features with the -g builds.

The optimized library is linked with the -g option by default.

Use vars. {sh|csh} [debug|debug mt] at runtime to load a particular libmpi.so configuration.

-link mpi=<arg>

Use this option to always link the specified version of the Intel MPI Library. See the I_MPI_LINK environment variable for detailed argument descriptions. This option overrides all other options that select a specific library .

Use vars. {sh|csh} [debug|debug mt] during runtime to load particular libmpi.so configuration.

-0

Use this option to enable compiler optimization.

ce **1**

-fast

Use this option to maximize speed across the entire program. This option forces static linkage method for the Intel MPI Library.

This option is supported only by the mpiicc, mpiicpc, and mpiifort Intel® compiler wrappers.

-echo

Use this option to display everything that the command script does.

-show

Use this option to learn how the underlying compiler is invoked, without actually running it. Use the following command to see the required compiler flags and options:

```
$ mpiicc -show -c test.c
```

Use the following command to see the required link flags, options, and libraries:

```
$ mpiicc -show -o a.out test.o
```

This option is particularly useful for determining the command line for a complex build procedure that directly uses the underlying compilers.

-show env

Use this option to see the environment settings in effect when the underlying compiler is invoked.

-{cc,cxx,fc}=<compiler>

Use this option to select the underlying compiler. The tables below list the available LLVM and ILO compiler options and commands used to invoke them.

LLVM Compiler Options for Intel® oneAPI

Language	Product Name	Compiler Driver	Compiler Wrapper	Command	Example
С	>Intel® oneAPI DPC++/C++ Compiler	icx	mpiicc	-cc=icx	\$ mpiicc -cc=icx -c test.c
C++	Intel [©] oneAPI DPC++/C++ C	icpx	mpiicpc	-cxx=icpx	\$ mpiicpc - cxx=icpx -c test.cpp
DPC++	Intel [©] oneAPI DPC++/C++ Compiler	dpcpp	mpiicpc	-cxx=dpcpp	\$ mpiicpc - cxx=dpcpp - c test.cpp
Fortran	Intel [®] oneAPI Fortran Compiler	ifx	mpiifort	-fc=ifx	\$ mpiifort -fc=ifx -c test.

NOTE Make sure that the wrapper name is in your PATH. Alternatively, you can specify the full path to the compiler.

-nofortbind, -nofortran

Use this option to disable mpiicc linking with Fortran bindings. This has the same effect as the I MPI FORT BIND variable.

-v

Use this option to print the compiler wrapper script version and its underlying compiler version.

-norpath

Use this option to disable rpath for the compiler wrapper for the Intel® MPI Library.

mpirun

Launches an MPI job and provides integration with job schedulers.

Syntax

mpirun <options>

Arguments

<options></options>	mpiexec.hydra options as described in the	
	<pre>mpiexec.hydra section. This is the default operation</pre>	
	mode.	

Description

Use this command to launch an MPI job. The mpirun command uses Hydra as the underlying process manager.

The mpirun command detects if the MPI job is submitted from within a session allocated using a job scheduler like Torque*, PBS Pro*, LSF*, Parallelnavi* NQS*, Slurm*, Univa* Grid Engine*, or LoadLeveler*. The mpirun command extracts the host list from the respective environment and uses these nodes automatically according to the above scheme.

In this case, you do not need to create a host file. Allocate the session using a job scheduler installed on your system, and use the mpirun command inside this session to run your MPI job.

Example

```
$ mpirun -n <# of processes> ./myprog
```

This command invokes the mpiexec.hydra command (Hydra process manager), which launches the myprog executable.

mpiexec.hydra

Launches an MPI job using the Hydra process manager.

Syntax

```
mpiexec.hydra<g-options> <l-options> <executable>
```

or

mpiexec.hydra<g-options> <1-options> <executable1> : <1-options> <executable2>

Arguments

<g-options></g-options>	Global options that apply to all MPI processes
<i-options></i-options>	Local options that apply to a single argument set
<executable></executable>	./a.out or path/ name of the executable file

Description

Use the mpiexec.hydra utility to run MPI applications using the Hydra process manager.

Command Reference

Use the first short command-line syntax to start all MPI processes of the <executable> with the single set of arguments. For example, the following command executes a.out over the specified processes and hosts:

```
$ mpiexec.hydra -f <hostfile> -n <# of processes> ./a.out
```

where:

- <# of processes> specifies the number of processes on which to run the a.out executable
- <hostfile> specifies a list of hosts on which to run the a.out executable

Use the second long command-line syntax to set different argument sets for different MPI program runs. For example, the following command executes two different binaries with different argument sets:

```
$ mpiexec.hydra -f <hostfile> -env <VAR1> <VAL1> -n 2 ./a.out : \
-env <VAR2> <VAL2> -n 2 ./b.out
```

NOTE You need to distinguish global options from local options. In a command-line syntax, place the local options after the global options.

Global Hydra Options

This section describes the global options of the Intel® MPI Library's Hydra process manager. Global options are applied to all arguments sets in the launch command. Argument sets are separated by a colon ':'.

-tune <filename>

Use this option to specify the file name that contains the tuning data in a binary format.

-usize <usize>

Use this option to set MPI UNIVERSE SIZE, which is available as an attribute of the MPI COMM WORLD.

<size></size>	Define the universe size
SYSTEM	Set the size equal to the number of cores passed to mpiexec through the
	hostfile or the resource manager.
INFINITE	Do not limit the size. This is the default value.
<value></value>	Set the size to a numeric value ≥ 0 .

-hostfile <hostfile> or -f <hostfile>

Use this option to specify host names on which to run the application. If a host name is repeated, this name is used only once.

See also the I MPI HYDRA HOST FILE environment variable for more details.

NOTE Use the following options to change the process placement on the cluster nodes:

- Use the -perhost, -ppn, and -grr options to place consecutive MPI processes on every host using the round robin scheduling.
- Use the -rr option to place consecutive MPI processes on different hosts using the round robin scheduling.

-machinefile <machine file> or -machine <machine file>

Use this option to control process placement through a machine file. To define the total number of processes to start, use the -n option. For example:

```
$ cat ./machinefile
node0:2
node1:2
node0:1
```

-hosts-group

Use this option to set node ranges using brackets, commas, and dashes (like in Slurm* Workload Manager). For more details, see the I_MPI_HYDRA_HOST_FILE environment variable in Hydra Environment Variables.

-silent-abort

Use this option to disable abort warning messages.

For more details, see the I MPI SILENT ABORT environment variable in Hydra Environment Variables.

-nameserver

Use this option to specify the nameserver in the hostname:port format.

For more details, see the I MPI HYDRA NAMESERVER environment variable in Hydra Environment Variables.

-genv <ENVVAR> <value>

Use this option to set the <ENVVAR> environment variable to the specified <value> for all MPI processes.

-genvall

Use this option to enable propagation of all environment variables to all MPI processes.

-genvnone

Use this option to suppress propagation of any environment variables to any MPI processes.

NOTE The option does not work for localhost.

-genvexcl < list of env var names>

Use this option to suppress propagation of the listed environment variables to any MPI processes.

-genvlist < list >

Use this option to pass a list of environment variables with their current values. <1ist> is a comma separated list of environment variables to be sent to all MPI processes.

-pmi-connect <mode>

Use this option to choose the caching mode of process management interface (PMI) message. Possible values for <mode> are:

<mode></mode>	The caching mode to be used
nocache	Do not cache PMI messages.

cache	Cache PMI messages on the local pmi_proxy management processes to minimize the number of PMI requests. Cached information is automatically propagated to child management processes.
lazy-cache	cache mode with on-request propagation of the PMI information.
alltoall	Information is automatically exchanged between all pmi_proxy before any get request can be done. This is the default mode.

See the I MPI HYDRA PMI CONNECT environment variable for more details.

-perhost <# of processes >, -ppn <# of processes >, or -grr <# of processes>

Use this option to place the specified number of consecutive MPI processes on every host in the group using round robin scheduling. See the I MPI PERHOST environment variable for more details.

NOTE When running under a job scheduler, these options are ignored by default. To be able to control process placement with these options, disable the <code>I_MPI_JOB_RESPECT_PROCESS_PLACEMENT</code> variable.

-rr

Use this option to place consecutive MPI processes on different hosts using the round robin scheduling. This option is equivalent to "-perhost 1". See the I MPI PERHOST environment variable for more details.

-trace [<profiling library>] or -t [<profiling library>]

Use this option to profile your MPI application with Intel® Trace Collector using the indicated cprofiling_library>. If you do not specify cprofiling_library>, the default profiling library libVT.so is used.

Set the I MPI JOB TRACE LIBS environment variable to override the default profiling library.

-trace-imbalance

Use this option to profile your MPI application with Intel® Trace Collector using the libVTim.so library.

-aps

Use this option to collect statistics from your MPI application using Application Performance Snapshot. The collected data includes hardware performance metrics, memory consumption data, internal MPI imbalance and OpenMP* imbalance statistics. When you use this option, a new folder <code>aps_result_<date>-<time></code> with statistics data is generated. You can analyze the collected data with the <code>aps</code> utility, for example:

```
$ mpirun -aps -n 2 ./myApp
$ aps aps_result_20171231_235959
```

NOTE

- 1. To use this option, set up the Application Performance Snapshot environment beforehand. See the tool's https://software.intel.com/content/www/us/en/develop/documentation/application-snapshot-user-guide/top.html*User Guide*.
- 2. If you use the options -trace or -check mpi, the -aps option is ignored.

-mps

Use this option to collect only MPI and OpenMP* statistics from your MPI application using Application Performance Snapshot. Unlike the <code>-aps</code> option, <code>-mps</code> doesn't collect hardware metrics. The option is equivalent to:

```
$ mpirun -n 2 aps -c mpi,omp ./myapp
```

-trace-pt2pt

Use this option to collect the information about point-to-point operations using Intel® Trace Analyzer and Collector. The option requires that you also use the -trace option.

-trace-collectives

Use this option to collect the information about collective operations using Intel® Trace Analyzer and Collector. The option requires that you also use the -trace option.

NOTE

Use the -trace-pt2pt and -trace-collectives to reduce the size of the resulting trace file or the number of message checker reports. These options work with both statically and dynamically linked applications.

-configfile <filename>

Use this option to specify the file $\langle filename \rangle$ that contains the command-line options with one executable per line. Blank lines and lines that start with '#' are ignored. Other options specified in the command line are treated as global.

You can specify global options in configuration files loaded by default (mpiexec.conf in <installdir>/etc, ~/.mpiexec.conf, and mpiexec.conf in the working directory). The remaining options can be specified in the command line.

-branch-count < num>

Use this option to restrict the number of child management processes launched by the Hydra process manager, or by each $pmi\ proxy$ management process.

See the I MPI HYDRA BRANCH COUNT environment variable for more details.

-pmi-aggregate or -pmi-noaggregate

Use this option to switch on or off, respectively, the aggregation of the PMI requests. The default value is -pmi-aggregate, which means the aggregation is enabled by default.

See the I MPI HYDRA PMI AGGREGATE environment variable for more details.

-gdb

Use this option to run an executable under GDB* (GNU debugger). You can use the following command:

```
$ mpiexec.hydra -gdb -n <# of processes><executable>
```

-gdba <pid>

Use this option to attach the GNU* debugger to the existing MPI job. You can use the following command:

```
$ mpiexec.hydra -gdba <pid>
```

Command Reference 1

-nolocal

Use this option to avoid running the <executable> on the host where mpiexec.hydra is launched. You can use this option on clusters that deploy a dedicated main node for starting the MPI jobs and a set of dedicated compute nodes for running the actual MPI processes.

-hosts < nodelist>

Use this option to specify a particular <nodelist> on which the MPI processes should be run. For example, the following command runs the executable a.out on the hosts host1 and host2:

```
$ mpiexec.hydra -n 2 -ppn 1 -hosts host1,host2 ./a.out
```

NOTE If <nodelist> contains only one node, this option is interpreted as a local option. See Local Options for details.

-iface <interface>

Use this option to choose the appropriate network interface. For example, if the IP emulation of your InfiniBand* network is configured to ib0, you can use the following command.

```
$ mpiexec.hydra -n 2 -iface ib0 ./a.out
```

See the I MPI HYDRA IFACE environment variable for more details.

-demux <mode>

Use this option to set the polling mode for multiple I/O. The default value is poll.

Arguments

<spec></spec>	Define the polling mode for multiple I/O
poll	Set poll as the polling mode. This is the default value.
select	Set select as the polling mode.

See the I_MPI_HYDRA_DEMUX environment variable for more details.

-enable-x or -disable-x

Use this option to control the Xlib* traffic forwarding. The default value is -disable-x, which means the Xlib traffic is not forwarded.

-l, -prepend-rank

Use this option to insert the MPI process rank at the beginning of all lines written to the standard output.

-ilp64

Use this option to preload the ILP64 interface.

-s <spec>

Use this option to direct standard input to the specified MPI processes.

Arguments

<spec></spec>	Define MPI process ranks
all	Use all processes.

none	Do not direct standard output to any processes.
<1>, <m>,<n></n></m>	Specify an exact list and use processes $<1>$, $$ and $$ only. The default value is zero.
<k>,<1>-<m>,<n></n></m></k>	Specify a range and use processes $< k>$, $<1>$ through $< m>$, and $< n>$.

-noconf

Use this option to disable processing of the mpiexec.hydra configuration files.

-ordered-output

Use this option to avoid intermingling of data outut from the MPI processes. This option affects both the standard output and the standard error streams.

NOTE When using this option, end the last output line of each process with the end-of-line \n' character. Otherwise the application may stop responding.

-path <directory>

Use this option to specify the path to the executable file.

-tmpdir <dir>

Use this option to set a directory for temporary files. See the I_MPI_TMPDIR environment variable for more details.

-version or -V

Use this option to display the version of the Intel® MPI Library.

-info

Use this option to display build information of the Intel® MPI Library. When this option is used, the other command line arguments are ignored.

-localhost

Use this option to explicitly specify the local host name for the launching node.

-rmk <RMK>

Use this option to select a resource management kernel to be used. Intel® MPI Library only supports pbs.

See the I MPI HYDRA RMK environment variable for more details.

-outfile-pattern <file>

Use this option to redirect stdout to the specified file.

-errfile-pattern <file>

Use this option to redirect stderr to the specified file.

-gpath <path>

Use this option to specify the path to the executable file.

-gwdir <dir>

Use this option to specify the working directory in which the executable file runs.

-gumask <umask>

Use this option to perform the "umask <umask>" command for the remote executable file.

-gdb-ia

Use this option to run processes under Intel® architecture specific GNU* debugger.

-prepend-pattern

Use this option to specify the pattern that is prepended to the process output.

-verbose or -v

Use this option to print debug information from mpiexec.hydra, such as:

- Service processes arguments
- Environment variables and arguments passed to start an application
- PMI requests/responses during a job life cycle

See the I MPI HYDRA DEBUG environment variable for more details.

-print-rank-map

Use this option to print out the MPI rank mapping.

-print-all-exitcodes

Use this option to print the exit codes of all processes.

-bootstrap
 bootstrap server>

Use this option to select a built-in bootstrap server to use. A bootstrap server is the basic remote node access mechanism that is provided by the system. Hydra supports multiple runtime bootstrap servers such as ssh, rsh, pdsh, fork, persist,slurm, ll, lsf, or sge to launch MPI processes. The default bootstrap server is ssh. By selecting slurm, ll, lsf, or sge, you use the corresponding srun, llspawn.stdio, blaunch, or qrsh internal job scheduler utility to launch service processes under the respective selected job scheduler (Slurm*, LoadLeveler*, LSF*, and SGE*).

Arguments

<arg></arg>	String parameter
ssh	Use secure shell. This is the default value.
rsh	Use remote shell.
pdsh	Use parallel distributed shell.
pbs	Use Torque* pbsdsh command.
pbsdsh	Alias for pbs bootstrap.
fork	Use fork call.
persist	Use Hydra persist server. See below for details.
slurm	Use Slurm* srun command.
11	Use LoadLeveler* llspawn.stdio command.
lsf	Use LSF blaunch command.
sge	Use Univa* Grid Engine* qrsh command.

See I MPI HYDRA BOOTSTRAP for details.

-bootstrap-exec <bootstrap server>

Use this option to set the executable to be used as a bootstrap server. The default bootstrap server is ssh. For example:

See I MPI HYDRA BOOTSTRAP for more details.

-bootstrap-exec-args <args>

Use this option to provide the additional parameters to the bootstrap server executable file.

```
$ mpiexec.hydra -bootstrap-exec-args <arguments> -n 2 ./a.out
```

For tight integration with the Slurm* scheduler (including support for suspend/resume), use the method outlined on the Slurm page here: http://www.schedmd.com/slurmdocs/mpi_guide.html#intel_mpi

See I MPI HYDRA BOOTSTRAP EXEC EXTRA ARGS for more details.

-v6

Use this option to force using the IPv6 protocol.

Local Hydra Options

This section describes the local options of the Intel® MPI Library's Hydra process manager. Local options are applied only to the argument set they are specified in. Argument sets are separated by a colon ':'.

-n <number-of-processes> Or -np <number-of-processes>

Use this option to set the number of MPI processes to run with the current argument set.

-env <envar> <value>

Use this option to set the $\langle envar \rangle$ environment variable to the specified $\langle value \rangle$ for all MPI processes in the current argument set.

-envall

Use this option to propagate all environment variables in the current argument set. See the I MPI HYDRA ENV environment variable for more details.

-envnone

Use this option to suppress propagation of any environment variables to the MPI processes in the current argument set.

NOTE The option does not work for localhost.

-envexcl <list-of-envvar-names>

Use this option to suppress propagation of the listed environment variables to the MPI processes in the current argument set.

Command Reference

-envlist <list>

Use this option to pass a list of environment variables with their current values. <1ist> is a comma separated list of environment variables to be sent to the MPI processes.

-host <nodename>

Use this option to specify a particular <nodename> on which the MPI processes are to be run. For example, the following command executes a.out on hosts host1 and host2:

```
$ mpiexec.hydra -n 2 -host host1 ./a.out : -n 2 -host host2 ./a.out
```

-path <directory>

Use this option to specify the path to the <executable> file to be run in the current argument set.

-wdir <directory>

Use this option to specify the working directory in which the <executable> file runs in the current argument set.

gtool Options

-gtool

Use this option to launch such tools as Intel® VTune™ Amplifier XE, Intel® Advisor, Valgrind*, and the GDB* (GNU Debugger) for the specified processes through the mpiexec.hydra and mpirun commands. An alternative to this option is the I MPI GTOOL environment variable.

Syntax

-gtool "<command line for tool 1>:<ranks set 1>[=launch mode 1][@arch 1]; <command line for tool 2>:<ranks set 2>[=exclusive][@arch 2]; ...; <command line for a tool n>:<ranks set n>[=exclusive][@arch n]" <executable>

or:

\$ mpirun -n <# of processes> -gtool "<command line for tool 1>:<ranks set 1>[=launch mode 1][@arch 1]" -gtool "<command line for a tool 2>:<ranks set 2>[=launch mode 2] [@arch 2]" ... -gtool "<command line for a tool n>:<ranks set n>[=launch mode 3][@arch n]" <executable>

In the syntax, the separator ';' and the -gtool option are interchangeable.

Arguments

<arg></arg>	Parameters
<rank set=""></rank>	Specify the range of ranks that are involved in the tool execution. Separate ranks with a comma or use the '-' symbol for a set of contiguous ranks. To run the tool for all ranks, use the all argument.
	NOTE If you specify incorrect rank index, the corresponding warning is printed and the tool continues working for valid ranks.
[=launch mode]	Specify the launch mode (optional). See below for the available values.

[@arch]	Specify the architecture on which the tool runs (optional). For a given < rank
	set>, if you specify this argument, the tool is launched only for the processes
	residing on hosts with the specified architecture. This parameter is optional.

NOTE Rank sets cannot overlap for the same @arch parameter. Missing @arch parameter is also considered a different architecture. Thus, the following syntax is considered valid: -gtool "gdb: 0-3=attach; gdb: 0-3=attach@hsw; /usr/bin/gdb: 0-3=attach@knl"Also, note that some tools cannot work together or their simultaneous use may lead to incorrect results. The following table lists the parameter values for [=launch mode]:

[=launch mode]	Tool launch mode (optional). You can specify several values for each tool, which are separated with a comma ','.	
exclusive	Specify this value to prevent the tool from launching for more than one rank per host.	
attach	Specify this value to attach the tool from <code>-gtool</code> to the executable. If you use debuggers or other tools that can attach to a process in a debugger manner, you need to specify this value. This mode has been tested with debuggers only.	
node-wide	Specify this value to apply the tool from <code>-gtool</code> to all ranks where the <code><rank set=""></rank></code> resides or for all nodes in the case of all ranks. That is, the tool is applied to a higher level than the executable (to the <code>pmi_proxy</code> daemon).	
	Use the <code>-remote</code> argument for ranks to use the tool on remote nodes only.	

NOTE The tool attached to an MPI process may be executed without having access to stdin. To pass input to it, run a rank under the tool directly, for example: -gtool "gdb --args:0"

Examples

The following examples demonstrate different scenarios of using the -gtool option.

Example 1

Launch the Intel® VTune™ Amplifier XE and Valgrind* through the mpirun command:

```
$ mpirun -n 16 -gtool "vtune -collect hotspots -analyze-system \
-r result1:5,3,7-9=exclusive@bdw;valgrind -log-file=log_%p:0,1,10-12@hsw" a.out
```

This command launches <code>vtune</code> for the processes that are run on the Intel® microarchitecture codenamed Broadwell. Only one copy of <code>vtune</code> is launched for each host, the process with the minimal index is affected. At the same time, <code>Valgrind*</code> is launched for all specified processes that are run on the Intel® microarchitecture codenamed Haswell. <code>Valgrind's</code> results are saved to the files <code>log <process ID></code>.

Example 2

Set different environment variables for different rank sets:

```
$ mpirun -n 16 -gtool "env VARIABLE1=value1 VARIABLE2=value2:3,5,7-9; env VARIABLE3=value3:0,11"
a.out
```

Example 3

Apply a tool for a certain process through the -machinefile option.

In this example, suppose ${\tt m_file}$ has the following content:

```
$ cat ./m_file
hostname_1:2
hostname_2:3
hostname_3:1
```

The following command line demonstrates how to use the -machinefile option to apply a tool:

```
$ mpirun -n 6 -machinefile m_file -gtool "vtune -collect hotspots -analyze-system \
-r result1:5,3=exclusive@hsw;valgrind:0,1@bdw" a.out
```

In this example, the use of -machinefie option means that processes with indices 0 and 1 are located on the hostname_1 machine, process 3 is located on the hostname_2 machine, and process 5 - on the hostname_3 machine. After that, vtune is applied only ranks 3 and 5 (since these ranks belong to different machines, the exclusive option matches both of them) in case if hostname_2 and hostname_3 machines have Intel® microarchitecture codenamed Haswell. At the same time, the Valgrind* tool is applied to both ranks allocated on hostname 1 machine in case if it has Intel® microarchitecture codenamed Broadwell.

```
-gtoolfile <gtool_config_file>
```

Use this option to specify the -gtool parameters in a configuration file. All the same rules apply. Additionally, you can separate different command lines with section breaks.

For example, if gtool config file contains the following settings:

```
env VARIABLE1=value1 VARIABLE2=value2:3,5,7-9; env VARIABLE3=value3:0,11 env VARIABLE4=value4:1,12
```

The following command sets VARIABLE1 and VARIABLE2 for processes 3, 5, 7, 8, and 9 and sets VARIABLE3 for processes 0 and 11, while VARIABLE4 is set for processes 1 and 12:

```
$ mpirun -n 16 -gtoolfile gtool_config_file a.out
```

NOTE The options and the environment variable <code>-gtool</code>, <code>-gtoolfile</code> and <code>I_MPI_GTOOL</code> are mutually exclusive. The options <code>-gtool</code> and <code>-gtoolfile</code> are of the same priority and have higher priority than <code>I_MPI_GTOOL</code>. The first specified option in a command line is effective and the second one is ignored. Therefore, use <code>I_MPI_GTOOL</code> if you do not specify <code>-gtool</code> or <code>-gtoolfile</code>.

cpuinfo

Provides information on processors used in the system.

Syntax

cpuinfo [[-]<options>]

Arguments

<pre><options></options></pre>	Sequence of one-letter options. Each option controls a specific part of the output data.
g	General information about single cluster node shows:
	 the processor product name the number of packages/sockets on the node core and threads numbers on the node and within each package SMT mode enabling
i	Logical processors identification table identifies threads, cores, and packages of each logical processor accordingly.

	 Processor - logical processor number. ThreadId - unique processor identifier within a core. CoreId - unique core identifier within a package. PackageId - unique package identifier within a node.
d	Node decomposition table shows the node contents. Each entry contains the information on packages, cores, and logical processors.
	 Package Id - physical package identifier. Cores Id - list of core identifiers that belong to this package. Processors Id - list of processors that belong to this package. This list order directly corresponds to the core list. A group of processors enclosed in brackets belongs to one core.
С	Cache sharing by logical processors shows information of sizes and processors groups, which share particular cache level.
	 Size - cache size in bytes. Processors - a list of processor groups enclosed in the parentheses those share this cache or no sharing otherwise.
s	Microprocessor signature hexadecimal fields (Intel platform notation) show signature values:
	 extended family extended model family model type stepping
f	Microprocessor feature flags indicate what features the microprocessor supports. The Intel platform notation is used.
n	Table shows the following information about NUMA nodes:
	NUMA Id - NUMA node identifier.Processors - a list of processors in this node.
	If the node has no processors, the node is not shown.
A	Equivalent to gidesf
gidc	Default sequence
?	Utility usage info

Description

The <code>cpuinfo</code> utility prints out the processor architecture information that can be used to define suitable process pinning settings. The output consists of a number of tables. Each table corresponds to one of the single options listed in the arguments table.

NOTE

The architecture information is available on systems based on the Intel® 64 architecture.

The cpuinfo utility is available for both Intel microprocessors and non-Intel microprocessors, but it may provide only partial information about non-Intel microprocessors.

An example of the cpuinfo output:

```
$ cpuinfo -gdcs
==== Processor composition =====
Processor name : Intel(R) Xeon(R) X5570
Packages (sockets) : 2
Cores
          : 8
Processors (CPUs) : 8
Cores per package: 4
Threads per core : 1
==== Processor identification =====
Processor Thread Id. Core Id.
                                          Package Id.
0
              0
                             0
1
              0
                            0
                                           1
2
              0
                            1
3
              0
                            1
                                            1
                             2
4
              0
                                            0
5
              0
                             2
              0
                            3
                                            0
              0
                            3
==== Placement on packages =====
Package Id. Core Id. Processors 0 0,1,2,3 0,2,4,6
1
             0,1,2,3
                           1,3,5,7
==== Cache sharing =====
Cache Size
                     Processors
L1 32 KB
                    no sharing
      256 KB no sharing
8 MB (0,2,4,6) (3
L2
                     (0,2,4,6)(1,3,5,7)
==== Processor Signature =====
| xFamily | xModel | Type | Family | Model | Stepping |
1 00
         | 1
                 | 0
                        | 6
                                | a
                                        | 5
```

impi_info

Provides information on available Intel® MPI Library environment variables.

Syntax

impi info <options>

Arguments

<pre><options></options></pre>	List of options.
-a -all	Show all IMPI variables.
-h -help	Show a help message.
-v -variable	Show all available variables or description of the specified variable.
-c -category	Show all available categories or variables of the specified category.
-e -expert	Show all expert variables.

Description

The impi_info utility provides information on environment variables available in the Intel MPI Library. For each variable, it prints out the name, the default value, and the value data type. By default, a reduced list of variables is displayed. Use the -all option to display all available variables with their descriptions.

The example of the impi info output:

mpitune

Tunes the Intel® MPI Library parameters for the given MPI application.

Syntax

mpitune <options>

Arguments

mpitune Options	Options
-c config-file <file></file>	Specify a configuration file to run a tuning session.
-d dump-file <file></file>	Specify a file that stores the collected results. The option is used in the analyze mode.
-m mode {collect analyze}	Specify the mpitune mode. The supported modes are collect and analyze:
	 the collect mode runs the tuning process and saves results in temporary files; the analyze mode transforms temporary files into a JSON-tree, which is used by the Intel MPI Library, and generates a table that represents algorithm values in a human-readable format.
-h help	Display the help message.
-v version	Display the product version.

Description

The mpitune utility allows you to automatically adjust Intel MPI Library parameters, such as collective operation algorithms, to your cluster configuration or application.

The tuner iteratively launches a benchmarking application with different configurations to measure performance and stores the results of each launch. Based on these results, the tuner generates optimal values for the parameters being tuned.

NOTE Starting with the Intel MPI Library 2019 Update 4 release, you must specify two mpitune configuration files, which differ in their mode and dump-file fields. A simpler alternative may be to use one of the single configuration file templates shipped with the Intel MPI Library. In this case, you must use the command line to define the mode and dump-file fields.

• The -mode option defines one of two possible MPI tune modes: collect or analyze.

• The -dump-file option defines the path to the temporary files when in analyze mode. This path is returned by mpitune after the first iteration.

The configuration files should specify all tuner parameters, which are passed to the tuner with the -config-file option. A typical configuration file consists of the main section, specifying generic options, and search space sections for specific library parameters (for example, for specific collective operations). To comment a line, use the hash symbol #. All configuration file examples are available at $<installdir>/etc/tune_cfg$. Please note that configuration files for Intel® MPI Benchmarks are already created.

The tuning process consists of two steps: data collection (the collect mode) and data analysis (the analyze mode):

```
$ mpitune -m collect -c <path-to-config-file2>
$ mpitune -m analyze -c <path-to-config-file1>
```

Another variant of the launch is:

```
$ mpitune -m analyze -c <path-to-config-file1>
```

where the path to the dump-file received in the first step is used in the config file with templates inside.

The tuning results are presented as a JSON tree and can be added to the library with the I_MPI_TUNING environment variable.

MPI Options Support

The following MPI options are available within the utility:

MPI Options	Description	
-f <filename></filename>	Specify a file containing host names.	
-hosts <hostlist></hostlist>	Specify a comma-separated list of hosts.	
-np < <i>value</i> >	Specify the number of processes.	

Examples

```
$ mpitune -np 2 -ppn 1 -hosts HOST1,HOST2 -m collect -c <path-to-config-file2>
$ mpitune -np 2 -ppn 1 -hosts HOST1,HOST2 -m analyze -c <path-to-config-file1>
```

See Also

MPI Tuning in the Developer Guide.

For available configuration options, refer to mpitune Configuration Options.

mpitune Configuration Options

Application Options

-app

Sets a template for the command line to be launched to gather tuning results. The command line can contain variables declared as @<var name>@. The variables are defined further on using other options.

For example:

```
-app: mpirun -np @np@ -ppn @ppn@ IMB-MPI1 -msglog 0:@logmax@ -npmin @np@ @func@
```

NOTE The application must produce output (in stdout or file or any other destination) that can be parsed by the tuner to pick the value to be tuned and other variables. See the <code>-app-regex</code> and <code>-app-regex-legend</code> options below for details.

-app-regex

Sets a regular expression to be evaluated to extract the required values from the application output. Use regular expression groups to assign the values to variables. Variables and groups associations are set using the <code>-app-regex-legend</code> option.

For example, to extract the #bytes and t max[usec] values from this output:

```
#bytes #repetitions t_min[usec] t_max[usec] t_avg[usec]
0 1000 0.06 0.06 0.06
1 1000 0.10 0.10 0.10
```

use the following configuration:

```
-app-regex: (\d+)\s+\d+\s+[\d.+-]+\s+([\d.+-]+)
```

-app-regex-legend

Specifies a list of variables extracted from the regular expression. Variables correspond to the regular expression groups. The tuner uses the last variable as the performance indicator of the launch. Use the -tree-opt to set the optimization direction of the indicator.

For example:

```
-app-regex-legend: size, time
```

-iter

Sets the number of iterations for each launch with a given set of parameters. Higher numbers of iterations increase accuracy of results.

For example: -iter: 3

Search Space Options

Use these options to define a search space, which is a set of combinations of Intel® MPI Library parameters that the target application uses for launches. The library parameters are generally configured using run-time options or environment variables.

NOTE A search space line can be very long, so line breaking is available for all the search space options. Use a backslash to break a line (see examples below).

-search

Defines the search space by defining variables declared with the -app option and by setting environment variables for the application launch.

For example:

```
-search: func=BCAST, \
np=4,ppn={1,4},{,I_MPI_ADJUST_BCAST=[1,3]},logmax=5
```

The -app variables are defined as <var1>=<value1>[,<var2>=<value2>][,...]. The following syntax is available for setting values:

Syntax	Description	Examples
<value></value>	Single value. Can be a number or a string.	4
{ <value1>[,<value2 >][,]}</value2 </value1>	List of independent values.	{2,4}
[<start>,<end>[,<st ep>]]</st </end></start>	Linear range of values with the default step of 1.	$[1,8,2]$ — expands to $\{1,2,4,6,8\}$
(<start>,<end>[,<st ep>])</st </end></start>	Exponential range with the default step of 2.	$(1,16)$ — expands to $\{1,2,4,8,16\}$

To set environment variables for the command launch, use the following syntax:

Syntax	Description	Examples
<variable>=<value></value></variable>	Single variable definition. Any type of the syntax above can be used for the value: single values, lists or ranges.	I_MPI_ADJUST_BCAST=3 I_MPI_ADJUST_BCAST=[1,3]
{, <variable>=<value>}</value></variable>	A special case of the syntax above. When set this way, the variable default value is first used in an application launch.	<pre>{,I_MPI_ADJUST_BCAST=[1,3] }</pre>
<pre><prefix>{<value1> [,<value2>][,]}</value2></value1></prefix></pre>	Multi-value variable definition.	<pre>I_MPI_ADJUST_ALLREDUCE{=1,</pre>
[, <value2][,]}<="" td=""><td rowspan="2">Prefix is a common part for all the values, commonly the variable name.</td><td>=2, (=9,_KN_RADIX=(2,8))}</td></value2>	Prefix is a common part for all the values, commonly the variable name.	=2, (=9,_KN_RADIX=(2,8))}
		See below for a more complete example.
	A value can be a singular value or a combination of values in the format: <pre><prefix>(<value1>[, <value2>][,]). Prefix is optional and a value in the combination is a string, which can utilize the list and range syntax above.</value2></value1></prefix></pre>	

The following example shows a more complex option definition:

```
I MPI ADJUST BCAST\{=1,=2, (=9, \text{ KN RADIX}=(2,8)), (=\{10,11\}, \text{ SHM KN RADIX}=[2,8,2])\}
```

This directive consecutively runs the target application with the following environment variables set:

```
I_MPI_ADJUST_BCAST=1
I_MPI_ADJUST_BCAST=2
I_MPI_ADJUST_BCAST=9,I_MPI_ADJUST_BCAST_KN_RADIX=2
I_MPI_ADJUST_BCAST=9,I_MPI_ADJUST_BCAST_KN_RADIX=4
I_MPI_ADJUST_BCAST=9,I_MPI_ADJUST_BCAST_KN_RADIX=8
I_MPI_ADJUST_BCAST=10,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=2
I_MPI_ADJUST_BCAST=10,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=4
I_MPI_ADJUST_BCAST=10,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=6
I_MPI_ADJUST_BCAST=10,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=8
I_MPI_ADJUST_BCAST=11,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=2
I_MPI_ADJUST_BCAST=11,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=4
I_MPI_ADJUST_BCAST=11,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=6
I_MPI_ADJUST_BCAST=11,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=6
I_MPI_ADJUST_BCAST=11,I_MPI_ADJUST_BCAST_SHM_KN_RADIX=8
```

-search-excl

Excludes certain combinations from the search space. The syntax is identical to that of the <code>-search</code> option. For example:

```
-search-excl: I_MPI_ADJUST_BCAST={1,2}

or

-search-excl: func=BCAST,np=4,ppn=1,I MPI ADJUST BCAST=1
```

-search-only

Defines a subset of the search space to search in. Only this subset is used for application launches. The syntax is identical to the -search option.

This option is useful for the second and subsequent tuning sessions on a subset of parameters from the original session, without creating a separate configuration file.

Output Options

Use these options to customize the output. The tuner can produce output of two types:

- · table— useful for verifying the tuning results, contains values from all the application launches
- tree— an internal output format, contains the optimal values

-table

Defines the layout for the resulting output table. The option value is a list of variables declared with the -app option, which are joined in colon-separated groups. Each group denotes a specific part of the table.

For example:

```
-table: func:ppn,np:size:*:time
```

The last group variables (time) are rendered in table cells. The second last group variables are used for building table columns (*, denotes all the variables not present the other variable groups). The third last group variables are used for building table rows (size). All other variable groups are used to make up the table label. Groups containing several variables are complex groups and produce output based on all the value combinations.

For example, the option definition above can produce the following output:

```
Label: "func=BCAST, ppn=2, np=2"
Legend:
set 0: ""
set 1: "I MPI ADJUST BCAST=1"
set 2: "I MPI ADJUST BCAST=2"
set 3: "I MPI ADJUST_BCAST=3"
Table:
          | set 0
                    "size=0" | "time=0.10" | "time=0.08" | "time=0.11" | "time=0.10"
          | "time=0.12" | "time=0.09" | "time=0.12" | "time=0.11"
                      | "time=0.10" |
             -----|----|
"size=4" | "time=1.12" | "time=1.11" | "time=1.94" | "time=1.72"
         | "time=1.35" | "time=1.18" | "time=1.97" | "time=1.81"
         | "time=1.38" | "time=1.23" | "time=2.11" | "time=1.89"
```

Cells include only unique values from all the launches for the given parameter combination. The number of launches is set with the -iter option.

-table-ignore

Specifies the variables to ignore from the -table option definition.

-tree

Defines the layout for the resulting tree of optimal values of the parameter that is tuned (for example, collective operation algorithms). The tree is rendered as a JSON structure. The option value is a list of variables declared with the -app option, which are joined in colon-separated groups. Each group denotes a specific part of the tree. Groups containing several variables are complex groups and produce output based on all the value combinations.

Example:

```
-tree: func:ppn,np:size:*:time
```

The first two groups (func and ppn, np) make up the first two levels of the tree. The last group variables (time) are used as the optimization criteria and are not rendered. The second last group contains variables to be optimized (*, denotes all the variables not present the other variable groups). The third last group variables are used to split the search space into intervals based on the optimal values of parameters from the next group (for example, I MPI ADJUST operation > algorithm numbers).

For example, the option definition above can produce the following output:

This tree representation is an intermediate format of tuning results and is ultimately converted to a string that the library can understand. The conversion script is specified with -tree-postprocess option.

-tree-ignore

Specifies the variables to ignore from the -tree option definition.

-tree-intervals

Specifies the maximum number of intervals where the optimal parameter value is applied. If not specified, any number of intervals is allowed.

-tree-tolerance

Specifies the tolerance level. Non-zero tolerance (for example, 0.03 for 3%) joins resulting intervals with the performance indicator value varying by the specified tolerance.

-tree-postprocess

Specifies an executable to convert the resulting JSON tree to a custom format.

-tree-opt

Specifies the optimization direction. The available values are max (default) and min.

-tree-file

Specifies a log file where the tuning results are saved.

-tree-view

Specify the mode to present the json-tree. The available values are "simple" and "default". The "default" mode enables an interpolation mechanism; the "simple" mode disables the interpolation mechanism. The resulting tree contains message sizes used during the launch.

-mode

Specifies the mpitune mode. The available values are "collect" for gathering data and "analyze" for converting this data to a JSON-tree. Note that the -mode field can be defined in the configuration file as macros @-mode@, although the real value must be defined in the command line.

-dump-file

Specifies the path for the dump-file, which is returned by mpitune after the first iteration. The first iteration can be initialized by way of "" (an nempty string). Note that the -dump-file field can be defined in the configuration file as macros @-dump-file@, although the real value must be defined in the command line.

mpitune fast

This utility tunes the Intel® MPI Library to the cluster configuration using the Autotuning functionality.

Syntax

mpitune fast <options>

Arguments

Options	Description
-c colls	Set custom collective operations to tune, delimited by commas.
-d results_dir <path-to-results-dir></path-to-results-dir>	Set custom directory for tuning results, host files, and logs. Default: the current working directory.
-h help	Display the help message.
-n < <i>n</i> >	Specify the number of nodes. This can be a commadelimited set of values to set up several launches.
-pd <path-to-performance-results-dir>,</path-to-performance-results-dir>	Set a custom directory for validation performance results.

Options	Description
perf_results_dir <path-to-performance- results-dir></path-to-performance- 	<pre>Default: performance_results_<timestamp>.</timestamp></pre>
-ppn < <i>n</i> >	Specify the number of processes per node. This can be a comma-delimited set of values to set up several launches.
-val <path-to-tuning-file>,</path-to-tuning-file>	Run the tuning file validation cycle to validate the existing
validate <path-to-tuning-file></path-to-tuning-file>	tuning file.

Description

The mpitune_fast utility allows you to automatically set up the Intel MPI Library and launch with Autotuning enabled and configured for your cluster configuration.

The tool iteratively launches the Intel® MPI Benchmarks utility with the proper autotuner environment and generates a .dat file with the tuning parameters for your cluster configuration.

After generation the tuning file, set it as I MPI TUNING BIN:

```
$ export I_MPI_TUNING_BIN=./tuning_results.dat
$ mpiexec <args>
```

Workload Managers Support

mpitune_fast supports Slurm and LSF workload managers. It automatically defines job allocated hosts and performs launches.

MPI Options Support

The following MPI options are available within the utility:

Options	Description
-f <filename></filename>	Specify a file containing host names.
-hosts HOSTS,hosts	Set the host names, delimited by commas. Example:hosts
HOSTS	host1,host2

Example (default launch):

```
$ mpitune fast -f ./hostfile
```

Example (customized launch):

```
$ mpitune fast -ppn 8,4,2,1 -f ./hostfile -c alltoall,allreduce,barrier
```

See Also

Autotuning

mpitune

MPI Tuning in the Intel MPI Library Developer Guide

For available configuration options, refer to mpitune Configuration Options.

2

Environment Variable Reference

This section provides information on different variables:

- Compilation Environment Variables
- Hydra Environment Variables
- I_MPI_ADJUST Family Environment Variables
- Environment Variables for Process Pinning
- Environment Variables for Fabrics Control
- Environment Variables for Asynchronous Progress Control
- Environment Variables for Multi-EP
- Other Environment Variables

Compilation Environment Variables

I MPI {CC,CXX,FC,F77,F90} PROFILE

Specify the default profiling library.

Syntax

```
I_MPI_CC_PROFILE=<profile-name>
I_MPI_CXX_PROFILE=<profile-name>
I_MPI_FC_PROFILE=<profile-name>
I_MPI_F77_PROFILE=<profile-name>
I_MPI_F90_PROFILE=<profile-name>
```

Argument

cprofile-name>

Specify a default profiling library.

Description

Set this environment variable to select a specific MPI profiling library to be used by default. This has the same effect as using -profile=cprofile=name as an argument for mpiico or another Intel® MPI Library compiler wrapper.

I_MPI_TRACE_PROFILE

Specify the default profile for the -trace option.

Syntax

I_MPI_TRACE_PROFILE=profile-name>

Argument

cprofile-name>

Specify a tracing profile name. The default value is ${\tt vt}\, .$

Description

Set this environment variable to select a specific MPI profiling library to be used with the -trace option of mpiico or another Intel MPI Library compiler wrapper.

The I MPI {CC,CXX,F77,F90} PROFILE environment variable overrides I MPI TRACE PROFILE.

I_MPI_CHECK_PROFILE

Specify the default profile for the -check mpi option.

Syntax

```
I_MPI_CHECK_PROFILE=profile-name>
```

Argument

<pre><pre><pre><pre><pre><pre><pre><pre></pre></pre></pre></pre></pre></pre></pre></pre>	Specify the checking profile name. The default value is
	vtmc.

Description

Set this environment variable to select a specific MPI checking library to be used with the <code>-check_mpi</code> option to <code>mpiicc</code> or another Intel MPI Library compiler wrapper.

The I_MPI_{CC,CXX,F77,F90}_PROFILE environment variable overrides I_MPI_CHECK PROFILE.

I_MPI_CHECK_COMPILER

Turn on/off compiler compatibility check.

Syntax

```
I MPI CHECK COMPILER=<arg>
```

Arguments

<arg></arg>	Binary indicator.
enable yes on 1	Enable checking the compiler.
disable no off 0	Disable checking the compiler. This is the default value.

Description

If I_MPI_CHECK_COMPILER is set to enable, the Intel MPI Library compiler wrapper checks the underlying compiler for compatibility. Normal compilation requires using a known version of the underlying compiler.

I_MPI_{CC,CXX,FC,F77,F90}

Set the path/name of the underlying compiler to be used.

Syntax

```
I_MPI_CC=<compiler>
I_MPI_CXX=<compiler>
I_MPI_FC=<compiler>
I_MPI_F77=<compiler>
I_MPI_F90=<compiler>
```

Arguments

<compiler></compiler>	Specify the full path/name of compiler to be used.
-----------------------	--

Description

Set this environment variable to select a specific compiler to be used. Specify the full path to the compiler if it is not located in the search path.

NOTE Some compilers may require additional command line options.

NOTE The configuration file is sourced if it exists for a specified compiler. See -configfor details.

I_MPI_ROOT

Set the Intel MPI Library installation directory path.

Syntax

I MPI ROOT=<path>

Arguments

<path></path>	Specify the installation directory of the Intel MPI Library.
---------------	--

Description

Set this environment variable to specify the installation directory of the Intel MPI Library.

NOTE If you are using the Visual Studio integration, you may need to use I MPI ONEAPI ROOT.

VT_ROOT

Set Intel® Trace Collector installation directory path.

Syntax

VT ROOT=<path>

Arguments

<path></path>	Specify the installation directory of the Intel Trace
	Collector.

Description

Set this environment variable to specify the installation directory of the Intel Trace Collector.

I_MPI_COMPILER_CONFIG_DIR

Set the location of the compiler configuration files.

Syntax

I_MPI_COMPILER_CONFIG_DIR=<path>

Arguments

<path></path>	Specify the location of the compiler configuration files.
	The default value is <install-dir>/etc</install-dir>

Description

Set this environment variable to change the default location of the compiler configuration files.

I MPI LINK

Select a specific version of the Intel MPI Library for linking.

Syntax

I_MPI_LINK=<arg>

Arguments

Argument	Library Version
opt	Multi-threaded optimized library (with the global lock). This is the default value
dbg	Multi-threaded debug library (with the global lock)
opt_mt	Multi-threaded optimized library (with per-object lock for the thread-split model)
dbg_mt	Multi-threaded debug library (with per-object lock for the thread-split model)

Description

Set this variable to always link against the specified version of the Intel MPI Library.

I_MPI_DEBUG_INFO_STRIP-

Turn on/off the debug information stripping while linking applications statically.

Syntax

```
I MPI DEBUG INFO STRIP=<arg>
```

Arguments

<arg></arg>	Binary indicator
enable yes on 1	Turn on. This is the default value
disable no off 0	Turn off

Description

Use this option to turn on/off the debug information stripping while linking the Intel MPI Library statically. Debug information is stripped by default.

I_MPI_{C,CXX,FC,F}FLAGS

Set special flags needed for compilation.

Syntax

```
I_MPI_CFLAGS=<flags>
I_MPI_CXXFLAGS=<flags>
I_MPI_FCFLAGS=<flags>
I_MPI_FFLAGS=<flags>
```

Arguments

<flags></flags>	Flag list	
-----------------	-----------	--

Description

Use this environment variable to specify special compilation flags.

I_MPI_LDFLAGS

Set special flags needed for linking.

Syntax

I MPI LDFLAGS=<flags>

Arguments

<flags></flags>	Flag list
-----------------	-----------

Description

Use this environment variable to specify special linking flags.

I_MPI_FORT_BIND

Disable mpiicc linking with Fortran bindings.

Syntax

```
I_MPI_FORT_BIND=<arg>
```

Arguments

<arg></arg>	Binary indicator
enable yes on 1	Enable linking. This is the default value
disable no off 0	Disable linking

Description

By default, the mpiicc also links against the Fortran bindings even if Fortran is not used. Use this environment variable to change this default behavior. Has the same effect as the -nofortbind option.

Hydra Environment Variables

I_MPI_HYDRA_HOST_FILE

Set the host file to run the application.

Syntax

I MPI HYDRA HOST FILE=<arg>

Argument

<arg></arg>	String parameter
<hostsfile></hostsfile>	The full or relative path to the host file

Description

Set this environment variable to specify the hosts file.

I_MPI_HYDRA_HOSTS_GROUP

Set node ranges using brackets, commas, and dashes.

Syntax

```
I_MPI_HYDRA_HOSTS_GROUP=<arg>
```

Argument

<arg></arg>	Set a node range.
-------------	-------------------

Description

Set this variable to be able to set node ranges using brackets, commas, and dashes (like in Slurm* Workload Manager). For example:

```
I MPI HYDRA HOSTS GROUP="hostA[01-05], hostB, hostC[01-05,07,09-11]"
```

You can set node ranges with the -hosts-group option.

I MPI HYDRA DEBUG

Print out the debug information.

Syntax

```
I_MPI_HYDRA_DEBUG=<arg>
```

Argument

<arg></arg>	Binary indicator
enable yes on 1	Turn on the debug output
disable no off 0	Turn off the debug output. This is the default value

Description

Set this environment variable to enable the debug mode.

I_MPI_HYDRA_ENV

Control the environment propagation.

Syntax

```
I MPI HYDRA ENV=<arg>
```

Argument

<arg></arg>	String parameter
all	Pass all environment to all MPI processes

Description

Set this environment variable to control the environment propagation to the MPI processes. By default, the entire launching node environment is passed to the MPI processes. Setting this variable also overwrites environment variables set by the remote shell.

I_MPI_JOB_TIMEOUT

Set the timeout period for mpiexec.hydra.

Syntax

```
I_MPI_JOB_TIMEOUT=<timeout>
I_MPI_MPIEXEC_TIMEOUT=<timeout>
```

Argument

<timeout></timeout>	Define mpiexec.hydra timeout period in seconds
<n> ≥ 0</n>	The value of the timeout period. The default timeout value is zero, which
	means no timeout.

Description

Set this environment variable to make mpiexec.hydra terminate the job in <timeout> seconds after its launch. The <timeout> value should be greater than zero. Otherwise the environment variable setting is ignored.

I_MPI_JOB_STARTUP_TIMEOUT

Set the mpiexec.hydra job startup timeout.

Syntax

```
I_MPI_JOB_STARTUP_TIMEOUT=<timeout>
```

Argument

<timeout></timeout>	Define mpiexec.hydra startup timeout period in seconds
< <i>n></i> ≥ 0	The value of the timeout period. The default timeout value is zero, which
	means no timeout.

Description

Set this environment variable to make mpiexec.hydra terminate the job in <timeout> seconds if some processes are not launched. The <timeout> value should be greater than zero.

I_MPI_JOB_TIMEOUT_SIGNAL

Define the signal to be sent when a job is terminated because of a timeout.

Syntax

I_MPI_JOB_TIMEOUT_SIGNAL=<number>

Argument

<number></number>	Define the signal number
<n>> 0</n>	The signal number. The default value is 9 (SIGKILL)

Description

Define a signal number to be sent to stop the MPI job if the timeout period specified by the I_MPI_JOB_TIMEOUT environment variable expires. If you set a signal number unsupported by the system, the mpiexec.hydra command prints a warning message and continues the task termination using the default signal number 9 (SIGKILL).

I_MPI_JOB_ABORT_SIGNAL

Define a signal to be sent to all processes when a job is terminated unexpectedly.

Syntax

I MPI JOB ABORT SIGNAL=<number>

Argument

<number></number>	Define signal number
<n>> 0</n>	The default value is 9 (SIGKILL)

Description

Set this environment variable to define a signal for task termination. If you set an unsupported signal number, mpiexec.hydra prints a warning message and uses the default signal 9 (SIGKILL).

I_MPI_JOB_SIGNAL_PROPAGATION

Control signal propagation.

Syntax

I MPI JOB SIGNAL PROPAGATION=<arg>

Argument

<arg></arg>	Binary indicator
enable yes on 1	Turn on propagation
disable no off 0	Turn off propagation. This is the default value

Description

Set this environment variable to control propagation of the signals (SIGINT, SIGALRM, and SIGTERM). If you enable signal propagation, the received signal is sent to all processes of the MPI job. If you disable signal propagation, all processes of the MPI job are stopped with the default signal 9 (SIGKILL).

I_MPI_HYDRA_BOOTSTRAP

Set the bootstrap server.

Syntax

I MPI HYDRA BOOTSTRAP=<arg>

Argument

<arg></arg>	String parameter
ssh	Use secure shell. This is the default value
rsh	Use remote shell
pdsh	Use parallel distributed shell
pbsdsh	Use Torque* and PBS* pbsdsh command
fork	Use fork call
slurm	Use Slurm* srun command
11	Use LoadLeveler* llspawn.stdio command
lsf	Use LSF* blaunch command
sge	Use Univa* Grid Engine* qrsh command

Description

Set this environment variable to specify the bootstrap server.

I_MPI_HYDRA_BOOTSTRAP_EXEC

Set the executable file to be used as a bootstrap server.

Syntax

I MPI HYDRA BOOTSTRAP EXEC=<arg>

Argument

<arg></arg>	String parameter
<executable></executable>	The name of the executable file

Description

Set this environment variable to specify the executable file to be used as a bootstrap server.

I_MPI_HYDRA_BOOTSTRAP_EXEC_EXTRA_ARGS

Set additional arguments for the bootstrap server.

Syntax

I_MPI_HYDRA_BOOTSTRAP_EXEC_EXTRA_ARGS=<arg>

Argument

<arg></arg>	String parameter
<args></args>	Additional bootstrap server arguments

Description

Set this environment variable to specify additional arguments for the bootstrap server.

NOTE If the launcher (blaunch, lsf, pdsh, pbsdsh) falls back to ssh, pass the arguments with the invocation of ssh.

I_MPI_HYDRA_BOOTSTRAP_AUTOFORK

Control the usage of fork call for local processes.

I MPI HYDRA BOOTSTRAP AUTOFORK = <arg>

Argument

<arg></arg>	String parameter
enable yes on 1	Use fork for the local processes. This is default value for ssh, rsh, ll, lsf,
	and pbsdsh bootstrap servers
disable no off 0	Do not use fork for the local processes. This is default value for the sge
	bootstrap server

Description

Set this environment variable to control usage of fork call for the local processes.

NOTE This option is not applicable to slurm and pdsh bootstrap servers.

I_MPI_HYDRA_RMK

Use the specified value as the resource management kernel to obtain data about available nodes, externally set process counts.

Syntax

I_MPI_HYDRA_RMK=<arg>

Argument

<arg></arg>	String parameter
<rmk></rmk>	Resource management kernel. The supported values are slurm, 11, 1sf,
	sge, pbs, cobalt.

Description

Set this environment variable to use the resource management kernel.

I_MPI_HYDRA_PMI_CONNECT

Define the processing method for PMI messages.

Syntax

I MPI HYDRA PMI CONNECT=<value>

Argument

<value></value>	The algorithm to be used
nocache	Do not cache PMI messages
cache	Cache PMI messages on the local pmi_proxy management processes to
	minimize the number of ${\tt PMI}$ requests. Cached information is automatically
	propagated to child management processes.
lazy-cache	cache mode with on-demand propagation.

alltoall	Information is automatically exchanged between all pmi_proxy before any
	get request can be done. This is the default value.

Description

Use this environment variable to select the PMI messages processing method.

I_MPI_PERHOST

Define the default behavior for the -perhost option of the mpiexec.hydra command.

Syntax

I MPI PERHOST=<value>

Argument

<value></value>	Define a value used for -perhost by default
integer > 0	Exact value for the option
all	All logical CPUs on the node
allcores	All cores (physical CPUs) on the node. This is the default value.

Description

Set this environment variable to define the default behavior for the -perhost option. Unless specified explicitly, the -perhost option is implied with the value set in I MPI PERHOST.

NOTE

When running under a job scheduler, this environment variable is ignored by default. To control process placement with <code>I_MPI_PERHOST</code>, disable the <code>I_MPI_JOB_RESPECT_PROCESS_PLACEMENT</code> variable.

I MPI JOB TRACE LIBS

Choose the libraries to preload through the -trace option.

Syntax

I_MPI_JOB_TRACE_LIBS=<arg>

Argument

<arg></arg>	String parameter
	Blank separated list of the libraries to preload. The default value is ${\tt vt}$

Description

Set this environment variable to choose an alternative library for preloading through the -trace option.

I_MPI_JOB_CHECK_LIBS

Choose the libraries to preload through the -check mpi option.

Syntax

I MPI JOB CHECK LIBS=<arg>

Argument

<arg></arg>	String parameter
	Blank separated list of the libraries to preload. The default value is ${\tt vtmc}$

Description

Set this environment variable to choose an alternative library for preloading through the -check mpi option.

I_MPI_HYDRA_BRANCH_COUNT

Set the hierarchical branch count.

Syntax

I_MPI_HYDRA_BRANCH_COUNT =<num>

Argument

<num></num>	Number
<n> >= 0</n>	The default value is 16. This value means that hierarchical structure is enabled if the number of nodes is more than 16.
	If $I_MPI_HYDRA_BRANCH_COUNT=0$, then there is no hierarchical structure.
	If $I_MPI_HYDRA_BRANCH_COUNT=-1$, then branch count is equal to default value.

Description

Set this environment variable to restrict the number of child management processes launched by the mpiexec.hydra operation or by each pmi_proxy management process.

I_MPI_HYDRA_PMI_AGGREGATE

Turn on/off aggregation of the PMI messages.

Syntax

I MPI HYDRA PMI AGGREGATE=<arg>

Argument

<arg></arg>	Binary indicator
enable yes on 1	Enable PMI message aggregation. This is the default value.
disable no off 0	Disable PMI message aggregation.

Description

Set this environment variable to enable/disable aggregation of PMI messages.

I_MPI_HYDRA_GDB_REMOTE_SHELL

Set the remote shell command to run the GDB debugger. This command uses the Intel® Distribution for GDB.

Syntax

I_MPI_HYDRA_GDB_REMOTE_SHELL=<arg>

Argument

<arg></arg>	String parameter
ssh	Secure Shell (SSH). This is the default value
rsh	Remote shell (RSH)

Description

Set this environment variable to specify the remote shell command to run the GNU* debugger on the remote machines. You can use this environment variable to specify any shell command that has the same syntax as SSH or RSH.

I_MPI_HYDRA_IFACE

Set the network interface.

Syntax

I MPI HYDRA IFACE=<arg>

Argument

<arg></arg>	String parameter
<network interface=""></network>	The network interface configured in your system

Description

Set this environment variable to specify the network interface to use. For example, use "-iface ib0", if the IP emulation of your InfiniBand* network is configured on ib0.

I_MPI_HYDRA_DEMUX

Set the demultiplexer (demux) mode.

Syntax

I MPI HYDRA DEMUX=<arg>

Argument

<arg></arg>	String parameter
poll	Set poll as the multiple I/O demultiplexer ($demux$) mode engine. This is the default value.
select	Set select as the multiple I/O demultiplexer (demux) mode engine

Description

Set this environment variable to specify the multiple I/O demux mode engine. The default value is poll.

I_MPI_TMPDIR

Specify a temporary directory.

Syntax

I_MPI_TMPDIR=<arg>

Argument

<arg></arg>	String parameter
<path></path>	Temporary directory. The default value is /tmp

Description

Set this environment variable to specify a directory for temporary files.

I_MPI_JOB_RESPECT_PROCESS_PLACEMENT

Specify whether to use the process-per-node placement provided by the job scheduler, or set explicitly.

Syntax

```
I_MPI_JOB_RESPECT_PROCESS_PLACEMENT=<arg>
```

Argument

<value></value>	Binary indicator
enable yes on 1	Use the process placement provided by job scheduler. This is the default value
disable no off 0	Do not use the process placement provided by job scheduler

Description

If the variable is set, the Hydra process manager uses the process placement provided by job scheduler (default). In this case the -ppn option and its equivalents are ignored. If you disable the variable, the Hydra process manager uses the process placement set with -ppn or its equivalents.

I_MPI_GTOOL

Specify the tools to be launched for selected ranks. An alternative to this variable is the -gtool option.

Syntax

I_MPI_GTOOL="<command line for a tool 1>:<ranks set 1>[=exclusive][@arch 1]; <command
line for a tool 2>:<ranks set 2>[=exclusive][@arch 2]; ...; <command line for a tool
n>:<ranks set n>[=exclusive][@arch n]"

Argument

_	
<arg></arg>	Parameters
<pre><command-line-for-a-tool></command-line-for-a-tool></pre>	Specify a tool's launch command, including parameters.
<rank set=""></rank>	Specify the range of ranks that are involved in the tool execution. Separate ranks with a comma or use the '-' symbol for a set of contiguous ranks. To run the tool for all ranks, use the all argument.
	NOTE If you specify incorrect rank index, the corresponding warning is printed and the tool continues working for valid ranks.
[=exclusive]	Specify this parameter to prevent launching a tool for more than one rank per host. This parameter is optional.
[@arch]	Specify the architecture on which the tool runs (optional). For a given < <i>rank set</i> >, if you specify this argument, the tool is launched only for the processes residing on hosts with the specified architecture. This parameter is optional.

Description

Use this option to launch the tools such as Intel® VTune™ Amplifier XE, Valgrind*, and GNU* Debugger for the specified processes.

Examples

The following command line examples demonstrate different scenarios of using the I_MPI_GTOOL environment variable.

Launch Intel® VTune™ Amplifier XE and Valgrind* by setting the I_MPI_GTOOL environment variable:

```
$ export I_MPI_GTOOL="vtune -collect hotspots -analyze-system -r result1:5,3,7-9=exclusive@bdw;\
valgrind -log-file=log_%p:0,1,10-12@hsw"
$ mpiexec.hydra -n 16 a.out
```

This command launches <code>vtune</code> for the processes that are run on the Intel® microarchitecture codenamed Broadwell. Only one copy of <code>vtune</code> is launched for each host, the process with the minimal index is affected. At the same time, <code>Valgrind*</code> is launched for all specified processes that are run on the Intel® microarchitecture codenamed Haswell. <code>Valgrind's</code> results are saved to the files <code>log <process ID></code>.

Launch GDB by setting the I_MPI_GTOOL environment variable (for Intel® oneAPI, this launches the Intel® Distribution for GDB):

```
$ mpiexec.hydra -n 16 -genv I MPI GTOOL="gdb:3,5,7-9" a.out
```

Use this command to apply GDB to the given rank set.

NOTE The options and the environment variable <code>-gtool</code>, <code>-gtoolfile</code> and <code>I_MPI_GTOOL</code> are mutually exclusive. The options <code>-gtool</code> and <code>-gtoolfile</code> are of the same priority and have higher priority than <code>I_MPI_GTOOL</code>. The first specified option in a command line is effective and the second one is ignored. Therefore, use <code>I_MPI_GTOOL</code> if you do not specify <code>-gtool</code> or <code>-gtoolfile</code>.

I_MPI_HYDRA_TOPOLIB

Set the interface for topology detection.

Syntax

I MPI HYDRA TOPOLIB=<arg>

Argument

<arg></arg>	String parameter
hwloc	The hwloc* library functions are invoked for topology detection.

Description

Set this environment variable to define the interface for platform detection. The hwloc* interface is used by default, but you may explicitly set the variable to use the native Intel MPI Library interface:

I MPI HYDRA TOPOLIB=ipl.

I_MPI_PORT_RANGE

Specify a range of allowed port numbers.

Syntax

I MPI PORT RANGE=<range>

Argument

<range></range>	String parameter
<min>:<max></max></min>	Allowed port range

Description

Set this environment variable to specify a range of the allowed port numbers for the Intel® MPI Library.

I MPI SILENT ABORT

Control abort warning messages.

Syntax

I MPI SILENT ABORT=<arg>

Argument

<arg></arg>	Binary indicator
enable yes on 1	Do not print abort warning message
disable no off 0	Print abort warning message. This is the default value

Description

Set this variable to disable printing of abort warning messages. The messages are printed in case of the MPI Abort call.

You can also disable printing of these messages with the -silent-abort option.

I_MPI_HYDRA_NAMESERVER

Specify the nameserver.

Syntax

I MPI HYDRA NAMESERVER=<arg>

Argument

<arg></arg>	String parameter
<hostname>:<port></port></hostname>	Set the hostname and the port.

Description

Set this variable to specify the nameserver for your MPI application in the following format:

```
I MPI HYDRA NAMESERVER = hostname:port
```

You can set the nameserver with the -nameserver option.

I_MPI_ADJUST Family Environment Variables

inclusive

I_MPI_ADJUST_<opname>

Control collective operation algorithm selection.

Syntax

 $\label{local_adjust_solutions} $$I_MPI_ADJUST_<opname>="<algid>[:<conditions>][;<algid>:<conditions>[...]]"$$

Arguments

<algid></algid>	Algorithm identifier
>= 0	Set a number to select the desired algorithm. The value 0 uses basic logic of the collective algorithm selection.
<conditions></conditions>	A comma separated list of conditions. An empty list selects all message sizes and process combinations
<1>	Messages of size <1>
<1>- <m></m>	Messages of size from <1> to <m>, inclusive</m>
<1>@	Messages of size $<1>$ and number of processes

Messages of size from <1> to <m> and number of processes from to <q>,

Description

<1>-<m>@-<q>

Set this environment variable to select the desired algorithm(s) for the collective operation *<opname>* under particular conditions. Each collective operation has its own environment variable and algorithms.

Environment Variables, Collective Operations, and Algorithms

Environment Variable	Collective Operation	Algorithms
I_MPI_ADJUST_ALLGATHER	MPI_Allgather	 Recursive doubling Bruck's Ring Topology aware Gatherv + Bcast Knomial
I_MPI_ADJUST_ALLGATHERV	MPI_Allgatherv	 Recursive doubling Bruck's Ring Topology aware Gatherv + Bcast
I_MPI_ADJUST_ALLREDUCE	MPI_Allreduce	 Recursive doubling Rabenseifner's Reduce + Bcast Topology aware Reduce + Bcast Binomial gather + scatter Topology aware binominal gather + scatter Shumilin's ring Ring Knomial Topology aware SHM-based flat Topology aware SHM-based Knomial Topology aware SHM-based Knomial Topology aware SHM-based Knomial
I_MPI_ADJUST_ALLTOALL	MPI_Alltoall	 Bruck's Isend/Irecv + waitall Pair wise exchange Plum's
I_MPI_ADJUST_ALLTOALLV	MPI_Alltoallv	 Isend/Irecv + waitall Plum's
I_MPI_ADJUST_ALLTOALLW	MPI_Alltoallw	Isend/Irecv + waitall
I_MPI_ADJUST_BARRIER	MPI_Barrier	 Dissemination Recursive doubling Topology aware dissemination Topology aware recursive doubling Binominal gather + scatter Topology aware binominal gather + scatter Topology aware SHM-based flat

Environment Variable	Collective Operation	Algorithms
		8. Topology aware SHM-based Knomial9. Topology aware SHM-based
I_MPI_ADJUST_BCAST	MPI_Bcast	 Knary Binomial Recursive doubling Ring Topology aware binomial Topology aware recursive doubling Topology aware ring Shumilin's Knomial Topology aware SHM-based flat Topology aware SHM-based Knomial Topology aware SHM-based Knary NUMA aware SHM-based (SSE4.2)
		13. NUMA aware SHM-based (AVX2)14. NUMA aware SHM-based (AVX512)
I_MPI_ADJUST_EXSCAN	MPI_Exscan	 Partial results gathering Partial results gathering regarding layout of processes
I_MPI_ADJUST_GATHER	MPI_Gather	 Binomial Topology aware binomial Shumilin's Binomial with segmentation
I_MPI_ADJUST_GATHERV	MPI_Gatherv	 Linear Topology aware linear Knomial
I_MPI_ADJUST_REDUCE_SCATTE R	MPI_Reduce_scatter	 Recursive halving Pair wise exchange Recursive doubling Reduce + Scatterv Topology aware Reduce + Scatterv
I_MPI_ADJUST_REDUCE	MPI_Reduce	 Shumilin's Binomial Topology aware Shumilin's Topology aware binomial Rabenseifner's

Environment Variable	Collective Operation	Algorithms
		 6. Topology aware Rabenseifner's 7. Knomial 8. Topology aware SHM-based flat 9. Topology aware SHM-based Knomial 10. Topology aware SHM-based Knary 11. Topology aware SHM-based binomial
I_MPI_ADJUST_SCAN	MPI_Scan	 Partial results gathering Topology aware partial results gathering
I_MPI_ADJUST_SCATTER	MPI_Scatter	 Binomial Topology aware binomial Shumilin's
I_MPI_ADJUST_SCATTERV	MPI_Scatterv	 Linear Topology aware linear
I_MPI_ADJUST_SENDRECV_REPL ACE	MPI_Sendrecv_replace	 Generic Uniform (with restrictions)
I_MPI_ADJUST_IALLGATHER	MPI_Iallgather	 Recursive doubling Bruck's Ring
I_MPI_ADJUST_IALLGATHERV	MPI_Iallgatherv	 Recursive doubling Bruck's Ring
I_MPI_ADJUST_IALLREDUCE	MPI_Iallreduce	 Recursive doubling Rabenseifner's Reduce + Bcast Ring (patarasuk) Knomial Binomial Reduce scatter allgather SMP Nreduce
I_MPI_ADJUST_IALLTOALL	MPI_Ialltoall	 Bruck's Isend/Irecv + Waitall Pairwise exchange
I_MPI_ADJUST_IALLTOALLV	MPI_Ialltoallv	Isend/Irecv + Waitall
I_MPI_ADJUST_IALLTOALLW	MPI_Ialltoallw	Isend/Irecv + Waitall
I_MPI_ADJUST_IBARRIER	MPI_Ibarrier	Dissemination
I_MPI_ADJUST_IBCAST	MPI_Ibcast	 Binomial Recursive doubling

Environment Variable	Collective Operation	Algorithms
		 Ring Knomial SMP Tree knominal Tree kary
I_MPI_ADJUST_IEXSCAN	MPI_Iexscan	 Recursive doubling SMP
I_MPI_ADJUST_IGATHER	MPI_Igather	 Binomial Knomial
I_MPI_ADJUST_IGATHERV	MPI_Igatherv	 Linear Linear ssend
I_MPI_ADJUST_IREDUCE_SCATT ER	MPI_Ireduce_scatter	 Recursive halving Pairwise Recursive doubling
I_MPI_ADJUST_IREDUCE	MPI_Ireduce	 Rabenseifner's Binomial Knomial
I_MPI_ADJUST_ISCAN	MPI_Iscan	 Recursive Doubling SMP
I_MPI_ADJUST_ISCATTER	MPI_Iscatter	 Binomial Knomial
I_MPI_ADJUST_ISCATTERV	MPI_Iscatterv	Linear

The message size calculation rules for the collective operations are described in the table. In the following table, "n/a" means that the corresponding interval <1>-<m> should be omitted.

NOTE The I_MPI_ADJUST_SENDRECV_REPLACE=2 ("Uniform") algorithm can be used only in the case when datatype and objects count are the same across all ranks.

Message Collective Functions

Collective Function	Message Size Formula
MPI_Allgather	recv_count*recv_type_size
MPI_Allgatherv	total_recv_count*recv_type_size
MPI_Allreduce	count*type_size
MPI_Alltoall	send_count*send_type_size
MPI_Alltoallv	n/a
MPI_Alltoallw	n/a
MPI_Barrier	n/a
MPI_Bcast	count*type_size
MPI_Exscan	count*type_size
MPI_Gather	<pre>recv_count*recv_type_size if MPI_IN_PLACE is used, otherwise send_count*send_type_size</pre>

Collective Function	Message Size Formula
MPI_Gatherv	n/a
MPI_Reduce_scatter	total_recv_count*type_size
MPI_Reduce	count*type_size
MPI_Scan	count*type_size
MPI_Scatter	<pre>send_count*send_type_size if MPI_IN_PLACE is used, otherwise recv_count*recv_type_size</pre>
MPI_Scatterv	n/a

Examples

Use the following settings to select the second algorithm for MPI Reduce operation:

```
I MPI ADJUST REDUCE=2
```

Use the following settings to define the algorithms for MPI Reduce scatter operation:

```
I MPI ADJUST REDUCE SCATTER="4:0-100,5001-10000;1:101-3200;2:3201-5000;3"
```

In this case. algorithm 4 is used for the message sizes between 0 and 100 bytes and from 5001 and 10000 bytes, algorithm 1 is used for the message sizes between 101 and 3200 bytes, algorithm 2 is used for the message sizes between 3201 and 5000 bytes, and algorithm 3 is used for all other messages.

I_MPI_ADJUST_<opname>_LIST

Syntax

```
I MPI ADJUST <opname> LIST=<algid1>[-<algid2>][,<algid3>][,<algid4>-<algid5>]
```

Description

Set this environment variable to specify the set of algorithms to be considered by the Intel MPI runtime for a specified <opname>. This variable is useful in autotuning scenarios, as well as tuning scenarios where users would like to select a certain subset of algorithms.

NOTE Setting an empty string disables autotuning for the <opname> collective.

I_MPI_COLL_INTRANODE

Syntax

I MPI COLL INTRANODE=<mode>

Arguments

<mode></mode>	Intranode collectives type
pt2pt	Use only point-to-point communication-based collectives
shm	Enables shared memory collectives. This is the default value

Description

Set this environment variable to switch intranode communication type for collective operations. If there is large set of communicators, you can switch off the SHM-collectives to avoid memory overconsumption.

I MPI COLL INTRANODE SHM THRESHOLD

Syntax

I MPI COLL INTRANODE SHM THRESHOLD=<nbytes>

Arguments

<nbytes></nbytes>	Define the maximal data block size processed by shared memory collectives.
> 0	Use the specified size. The default value is 16384 bytes.

Description

Set this environment variable to define the size of shared memory area available for each rank for data placement. Messages greater than this value will *not* be processed by SHM-based collective operation, but will be processed by point-to-point based collective operation. The value must be a multiple of 4096.

I_MPI_COLL_EXTERNAL

Syntax

I MPI COLL EXTERNAL=<arg>

Arguments

<arg></arg>	Binary indicator.
enable yes on 1	Enable the external collective operations functionality.
disable no off 0	Disable the external collective operations functionality. This is the default value.

Description

Set this environment variable to enable external collective operations. The mechanism allows to enable HCOLL. The functionality enables the following collective operations: I_MPI_ADJUST_ALLREDUCE=24, I_MPI_ADJUST_BARRIER=11, I_MPI_ADJUST_BCAST=16, I_MPI_ADJUST_REDUCE=13, I_MPI_ADJUST_ALLGATHER=6, I_MPI_ADJUST_ALLTOALL=5, I_MPI_ADJUST_ALLTOALLV=5, I_MPI_ADJUST_SCAN=3, I_MPI_ADJUST_EXSCAN=3, I_MPI_ADJUST_GATHER=5, I_MPI_ADJUST_GATHERV=4, I_MPI_ADJUST_SCATTER=5, I_MPI_ADJUST_SCATTERV=4, I_MPI_ADJUST_ALLGATHERV=5, I_MPI_ADJUST_ALLTOALLW=2, I_MPI_ADJUST_REDUCE_SCATTER=6, I_MPI_ADJUST_IALLGATHERV=5, I_MPI_ADJUST_IALLGATHERV=5, I_MPI_ADJUST_IALLGATHERV=5, I_MPI_ADJUST_IALLGATHERV=5, I_MPI_ADJUST_IALLGATHERV=5, I_MPI_ADJUST_IALLGATHERV=2, I_MPI_ADJUST_IBCAST=5, I_MPI_ADJUST_IREDUCE=4.

To force HCOLL usage, use the I_MPI_ADJUST_<opname> values described above. For reaching better performance, use an autotuner as soon as I_MPI_COLL_EXTERNAL is enabled in order to get the optimal collectives settings. For more information on HCOLL tuning, refer to NVIDIA* documentation.

I MPI CBWR

Control reproducibility of floating-point operations results across different platforms, networks, and topologies in case of the same number of processes.

Syntax

I MPI CBWR=<arg>

Arguments

<arg></arg>	CBWR compatibility mode	Description
0	None	Do not use CBWR in a library-wide mode. CNR-safe communicators may be created with
		MPI_Comm_dup_with_info explicitly. This is the default value.

1	Weak mode	Disable topology aware collectives. The result of a collective operation does not depend on the rank placement. The mode guarantees results reproducibility across different runs on the same cluster (independent of the rank placement).
2	Strict mode	Disable topology aware collectives, ignore CPU architecture, and interconnect during algorithm selection. The mode guarantees results reproducibility across different runs on different clusters (independent of the rank placement, CPU architecture, and interconnection)

Description

Conditional Numerical Reproducibility (CNR) provides controls for obtaining reproducible floating-point results on collectives operations. With this feature, Intel MPI collective operations are designed to return the same floating-point results from run to run in case of the same number of MPI ranks.

Control this feature with the <code>I_MPI_CBWR</code> environment variable in a library-wide manner, where all collectives on all communicators are guaranteed to have reproducible results. To control the floating-point operations reproducibility in a more precise and per-communicator way, pass the $\{"I_MPI_CBWR", "yes"\}\$ key-value pair to the MPI Comm dup with info call.

NOTE

Setting the I_MPI_CBWR in a library-wide mode using the environment variable leads to performance penalty.

CNR-safe communicators created using $MPI_Comm_dup_with_info$ always work in the strict mode. For example:

```
MPI_Info hint;
MPI_Comm cbwr_safe_world, cbwr_safe_copy;
MPI_Info_create(&hint);
MPI_Info_set(hint, "I_MPI_CBWR", "yes");
MPI_Comm_dup_with_info(MPI_COMM_WORLD, hint, & cbwr_safe_world);
MPI_Comm_dup(cbwr_safe_world, & cbwr_safe_copy);
```

In the example above, both cbwr_safe_world and cbwr_safe_copy are CNR-safe. Use cbwr_safe_world and its duplicates to get reproducible results for critical operations.

Note that MPI_COMM_WORLD itself may be used for performance-critical operations without reproducibility limitations.

Tuning Environment Variables

I MPI TUNING MODE

Select the tuning method.

Syntax

I_MPI_TUNING_MODE=<arg>

Arguments

<arg></arg>	Description
none	Disable tuning modes. This is the default value.
auto	Enable autotuner.
auto:application Enable autotuner with application focused strategy (alias for auto).	
auto:cluster	Enable autotuner without application specific logic. This is typically performed with the help of benchmarks (for example, IMB-MPI1) and proxy applications.

Description

Set this environment variable to enable the autotuner functionality and set the autotuner strategy.

I_MPI_TUNING_BIN

Specify the path to tuning settings in a binary format.

Syntax

I MPI TUNING BIN=<path>

Argument

<path></path>	A path to a binary file with tuning settings. By default, Intel® MPI Library uses
	the binary tuning file located at <\$I_MPI_ONEAPI_ROOT/etc>.

Description

Set this environment variable to load tuning settings in a binary format.

I_MPI_TUNING_BIN_DUMP

Specify the file for storing tuning settings in a binary format.

Syntax

I MPI TUNING BIN DUMP=<filename>

Argument

<filename></filename>	A file name of a binary that stores tuning settings. By default, the path is not
	specified.

Description

Set this environment variable to store tuning settings in binary format.

I_MPI_TUNING

Load tuning settings in a JSON format.

Syntax

I_MPI_TUNING=<path>

Argument

<path></path>	A path to a JSON file with tuning settings.	
---------------	---	--

Description

Set this environment variable to load tuning settings in a JSON format.

NOTE The tuning settings in the JSON format are produced by the mpitune utility.

By default, the Intel® MPI Library loads tuning settings in a binary format. If it is not possible, the Intel MPI Library loads the tuning file in a JSON format specified through the <code>I_MPI_TUNING</code> environment variable. Thus, to enable JSON tuning, turn off the default binary tuning: <code>I_MPI_TUNING_BIN=""</code>. If it is not possible to load tuning settings from a JSON file and in a binary format, the default tuning values are used.

You do not need to turn off binary or JSON tuning settings if you use I_MPI_ADJUST family environment variables. The algorithms specified with I_MPI_ADJUST environment variables always have priority over binary and JSON tuning settings.

See Also

- Autotuning
- Environment Variables for Autotuning

Autotuning

Tuning is very dependent on the specifications of the particular platform. Intel carefully determines the tuning parameters, and makes them available for autotuning using $I_MPI_TUNING_MODE$ and the $I_MPI_TUNING_AUTO$ family environment variables to find the best settings (see Tuning Environment Variables and $I_MPI_TUNING_AUTO$ Family Environment Variables).

NOTEI_MPI_TUNING_MODE and the I_MPI_TUNING_AUTO family environment variables support only Intel processors, and cannot be used on other platforms.

The autotuner functionality lets you automatically find the best algorithms for collective operations. The autotuner search space can be modified by the $I_MPI_ADJUST_<opname>_LIST$ variable (see I_MPI_ADJUST Family Environment Variables).

The collectives currently available for autotuning are: MPI_Allreduce, MPI_Bcast, MPI_Barrier, MPI_Reduce, MPI_Gather, MPI_Scatter, MPI_Alltoall, MPI_Allgatherv, MPI_Reduce_scatter, MPI_Reduce_scatter_block, MPI_Scan, MPI_Exscan, MPI_Iallreduce, MPI_Ibcast, MPI_Ibarrier, MPI_Ireduce, MPI_Igather, MPI_Iscatter, MPI_Ialltoall, MPI_Iallgatherv, MPI_Ireduce_scatter, MPI_Ireduce_scatter_block, MPI_Iscan, and MPI_Iexscan.

To get started with autotuning, follow these steps:

1. Launch the application with autotuner enabled and specify the dump file, which stores results:

```
I_MPI_TUNING_MODE=auto
I MPI TUNING BIN DUMP=<tuning-results.dat>
```

2. Launch the application with the tuning results generated at the previous step:

```
I MPI TUNING BIN=<tuning-results.dat>
```

Or use the -tune Hydra option.

3. If you experience performance issues, see Environment Variables for Autotuning.

For example:

```
1$ export I_MPI_TUNING_MODE=auto
$ export I_MPI_TUNING_AUTO_SYNC=1
$ export I_MPI_TUNING_AUTO_ITER_NUM=5
$ export I_MPI_TUNING_BIN_DUMP=./tuning_results.dat
$ mpirun -n 128 -ppn 64 IMB-MPI1 allreduce -iter 1000,800 -time 4800
2$ export I_MPI_TUNING_BIN=./tuning_results.dat
$ mpirun -n 128 -ppn 64 IMB-MPI1 allreduce -iter 1000,800 -time 4800
```

NOTE To tune collectives on a communicator identified with the help of Application Performance Snapshot (APS), execute the following variable at step 1:

I MPI TUNING AUTO COMM LIST=comm id 1, ..., comm id n.

See Also

Environment Variables for Autotuning

mpitune_fast

I MPI TUNING AUTO Family Environment Variables

NOTE You must set I_MPI_TUNING_MODE to use any of the I_MPI_TUNING_AUTO family environment variables.

NOTE The I_MPI_TUNING_AUTO family environment variables support only Intel processors, and cannot be used on other platforms.

I_MPI_TUNING_AUTO_STORAGE_SIZE

Define size of the per-communicator tuning storage.

Syntax

I MPI TUNING AUTO STORAGE SIZE=<size>

Argument

<size></size>	Specify size of the communicator tuning storage. The
	default size of the storage is 512 Kb.

Description

Set this environment variable to change the size of the communicator tuning storage.

I_MPI_TUNING_AUTO_ITER_NUM

Specify the number of autotuner iterations.

Syntax

I MPI TUNING AUTO ITER NUM=<number>

Argument

Γ.	<number></number>	Define the number of iterations. By default, it is 1.

Description

Set this environment variable to specify the number of autotuner iterations. The greater iteration number produces more accurate results.

NOTE To check if all possible algorithms are iterated, make sure that the total number of collective invocations for a particular message size in a target application is at least equal the value of <code>I_MPI_TUNING_AUTO_ITER_NUM</code> multiplied by the number of algorithms.

I MPI TUNING AUTO WARMUP ITER NUM

Specify the number of warmup autotuner iterations.

Syntax

I_MPI_TUNING_AUTO_WARMUP_ITER_NUM=<number>

Argument

<number></number>	Define the number of iterations. By default, it is 1.
	, ,

Description

Set this environment variable to specify the number of autotuner warmup iterations. Warmup iterations do not impact autotuner decisions and allow to skip additional iterations, such as infrastructure preparation.

I_MPI_TUNING_AUTO_SYNC

Enable the internal barrier on every iteration of the autotuner.

Syntax

I_MPI_TUNING_AUTO_SYNC=<arg>

Argument

<arg></arg>	Binary indicator
enable yes on 1	Align the autotuner with the IMB measurement approach.
disable no off 0	Do not use the barrier on every iteration of the autotuner.
	This is the default value.

Description

Set this environment variable to control the IMB measurement logic. Setting this variable to 1 may lead to overhead due to an additional MPI_Barrier call.

I_MPI_TUNING_AUTO_COMM_LIST

Control the scope of autotuning.

Syntax

```
I_MPI_TUNING_AUTO_COMM_LIST=<comm_id_1, ..., comm_id_n>
```

Argument

<pre><comm id="" n,=""></comm></pre>	Specify communicators to be tuned.
/ _ / /	,

Description

Set this environment variable to specify communicators to be tuned using their unique id. By default, the variable is not specified. In this case, all communicators in the application are involved into the tuning process.

NOTE To get the list of communicators available for tuning, use the Application Performance Snapshot (APS) tool, which supports per communicator profiling starting with the 2019 Update 4 release. For example:

1. Source apsvars.sh:

```
$ source <path to aps>/apsvars.sh
```

2. Gather APS statistics:

```
$ export MPS_STAT_LEVEL=5
$ export APS_COLLECT_COMM_IDS=1
mpirun -aps -n 128 -ppn 64 IMB-MPI1 allreduce -npmin 128 -iter 1000,800 -time 4800
```

3. Generate an APS report:

```
$ aps-report aps result 20190228/ -1FE
```

4. Get the results:

5. Specify the communicators to be tuned:

```
$ export I_MPI_TUNING_AUTO_COMM_LIST=4611686018431582688
$ export MPS_STAT_LEVEL=5
$ export APS_COLLECT_COMM_IDS=1
$ export I_MPI_TUNING_AUTO=1
$ mpirun -aps -n 128 -ppn 64 IMB-MPI1 allreduce -iter 1000,800 -time 4800
```

I_MPI_TUNING_AUTO_COMM_DEFAULT

Mark all communicators with the default value.

Syntax

```
I_MPI_TUNING_AUTO_COMM_DEFAULT=<arg>
```

Argument

<arg></arg>	Binary indicator
enable yes on 1	Mark communicators.
disable no off 0	Do not mark communicators. This is the default value.

Description

Set this environment variable to mark all communicators in an application with the default value. In this case, all communicators will have the identical default comm_id equal to -1.

2

I MPI TUNING AUTO COMM USER

Enable communicator marking with a user value.

Syntax

I MPI TUNING AUTO COMM USER=<arg>

Argument

<arg></arg>	Binary indicator
enable yes on 1	Enable marking of communicators.
disable no off 0	Disable marking of communicators. This is the default value.

Description

Set this environment variable to enable communicator marking with a user value. To mark a communicator in your application, use the MPI_Info object for this communicator that contains a record with the comm_id key. The key must belong the 0...UINT64 MAX range.

I_MPI_TUNING_AUTO_ITER_POLICY

Control the iteration policy logic.

Syntax

MPI TUNING AUTO ITER POLICY=<arg>

Argument

<arg></arg>	Binary indicator
enable yes on 1	Reduce the number of iterations with a message size increase after 64Kb (by half). This is the default value.
disable no off 0	Use the <pre>I_MPI_TUNING_AUTO_ITER_NUM</pre> value.
	This value affects warmup iterations.

Description

Set this environment variable to control the autotuning iteration policy logic.

I_MPI_TUNING_AUTO_ITER_POLICY_THRESHOLD

Control the message size limit for the I MPI TUNING AUTO ITER POLICY environment variable.

Syntax

I MPI TUNING AUTO ITER POLICY THRESHOLD=<arg>

Argument

<arg></arg>	Define the value. By default, it is 64KB.

Description

Set this environment variable to control the message size limit for the autotuning iteration policy logic ($I_MPI_TUNING_AUTO_ITER_POLICY$).

I MPI TUNING AUTO POLICY

Choose the best algorithm identification strategy.

Syntax

I MPI TUNING AUTO POLICY=<arg>

Argument

<arg></arg>	Description
max	Choose the best algorithm based on a maximum time value. This is the default value.

min	Choose the best algorithm based on a minimum time value.
avg	Choose the best algorithm based on an average time value.

Description

Set this environment variable to control the autotuning strategy and choose the best algorithm based on the time value across ranks involved into the tuning process.

Process Pinning

Use this feature to pin a particular MPI process to a corresponding set of CPUs within a node and avoid undesired process migration. This feature is available on operating systems that provide the necessary kernel interfaces.

This page describes the pinning process. You can simulate your pinning configuration using the Pinning Simulator for Intel MPI Library.

Processor Identification

The following schemes are used to identify logical processors in a system:

- System-defined logical enumeration
- Topological enumeration based on three-level hierarchical identification through triplets (package/socket, core, thread)

The number of a logical CPU is defined as the corresponding position of this CPU bit in the kernel affinity bit-mask. Use the <code>cpuinfo</code> utility, provided with your Intel MPI Library installation or the <code>cat /proc/cpuinfo</code> command to find out the logical CPU numbers.

The three-level hierarchical identification uses triplets that provide information about processor location and their order. The triplets are hierarchically ordered (package, core, and thread).

See the example for one possible processor numbering where there are two sockets, four cores (two cores per socket), and eight logical processors (two processors per core).

NOTE Logical and topological enumerations are not the same.

Logical Enumeration

0	4	1	5		2	6	3	7	
Hierarchi	cal Levels	5							
Socket	0	0	0	0	1	1	1	1	
Core	0	0	1	1	0	0	1	1	ĺ
Thread	0	1	0	1	0	1	0	1	
Topological Enumeration									
0	1	2	3		4	5	6	7	

Use the <code>cpuinfo</code> utility to identify the correspondence between the logical and topological enumerations. See Processor Information Utility for more details.

Default Settings

If you do not specify values for any process pinning environment variables, the default settings below are used. For details about these settings, see Environment Variables and Interoperability with OpenMP API.

- I MPI PIN=on
- I MPI PIN RESPECT CPUSET=on

- I MPI PIN RESPECT HCA=on
- I MPI PIN CELL=unit
- I MPI PIN DOMAIN=auto:compact
- I MPI PIN ORDER=bunch

NOTE If hyperthreading is on, the number or processes on the node is greater than the number of cores and no one process pinning environment variable is set. For better performance, the "spread" order will automatically be used instead of the default "compact" order.

Environment Variables for Process Pinning

I_MPI_PIN

Turn on/off process pinning.

Syntax

```
I MPI PIN=<arg>
```

Arguments

Description

Set this environment variable to control the process pinning feature of the Intel® MPI Library.

I MPI PIN PROCESSOR LIST (I MPI PIN PROCS)

Define a processor subset and the mapping rules for MPI processes within this subset.

Syntax Forms

```
I MPI PIN PROCESSOR LIST=<value>
```

The environment variable value has three syntax forms:

- 1. clist>
- 2. [cset>][:[grain=<grain>][,shift=<shift>][,preoffset=<preoffset>]
 [,postoffset=<postoffset>]
- **3.** [< procset >] [:map = < map >]

The following paragraphs provide detailed descriptions for the values of these syntax forms.

NOTE The postoffset keyword has offset alias.

NOTE The second form of the pinning procedure has three steps:

- 1. Cyclic shift of the source processor list on preoffset*grain value.
- 2. Round robin shift of the list derived on the first step on shift*grain value.
- **3.** Cyclic shift of the list derived on the second step on the postoffset*grain value.

NOTE The grain, shift, preoffset, and postoffset parameters have a unified definition style.

This environment variable is available for both Intel and non-Intel microprocessors, but it may perform additional optimizations for Intel microprocessors than it performs for non-Intel microprocessors.

Syntax 1

I MPI PIN PROCESSOR LIST=proclist>

Arguments

<pre><pre><pre><pre></pre></pre></pre></pre>	A comma-separated list of logical processor numbers and/or ranges of processors. The process with the i-th rank is pinned to the i-th processor in the list. The number should not exceed the number of processors on a node.
<1>	Processor with logical number <1>.
<1>- <m></m>	Range of processors with logical numbers from $<1>$ to $$.
<k>,<1>-<m></m></k>	Processors $\langle k \rangle$, as well as $\langle 1 \rangle$ through $\langle m \rangle$.

Syntax 2

I_MPI_PIN_PROCESSOR_LIST=[procset>][:[grain=<grain>][,shift=<shift>]
[,preoffset=<preoffset>][,postoffset=<postoffset>]

Arguments

<pre><pre><pre><pre></pre></pre></pre></pre>	Specify a processor subset based on the topological numeration. The default value is allcores.
all	All logical processors. Specify this subset to define the number of CPUs on a node.
allcores	All cores (physical CPUs). Specify this subset to define the number of cores on a node. This is the default value.
	If Intel® Hyper-Threading Technology is disabled, allcores equals to all.
allsocks	All packages/sockets. Specify this subset to define the number of sockets on a node.
<grain></grain>	Specify the pinning granularity cell for a defined <pre>procset></pre> . The minimal <pre>cgrain>value</pre> is a single element of the <pre>procset></pre> . The maximal <pre>cgrain></pre> value is the number of <pre>procset></pre> elements in a socket. The <pre>cgrain>value</pre> must be a multiple of the <pre>procset></pre> value. Otherwise, the minimal <pre>cgrain></pre> value is assumed. The default value is the minimal <pre>cgrain></pre> value.
<shift></shift>	Specify the granularity of the round robin scheduling shift of the cells for the $$. $$ is measured in the defined $$ units. The $$ value must be positive integer. Otherwise, no shift is performed. The default value is no shift, which is equal to 1 normal increment.
<pre><pre><pre><pre></pre></pre></pre></pre>	Specify the cyclic shift of the processor subset $\langle procset \rangle$ defined before the round robin shifting on the $\langle preoffset \rangle$ value. The value is measured in the defined $\langle grain \rangle$ units. The $\langle preoffset \rangle$ value must be non-negative integer. Otherwise, no shift is performed. The default value is no shift.
<postoffset></postoffset>	Specify the cyclic shift of the processor subset $$ derived after round robin shifting on the $$ value. The value is measured in the defined $$ units. The $$ value must be non-negative integer. Otherwise no shift is performed. The default value is no shift.

The following table displays the values for <grain>, <shift>, <preoffset>, and <postoffset> options:

<n:< th=""><th>></th><th>Specify an explicit value of the corresponding parameters. <n>is non-negative</n></th></n:<>	>	Specify an explicit value of the corresponding parameters. <n>is non-negative</n>
		integer.
fir	ne	Specify the minimal value of the corresponding parameter.

core	Specify the parameter value equal to the amount of the corresponding parameter units contained in one core.
cache1	Specify the parameter value equal to the amount of the corresponding parameter units that share an L1 cache.
cache2	Specify the parameter value equal to the amount of the corresponding parameter units that share an L2 cache.
cache3	Specify the parameter value equal to the amount of the corresponding parameter units that share an L3 cache.
cache	The largest value among cache1, cache2, and cache3.
socket sock	Specify the parameter value equal to the amount of the corresponding parameter units contained in one physical package/socket.
half mid	Specify the parameter value equal to socket/2.
third	Specify the parameter value equal to socket/3.
quarter	Specify the parameter value equal to socket/4.
octavo	Specify the parameter value equal to socket/8.

Syntax 3

I MPI PIN PROCESSOR LIST=[procset>][:map=<map>]

Arguments

<map></map>	The mapping pattern used for process placement.
bunch	The processes are mapped as close as possible on the sockets.
scatter	The processes are mapped as remotely as possible so as not to share common resources: FSB, caches, and core.
spread	The processes are mapped consecutively with the possibility not to share common resources.

Description

Set the I_MPI_PIN_PROCESSOR_LIST environment variable to define the processor placement. To avoid conflicts with different shell versions, the environment variable value may need to be enclosed in quotes.

NOTE This environment variable is valid only if <code>I_MPI_PIN</code> is enabled.

The <code>I_MPI_PIN_PROCESSOR_LIST</code> environment variable has the following different syntax variants:

• Explicit processor list. This comma-separated list is defined in terms of logical processor numbers. The relative node rank of a process is an index to the processor list such that the i-th process is pinned on i-th list member. This permits the definition of any process placement on the CPUs.

For example, process mapping for I MPI PIN PROCESSOR LIST=p0,p1,p2,...,pn is as follows:

Rank on a node	0	1	2		n-1	N	
Logical CPU	0 q	p1	p2	***	pn-1	Pn	

• grain/shift/offset mapping. This method provides cyclic shift of a defined grain along the processor list with steps equal to shift*grain and a single shift on offset*grain at the end. This shifting action is repeated shift times.

For example: grain = 2 logical processors, shift = 3 grains, offset = 0.

Legend:

gray - MPI process grains

- A) red processor grains chosen on the 1st pass
- B) cyan processor grains chosen on the 2nd pass
- C) green processor grains chosen on the final 3rd pass
- D) Final map table ordered by MPI ranks

A)

0 1			2 3				2n-2 2n-1		
0 1	2 3	4 5	6 7	8 9	10 11		6n-6 6n-5	6n-4 6n-3	6n-2 6n-1
B)									
0 1	2n 2n+1		2 3	2n+2 2n+3			2n-2 2n-1	4n-2 4n-1	
0 1	2 3	4 5	6 7	8 9	10 11		6n-6 6n-5	6n-4 6n-3	6n-2 6n-1
C)									
0 1	2n 2n+1	4n 4n+1	2 3	2n+2 2n +3	4n+2 4r +3	١	2n-2 2n-1	4n-2 4n-1	6n-2 6n-1
0 1	2 3	4 5	6 7	8 9	10 11		6n-6 6n-5	6n-4 6n-3	6n-2 6n-1
D)									-
0 1	2 3	2n- 2n-		n 2n 2n+2 1 2n+3		4n-2 4n-1	4n 4n +1	4n+2 4n+3	6n-2 6n-1
0 1	6 7	6n- 6n-		3 89		6n-4 6n-3	4 5	10 11	6n-2 6n-1

 Predefined mapping scenario. In this case, popular process pinning schemes are defined as keywords selectable at runtime. There are two such scenarios: bunch and scatter.

In the bunch scenario the processes are mapped proportionally to sockets as closely as possible. This mapping makes sense for partial processor loading. In this case, the number of processes is less than the number of processors.

In the scatter scenario the processes are mapped as remotely as possible so as not to share common resources: FSB, caches, and cores.

In the example, there are two sockets, four cores per socket, one logical CPU per core, and two cores per shared cache.

Legend:

gray - MPI processes

cyan - 1st socket processors

green - 2nd socket processors

Same color defines a processor pair sharing a cache.

0	1	2		3	4			
0	1	2	3	4	5	6	7	l
bunch S	cenario for 5	processes						
0	4	2	6	1	5	3	7	
0	1	2	3	4	5	6	7	Ì

scatter scenario for full loading

Examples

To pin the processes to CPU0 and CPU3 on each node globally, use the following command:

\$ mpirun -genv I MPI PIN PROCESSOR LIST=0,3 -n <number-of-processes><executable>

To pin the processes to different CPUs on each node individually (CPU0 and CPU3 on host1 and CPU0, CPU1 and CPU3 on host2), use the following command:

\$ mpirun -host host1 -env I_MPI_PIN_PROCESSOR_LIST=0,3 -n <number-of-processes> <executable> : \
-host host2 -env I_MPI_PIN_PROCESSOR_LIST=1,2,3 -n <number-of-processes> <executable>

To print extra debugging information about process pinning, use the following command:

```
$ mpirun -genv I_MPI_DEBUG=4 -m -host host1 \
-env I_MPI_PIN_PROCESSOR_LIST=0,3 -n <number-of-processes> <executable> :\
-host host2 -env I_MPI_PIN_PROCESSOR_LIST=1,2,3 -n <number-of-processes> <executable>
```

NOTE If the number of processes is greater than the number of CPUs used for pinning, the process list is wrapped around to the start of the processor list.

I_MPI_PIN_PROCESSOR_EXCLUDE_LIST

Define a subset of logical processors to be excluded for the pinning capability on the intended hosts.

Syntax

```
I MPI PIN PROCESSOR EXCLUDE LIST=proclist>
```

Arguments

<pre><pre><pre><pre></pre></pre></pre></pre>	A comma-separated list of logical processor numbers and/or ranges of processors.
	Processor with logical number $<1>$.
- <m></m>	Range of processors with logical numbers from $<1>$ to $$.
<k>,<l>-<m></m></l></k>	Processors $\langle k \rangle$, as well as $\langle 1 \rangle$ through $\langle m \rangle$.

Description

Set this environment variable to define the logical processors that Intel® MPI Library does not use for pinning capability on the intended hosts. Logical processors are numbered as in /proc/cpuinfo.

I_MPI_PIN_CELL

Set this environment variable to define the pinning resolution granularity. I_MPI_PIN_CELL specifies the minimal processor cell allocated when an MPI process is running.

Syntax

```
I MPI PIN CELL=<cell>
```

Arguments

<cell></cell>	Specify the resolution granularity
unit	Basic processor unit (logical CPU)
core	Physical processor core

Description

Set this environment variable to define the processor subset used when a process is running. You can choose from two scenarios:

- all possible CPUs in a node (unit value)
- all cores in a node (core value)

The environment variable has effect on both pinning types:

- one-to-one pinning through the I MPI PIN PROCESSOR LIST environment variable
- one-to-many pinning through the I MPI PIN DOMAIN environment variable

The default value rules are:

- If you use I MPI PIN DOMAIN, the cell granularity is unit.
- If you use I MPI PIN PROCESSOR LIST, the following rules apply:

- When the number of processes is greater than the number of cores, the cell granularity is unit.
- When the number of processes is equal to or less than the number of cores, the cell granularity is core.

NOTE The core value is not affected by the enabling/disabling of Intel® Hyper-Threading Technology in a system.

I_MPI_PIN_RESPECT_CPUSET

Respect the process affinity mask.

Syntax

I MPI PIN RESPECT CPUSET=<value>

Arguments

<value></value>	Binary indicator
enable yes on	Respect the process affinity mask. This is the default value.
1	
disable no off	Do not respect the process affinity mask.
0	

Description

If you set <code>I_MPI_PIN_RESPECT_CPUSET=enable</code>, the Hydra process launcher uses job manager's process affinity mask on each intended host to determine logical processors for applying Intel MPI Library pinning capability.

If you set <code>I_MPI_PIN_RESPECT_CPUSET=disable</code>, the Hydra process launcher uses its own process affinity mask to determine logical processors for applying Intel MPI Library pinning capability.

I MPI PIN RESPECT HCA

In the presence of Infiniband architecture* host channel adapter (IBA* HCA), adjust the pinning according to the location of IBA HCA.

Syntax

I MPI PIN RESPECT HCA=<value>

Arguments

<value></value>	Binary indicator
enable yes on	Use the location of IBA HCA if available. This is the default value.
1	
disable no off	Do not use the location of IBA HCA.
0	

Description

If you set $I_MPI_PIN_RESPECT_HCA=enable$, the Hydra process launcher uses the location of IBA HCA on each intended host for applying Intel MPI Library pinning capability.

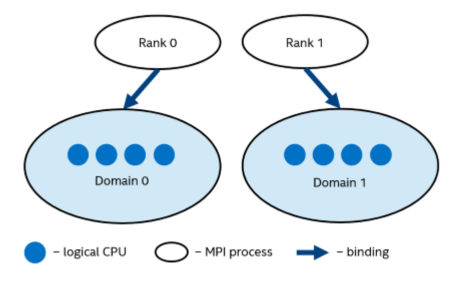
If you set $I_MPI_PIN_RESPECT_HCA=disable$, the Hydra process launcher does not use the location of IBA HCA on each intended host for applying Intel MPI Library pinning capability.

Interoperability with OpenMP* API

I_MPI_PIN_DOMAIN

Intel® MPI Library provides an additional environment variable to control process pinning for hybrid MPI/ OpenMP* applications. This environment variable is used to define a number of non-overlapping subsets (domains) of logical processors on a node, and a set of rules on how MPI processes are bound to these domains by the following formula: *one MPI process per one domain*. See the picture below.

Figure 1 Domain Example



Each MPI process can create a number of children threads for running within the corresponding domain. The process threads can freely migrate from one logical processor to another within the particular domain.

If the <code>I_MPI_PIN_DOMAIN</code> environment variable is defined, then the <code>I_MPI_PIN_PROCESSOR_LIST</code> environment variable setting is ignored.

If the $I_MPI_PIN_DOMAIN$ environment variable is not defined, then MPI processes are pinned according to the current value of the $I_MPI_PIN_PROCESSOR_LIST$ environment variable.

The I MPI PIN DOMAIN environment variable has the following syntax forms:

- Domain description through multi-core terms <mc-shape>
- Domain description through domain size and domain member layout <size>[:<layout>]
- Explicit domain description through bit mask <masklist>

The following tables describe these syntax forms.

Multi-Core Shape

I MPI PIN DOMAIN=<mc-shape>

<mc-shape></mc-shape>	Define domains through multi-core terms.
core	Each domain consists of the logical processors that share a particular core. The number of domains on a node is equal to the number of cores on the node.
socket sock	Each domain consists of the logical processors that share a particular socket. The number of domains on a node is equal to the number of sockets on the node. This is the recommended value.

numa	Each domain consists of the logical processors that share a particular NUMA node. The number of domains on a machine is equal to the number of NUMA nodes on the machine.
node	All logical processors on a node are arranged into a single domain.
cache1	Logical processors that share a particular level 1 cache are arranged into a single domain.
cache2	Logical processors that share a particular level 2 cache are arranged into a single domain.
cache3	Logical processors that share a particular level 3 cache are arranged into a single domain.
cache	The largest domain among cache1, cache2, and cache3 is selected.

NOTE If <code>Cluster</code> on <code>Die</code> is disabled on a machine, the number of NUMA nodes equals to the number of sockets. In this case, pinning for <code>I_MPI_PIN_DOMAIN</code> = <code>numa</code> is equivalent to pinning for <code>I_MPI_PIN_DOMAIN</code> = <code>socket</code>.

Explicit Shape

I_MPI_PIN_DOMAIN=<size>[:<layout>]

<size></size>	Define a number of logical processors in each domain (domain size)
omp	The domain size is equal to the <code>OMP_NUM_THREADS</code> environment variable
	value. If the OMP_NUM_THREADS environment variable is not set, each node
	is treated as a separate domain.
auto	The domain size is defined by the formula size=#cpu/#proc, where #cpu
	is the number of logical processors on a node, and #proc is the number of
	the MPI processes started on a node
<n></n>	The domain size is defined by a positive decimal number $\langle n \rangle$
<layout></layout>	Ordering of domain members. The default value is compact
<pre> platform</pre>	Ordering of domain members. The default value is compact Domain members are ordered according to their BIOS numbering (platform-depended numbering)
_	Domain members are ordered according to their BIOS numbering (platform-

Explicit Domain Mask

I_MPI_PIN_DOMAIN=<masklist>

<masklist></masklist>	Define domains through the comma separated list of hexadecimal numbers (domain masks)
[m ₁ ,,m _n]	For $< masklist >$, each m_i is a hexadecimal bit mask defining an individual domain. The following rule is used: the i^{th} logical processor is included into the domain if the corresponding mi value is set to 1. All remaining processors are put into a separate domain. BIOS numbering is used.

NOTE To ensure that your configuration in <masklist> is parsed correctly, use square brackets to enclose the domains specified by the <masklist>. For example: $I_MPI_PIN_DOMAIN = [55, aa]$

NOTE These options are available for both Intel® and non-Intel microprocessors, but they may perform additional optimizations for Intel microprocessors than they perform for non-Intel microprocessors.

To pin OpenMP* processes or threads inside the domain, the corresponding OpenMP feature (for example, the KMP AFFINITY environment variable for Intel® compilers) should be used.

NOTE The following configurations are effectively the same as if pinning is not applied:

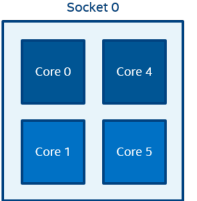
- I MPI PIN DOMAIN=node

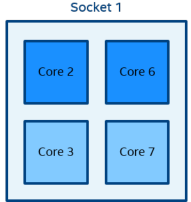
If you do not want the process to be migrated between sockets on a multi-socket platform, specify the domain size as $I_MPI_PIN_DOMAIN=socket$ or smaller.

You can also use I_MPI_PIN_PROCESSOR_LIST, which produces a single-cpu process affinity mask for each rank (the affinity mask is supposed to be automatically adjusted in presence of IBA* HCA).

See the following model of a symmetric multiprocessing (SMP) node in the examples:

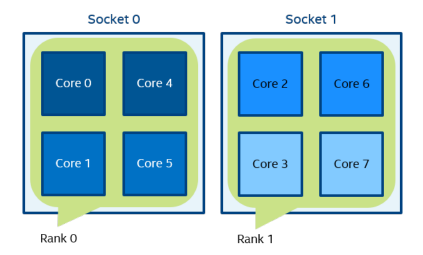
Figure 2 Model of a Node





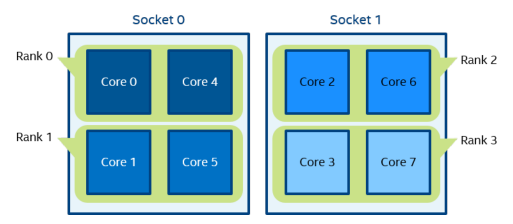
The figure above represents the SMP node model with a total of 8 cores on 2 sockets. Intel® Hyper-Threading Technology is disabled. Core pairs of the same color share the L2 cache.

Figure 3 mpi run -n 2 -env I_MPI_PIN_DOMAIN socket ./a.out



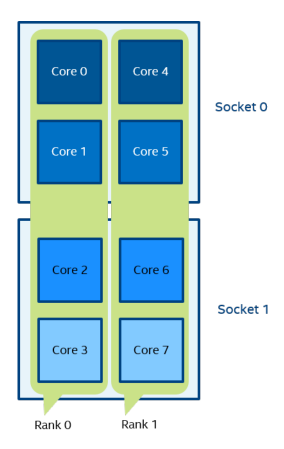
In Figure 3, two domains are defined according to the number of sockets. Process rank 0 can migrate on all cores on the 0-th socket. Process rank 1 can migrate on all cores on the first socket.

Figure 4 mpi run -n 4 -env I MPI PIN DOMAIN cache2 ./a.out



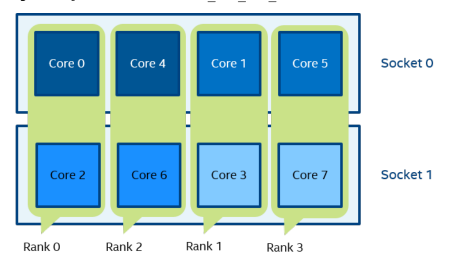
In Figure 4, four domains are defined according to the amount of common L2 caches. Process rank 0 runs on cores $\{0,4\}$ that share an L2 cache. Process rank 1 runs on cores $\{1,5\}$ that share an L2 cache as well, and so on.

Figure 5 mpi run -n 2 -env I MPI PIN DOMAIN 4:platform ./a.out



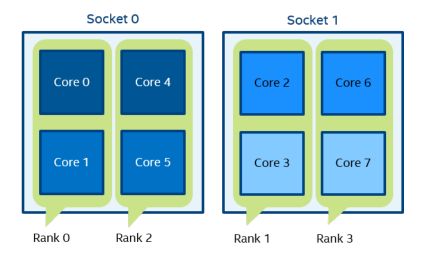
In Figure 5, two domains with size=4 are defined. The first domain contains cores $\{0,1,2,3\}$, and the second domain contains cores $\{4,5,6,7\}$. Domain members (cores) have consecutive numbering as defined by the platform option.

Figure 6 mpi run -n 4 -env I MPI PIN DOMAIN auto:scatter ./a.out



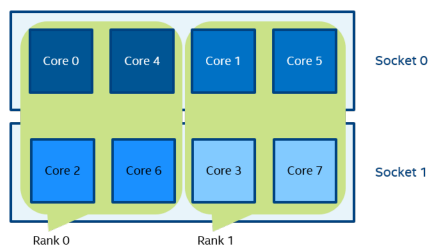
In Figure 6, domain size=2 (defined by the number of CPUs=8 / number of processes=4), scatter layout. Four domains $\{0,2\}$, $\{1,3\}$, $\{4,6\}$, $\{5,7\}$ are defined. Domain members do not share any common resources.

 $\textit{Figure 7} \textit{ setenv OMP_NUM_THREADS=2 mpi run -n 4 -env I_MPI_PIN_DOMAIN omp:platform ./a.out } \\$



In Figure 7, domain size=2 (defined by $OMP_NUM_THREADS=2$), platform layout. Four domains {0,1}, {2,3}, {4,5}, {6,7} are defined. Domain members (cores) have consecutive numbering.

Figure 8 mpi run -n 2 -env I_MPI_PIN_DOMAIN [55,aa] ./a.out



In Figure 8 (the example for <code>I_MPI_PIN_DOMAIN=<masklist></code>), the first domain is defined by the 55 mask. It contains all cores with even numbers $\{0,2,4,6\}$. The second domain is defined by the AA mask. It contains all cores with odd numbers $\{1,3,5,7\}$.

I_MPI_PIN_ORDER

Set this environment variable to define the mapping order for MPI processes to domains as specified by the $I_MPI_PIN_DOMAIN$ environment variable.

Syntax

I MPI PIN ORDER=<order>

Arguments

<order></order>	Specify the ranking order
range	The domains are ordered according to the processor's BIOS numbering. This is a platform-dependent numbering.
scatter	The domains are ordered so that adjacent domains have minimal sharing of common resources, whenever possible.

compact	The domains are ordered so that adjacent domains share common resources as much as possible.
spread	The domains are ordered consecutively with the possibility not to share common resources.
bunch	The processes are mapped proportionally to sockets and the domains are ordered as close as possible on the sockets. This is the default value.

The optimal setting for this environment variable is application-specific. If adjacent MPI processes prefer to share common resources, such as cores, caches, sockets, FSB, use the compact or bunch values. Otherwise, use the scatter or spread values. Use the range value as needed. For detail information and examples about these values, see the Arguments table and the Example section of I MPI PIN ORDER in this topic.

The options scatter, compact, spread and bunch are available for both Intel® and non-Intel microprocessors, but they may perform additional optimizations for Intel microprocessors than they perform for non-Intel microprocessors.

Examples

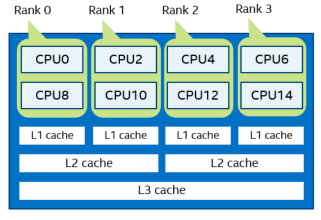
For the following configuration:

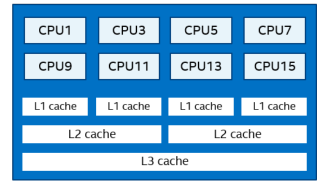
- Two socket nodes with four cores and a shared L2 cache for corresponding core pairs.
- 4 MPI processes you want to run on the node using the settings below.

Compact order:

I MPI PIN DOMAIN=2 I MPI PIN ORDER=compact

Figure 9 Compact Order Example

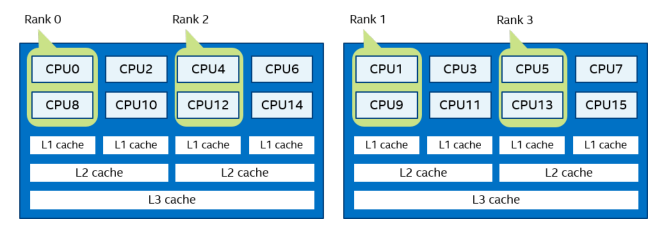




Scatter order:

I MPI PIN DOMAIN=2 I MPI PIN ORDER=scatter

Figure 10 Scatter Order Example

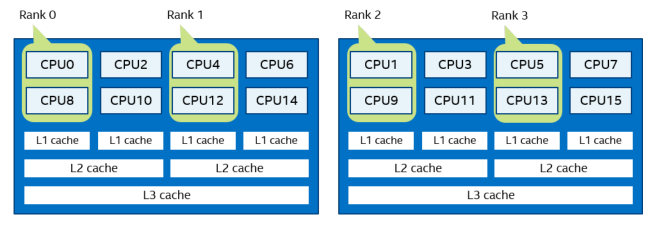


Spread order:

I MPI PIN DOMAIN=2 I MPI PIN ORDER=spread

NOTE For I_MPI_PIN_ORDER=spread, the order will be switched to 'compact' if there are not enough CPUs to emplace all domains.

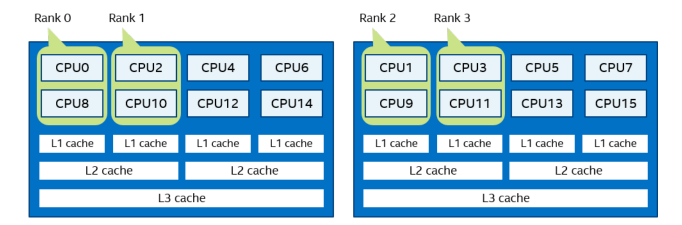
Figure 11 Spread Order Example



Bunch order:

I_MPI_PIN_DOMAIN=2 I_MPI_PIN_ORDER=bunch

Figure 12 Bunch Order Example



GPU Support

In addition to GPU pinning, the Intel MPI Library supports GPU buffers (see below).

GPU Pinning

Use this feature to distribute Intel GPU devices between MPI ranks.

To enable this feature, set <code>I_MPI_OFFLOAD_TOPOLIB=level_zero</code>. This feature requires that the Level-Zero* library be installed on the nodes. The device pinning information is printed out in the Intel MPI debug output at <code>I_MPI_DEBUG=3</code>.

Default settings:

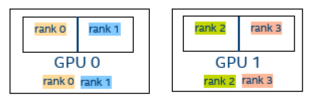
```
I_MPI_OFFLOAD_CELL=tile
I_MPI_OFFLOAD_DOMAIN_SIZE=-1
I_MPI_OFFLOAD_DEVICES=all
```

By default, all available resources are distributed between MPI ranks as equally as possible given the position of the ranks; that is, the distribution of resources takes into account on which NUMA node the rank and the resource are located. Ideally, the rank will have resources only on the same NUMA node on which the rank is located.

Examples

All examples below represent a machine configuration with two NUMA nodes and two GPUs with two tiles.

Figure 1. Four MPI Ranks



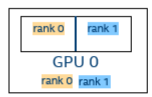
NUMA node 0 NUMA node 1

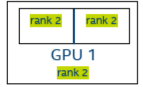
Debug output I_MPI_DEBUG=3:

```
[0] MPI startup(): ===== GPU pinning on host1 =====
[0] MPI startup(): Rank Pin tile
[0] MPI startup(): 0 {0}
```

```
[0] MPI startup(): 1 {1}
[0] MPI startup(): 2 {2}
[0] MPI startup(): 3 {3}
```

Figure 2. Three MPI Ranks





NUMA node 0

NUMA node 1

Debug output I MPI DEBUG=3:

I_MPI_OFFLOAD_TOPOLIB

Set the interface for GPU topology recognition.

Syntax

```
I MPI OFFLOAD TOPOLIB=<arg>
```

Arguments

<arg> String parameter.

level zero Use Level-Zero library for GPU topology recognition.

Description

Set this environment variable to define the interface for GPU topology recognition. Setting this variable enables the GPU Pinning feature.

I_MPI_OFFLOAD_LEVEL_ZERO_LIBRARY

Specify the name and full path to the Level-Zero library.

Syntax

I_MPI_OFFLOAD_LEVEL_ZERO_LIBRARY="<path>/<name>"

Arguments

<path> Full path to the Level-Zero library.

<name> Name of the Level-Zero library.

Description

Set this environment variable to specify the name and full path to Level-Zero library. Set this variable if Level-Zero is not located in the default path. Default value: libze_loader.so.

I_MPI_OFFLOAD_CELL

Set this variable to define the base unit: tile (subdevice) or device (GPU).

Syntax

I_MPI_OFFLOAD_CELL=<cell>

Arguments

<cell> Specify the base unit.

tile One tile (subdevice). Default value.

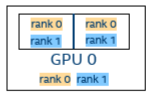
device Whole device (GPU) with all subdevices

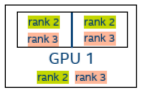
Description

Set this variable to define the base unit. This variable may affect other GPU pinning variables.

Example

Figure 3. Four MPI ranks, I_MPI_OFFLOAD_CELL=device





NUMA node 0

NUMA node 1

I_MPI_OFFLOAD_DOMAIN_SIZE

Control the number of base units per MPI rank.

Syntax

I MPI OFFLOAD DOMAIN SIZE=<value>

Arguments

<value> Integer number.

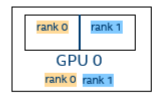
- -1 Auto. Default value. Each MPI rank may have a different domain size to use all available resources.
- > 0 Custom domain size.

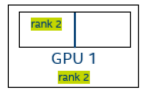
Description

Set this variable to define how many base units will be pinned to the MPI rank. I_MPI_OFFLOAD_CELL variable defines the base unit: tile or device.

Examples

Figure 4. Three MPI ranks, I_MPI_OFFLOAD_DOMAIN_SIZE=1





NUMA node 0

NUMA node 1

I_MPI_OFFLOAD_DEVICES

Define a list of available devices.

Syntax

I MPI OFFLOAD DEVICES=<devicelist>

Arguments

<devicelist> A comma-separated list of available devices.

all All devices are available. Default value.

<1> Device with logical number <1>.

<1>-<m> Range of devices with logical numbers from <1> to <m>.

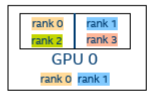
 $\langle k \rangle$, $\langle 1 \rangle$ - $\langle m \rangle$ Device $\langle k \rangle$ and devices from $\langle 1 \rangle$ to $\langle m \rangle$.

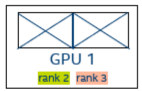
Description

Set this variable to define the available devices. This variable also gives you the ability to exclude devices.

Example

Figure 5. Four MPI ranks, I_MPI_OFFLOAD_DEVICES=0





NUMA node 0

NUMA node 1

I_MPI_OFFLOAD_DEVICE_LIST

Define a list of base units to pin for each MPI rank.

Syntax

```
I_MPI_OFFLOAD_DEVICE_LIST=<base_units_list>
```

Arguments

<base_units_list> A comma-separated list of base units. The process with the i-th rank is pinned to the ith base unit in the list.

<1> Base unit with logical number <1>.

<1>-<m> Range of base units with logical numbers from <1> to <m>.

<k>,<1>-<m> Base unit <k> and base units from <1> to <m>.

Description

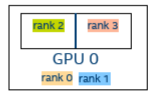
Set this variable to define the list of base units to pin for each MPI rank. The process with the i-th rank is pinned to the i-th base unit in the list.

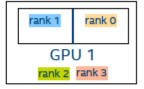
I MPI OFFLOAD CELL variable defines the base unit: tile or device.

I_MPI_OFFLOAD_DEVICE_LIST variable has less priority than the I_MPI_OFFLOAD_DOMAIN variable.

Example

Figure 6. Four MPI ranks, I_MPI_OFFLOAD_DEVICE_LIST=3,2,0,1





NUMA node 0

NUMA node 1

I_MPI_OFFLOAD_DOMAIN

Define domains through the comma separated list of hexadecimal numbers for each MPI rank.

Syntax

I MPI OFFLOAD DOMAIN=<masklist>

Arguments

<masklist> A comma-separated list of hexadecimal numbers.

[m1,...,mn] For <masklist>, each mi is a hexadecimal bit mask defining an individual domain.

The following rule is used: the i-th base unit is included into the domain if the corresponding bit in mi value is set to 1.

Description

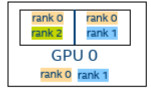
Set this variable to define the list of hexadecimal bit masks. For the i-th bit mask, if the j-th bit set to 1, then the j-th base unit will be pinned to the i-th MPI rank.

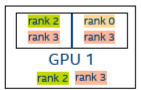
I_MPI_OFFLOAD_CELL variable defines the base unit: tile or device.

I_MPI_OFFLOAD_DOMAIN variable has higher priority than the I_MPI_OFFLOAD_DEVICE_LIST.

Example

Figure 7. Four MPI ranks, I_MPI_OFFLOAD_DOMAIN=[B,2,5,C]. Parsed bit masks: [1101,0100,1010,0011]





NUMA node 0

NUMA node 1

GPU Buffers Support

I MPI OFFLOAD

Syntax

I MPI OFFLOAD=<value>

Value	Integer Number
0	Disabled (default value)

Value	Integer Number
1	Auto. Intel MPI expects that libze_loader.so is already loaded and does not load it twice
2	Enabled. Intel MPI loads libze_loader.so

Set this variable to enable handling of device buffers in MPI functions such as MPI_Send, MPI_Recv, MPI_Bcast, MPI_Allreduce, and so on by using the Level Zero library specified in the I MPI OFFLOAD LEVEL ZERO LIBRARY variable.

In order to pass a pointer of an offloaded memory region to MPI, you may need to use specific compiler directives or get it from corresponding acceleration runtime API. For example, use_device_ptr and use_device_addr are useful keywords to obtain device pointers in the OpenMP environment, as shown in the following code sample:

```
/* Copy data from host to device */
#pragma omp target data map(to: rank, values[0:num_values]) use_device_ptr(values)
{
    /* Compute something on device */
    #pragma omp target parallel for is_device_ptr(values)
    for (unsigned i = 0; i < num_values; ++i) {
        values[i] *= (rank + 1);
    }
    /* Send device buffer to another rank */
    MPI_Send(values, num_values, MPI_INT, dest_rank, tag, MPI_COMM_WORLD);
}</pre>
```

Environment Variables for Fabrics Control

This section provides description of the general environment variables for controlling fabrics, as well as description of variables for controlling specific fabrics:

- Communication Fabrics Control
- Shared Memory Control
- OFI*-capable network fabrics

Communication Fabrics Control

I MPI FABRICS

Select the particular fabrics to be used.

Syntax

```
I MPI FABRICS=ofi | shm:ofi | shm
```

Arguments

<fabric></fabric>	Define a network fabric.
shm	Shared memory transport (used for intra-node communication only).
ofi	OpenFabrics Interfaces* (OFI)-capable network fabrics, such as Intel® True
	Scale Fabric, Intel® Omni-Path Architecture, InfiniBand*, and Ethernet
	(through OFI API).

Description

Set this environment variable to select a specific fabric combination.

The default values are <code>shm:ofi</code> for the regular mode and <code>ofi</code> for the multiple endpoints mode. In the multiple endpoints mode, the default value <code>ofi</code> cannot be changed.

NOTE

This option is not applicable to slurm and pdsh bootstrap servers.

NOTE

DAPL, TMI, and OFA fabrics are deprecated.

Shared Memory Control

I_MPI_SHM

Select a shared memory transport to be used.

Syntax

I_MPI_SHM=<transport>

<transport></transport>	Define a shared memory transport solution.
disable no off 0	Do not use shared memory transport.
auto	Select a shared memory transport solution automatically.
bdw_sse	The shared memory transport solution tuned for Intel $^{\odot}$ microarchitecture code name Broadwell. The SSE4.2. instruction set is used.
bdw_avx2	The shared memory transport solution tuned for Intel $^{\scriptsize \odot}$ microarchitecture code name Broadwell. The AVX2 instruction set is used.
skx_sse	The shared memory transport solution tuned for Intel® Xeon® processors based on Intel® microarchitecture code name Skylake. The CLFLUSHOPT and SSE4.2 instruction set is used.
skx_avx2	The shared memory transport solution tuned for Intel® Xeon® processors based on Intel® microarchitecture code name Skylake. The CLFLUSHOPT and AVX2 instruction set is used.
skx_avx512	The shared memory transport solution tuned for Intel® Xeon® processors based on Intel® microarchitecture code name Skylake. The CLFLUSHOPT and AVX512 instruction set is used.
knl_ddr	The shared memory transport solution tuned for Intel $\!\!^{\otimes}$ microarchitecture code name Knights Landing.
knl_mcdram	The shared memory transport solution tuned for Intel® microarchitecture code name Knights Landing. Shared memory buffers may be partially located in the Multi-Channel DRAM (MCDRAM).
clx_sse	The shared memory transport solution tuned for Intel® Xeon® processors based on Intel® microarchitecture code name Cascade Lake. The CLFLUSHOPT and SSE4.2 instruction set is used.
clx_avx2	The shared memory transport solution tuned for Intel® Xeon® processors based on Intel® microarchitecture code name Cascade Lake. The CLFLUSHOPT and AVX2 instruction set is used.

The shared memory transport solution tuned for Intel® Xeon® processors based
on Intel [®] microarchitecture code name Cascade Lake. The CLFLUSHOPT and
AVX512 instruction set is used.
The shared memory transport solution tuned for Intel® Xeon® processors based
on Intel [®] microarchitecture code name Cascade Lake Advanced Performance.
The shared memory transport solution tuned for Intel® Xeon® processors based on Intel® microarchitecture code name Ice Lake.

Set this environment variable to select a specific shared memory transport solution.

Automatically selected transports:

- icx for Intel® Xeon® processors based on Intel® microarchitecture code name Ice Lake
- clx-ap for Intel® Xeon® processors based on Intel® microarchitecture code name Cascade Lake Advanced Performance
- bdw avx2 for Intel® microarchitecture code name Haswell, Broadwell and Skylake
- skx avx2 for Intel® Xeon® processors based on Intel® microarchitecture code name Skylake
- ckx avx2 for Intel® Xeon® processors based on Intel® microarchitecture code name Cascade Lake
- knl mcdram for Intel® microarchitecture code name Knights Landing and Knights Mill
- bdw sse for all other platforms

The value of <code>I_MPI_SHM</code> depends on the value of <code>I_MPI_FABRICS</code> as follows: if <code>I_MPI_FABRICS</code> is <code>ofi</code>, <code>I_MPI_SHM</code> is disabled. If <code>I_MPI_FABRICS</code> is <code>shm:ofi</code>, <code>I_MPI_SHM</code> defaults to auto or takes the specified value.

I_MPI_SHM_CELL_FWD_SIZE

Change the size of a shared memory forward cell.

Syntax

I_MPI_SHM_CELL_FWD_SIZE=<nbytes>

Arguments

<nbytes></nbytes>	The size of a shared memory forward cell in bytes
> 0	The default <nbytes> value depends on the transport used and should</nbytes>
	normally range from 64K to 1024K.

Description

Forward cells are in-cache message buffer cells used for sending small amounts of data. Lower values are recommended. Set this environment variable to define the size of a forward cell in the shared memory transport.

I_MPI_SHM_CELL_BWD_SIZE

Change the size of a shared memory backward cell.

Syntax

I MPI SHM CELL BWD SIZE=<nbytes>

Arguments

<nbytes></nbytes>	The size of a shared memory backward cell in bytes
> 0	The default <nbytes> value depends on the transport used and should</nbytes>
	normally range from 64K to 1024K.

Description

Backward cells are out-of-cache message buffer cells used for sending large amounts of data. Higher values are recommended. Set this environment variable to define the size of a backwrad cell in the shared memory transport.

I_MPI_SHM_CELL_EXT_SIZE

Change the size of a shared memory extended cell.

Syntax

I MPI SHM CELL EXT SIZE=<nbytes>

Arguments

<nbytes></nbytes>	The size of a shared memory extended cell in bytes
> 0	The default <nbytes> value depends on the transport used and should</nbytes>
	normally range from 64K to 1024K.

Description

Extended cells are used in the imbalanced applications when forward and backward cells are run out. An extended cell does not have a specific owner - it is shared between all ranks on the computing node. Set this environment variable to define the size of an extended cell in the shared memory transport.

I_MPI_SHM_CELL_FWD_NUM

Change the number of forward cells in the shared memory transport (per rank).

Syntax

I_MPI_SHM_CELL_FWD_NUM=<num>

Arguments

<num></num>	The number of shared memory forward cells
> 0	The default value depends on the transport used and should normally range
	from 4 to 16.

Description

Set this environment variable to define the number of forward cells in the shared memory transport.

I_MPI_SHM_CELL_BWD_NUM

Change the number of backward cells in the shared memory transport (per rank).

Syntax

I MPI SHM CELL BWD NUM=<num>

Arguments

<num></num>	The number of shared memory backward cells
> 0	The default value depends on the transport used and should normally range
	from 4 to 64.

Description

Set this environment variable to define the number of backward cells in the shared memory transport.

I MPI SHM CELL EXT NUM TOTAL

Change the total number of extended cells in the shared memory transport.

Syntax

```
I_MPI_SHM_CELL_EXT_NUM_TOTAL=<num>
```

Arguments

<num></num>	The number of shared memory backward cells
> 0	The default value depends on the transport used and should normally range
	from 2K to 8K.

Description

Set this environment variable to define the number of extended cells in the shared memory transport.

NOTE

This is not "per rank" number, it is total number of extended cells on the computing node.

I_MPI_SHM_CELL_FWD_HOLD_NUM

Change the number of hold forward cells in the shared memory transport (per rank).

Syntax

I_MPI_SHM_CELL_FWD_HOLD_NUM=<num>

Arguments

<num></num>	The number of shared memory hold forward cells
> 0	The default value depends on the transport used and must be less than
	I_MPI_SHM_CELL_FWD_NUM.

Description

Set this environment variable to define the number of forward cells in the shared memory transport a rank can hold at the same time. Recommended values are powers of two in the range between 1 and 8.

I_MPI_SHM_MCDRAM_LIMIT

Change the size of the shared memory bound to the multi-channel DRAM (MCDRAM) (size per rank).

Syntax

I_MPI_SHM_MCDRAM_LIMIT=<nbytes>

Arguments

<nbytes></nbytes>	The size of the shared memory bound to MCDRAM per rank
1048576	This is the default value.

Description

Set this environment variable to define how much MCDRAM memory per rank is allowed for the shared memory transport. This variable takes effect with $I_MPI_SHM=knl_mcdram$ only.

I_MPI_SHM_SEND_SPIN_COUNT

Control the spin count value for the shared memory transport for sending messages.

Syntax

I_MPI_SHM_SEND_SPIN_COUNT=<count>

Arguments

<pre><count></count></pre> Define the spin count value. A typical value range is between 1 and 1000.
--

Description

If the recipient ingress buffer is full, the sender may be blocked until this spin count value is reached. It has no effect when sending small messages.

I_MPI_SHM_RECV_SPIN_COUNT

Control the spin count value for the shared memory transport for receiving messages.

Syntax

I MPI SHM RECV SPIN COUNT=<count>

Arguments

<pre><count></count></pre> Define the spin count value. A typical value range is between 1 and 1000)000.
---	-------

Description

If the receive is non-blocking, this spin count is used only for safe reorder of expected and unexpected messages. It has no effect on receiving small messages.

I_MPI_SHM_FILE_PREFIX_4K

Change the mount point of the 4 KB pages size file system (tmpfs) where the shared memory files are created.

Syntax

I MPI SHM FILE PREFIX 4K=<path>

Arguments

<path></path>	Define the path to the existed mount point of the 4 KB pages size file system
	(tmpfs). By default, the path is not set.

Description

Set this environment variable to define a new path to the shared memory files. By default, the shared memory files are created at /dev/shm/.

This variable affects shared memory transport buffers and RMA windows.

Example

```
I_MPI_SHM_FILE_PREFIX_4K=/dev/shm/intel/
```

I_MPI_SHM_FILE_PREFIX_2M

Change the mount point of the 2 MB pages size file system (hugetlbfs) where the shared memory files are created.

Syntax

I MPI SHM FILE PREFIX 2M=<path>

Arguments

<path></path>	Define the path to the existed mount point of the 2 MB pages size file system
	(hugetlbfs). By default, the path is not set.

Description

Set this environment variable to enable 2 MB huge pages on the Intel MPI Library.

The variable affects shared memory transport buffers. It may affect RMA windows as well if the windows size is greater than or equal to 2 MB.

Example

I MPI SHM FILE PREFIX 2M=/dev/hugepages

NOTE

The root privileges are required to configure the huge pages subsystem. Contact your system administrator to obtain permission.

I_MPI_SHM_FILE_PREFIX_1G

Change the mount point of the 1 GB pages size file system (hugetlbfs) where the shared memory files are created.

Syntax

I_MPI_SHM_FILE_PREFIX_1G=<path>

Arguments

<path></path>	Define the path to the existed mount point of the 1 GB pages size file system
	(hugetlbfs). By default, the path is not set.

Description

Set this environment variable to enable 1 GB huge pages on the Intel MPI Library.

The variable affects shared memory transport buffers. It may affect RMA windows as well if the windows size is greater than or equal to 1 GB.

Example

I MPI SHM FILE PREFIX 1G=/dev/hugepages1G

NOTE

The root privileges are required to configure the huge pages subsystem. Contact your system administrator to obtain permission.

OFI*-capable Network Fabrics Control

I_MPI_OFI_PROVIDER

Define the name of the OFI provider to load.

Syntax

I_MPI_OFI_PROVIDER=<name>

Arguments

<name> The name of the OFI provider to load

Description

Set this environment variable to define the name of the OFI provider to load. If you do not specify this variable, the OFI library chooses the provider automatically. You can check all available providers by using the <code>I_MPI_OFI_PROVIDER_DUMP</code> environment variable. If you set the wrong name for an available provider, use <code>FI_LOG_LEVEL=debug</code> to get a hint to set the name correctly.

I MPI OFI PROVIDER DUMP

Control the capability of printing information about all OFI providers and their attributes from an OFI library.

Syntax

```
I_MPI_OFI_PROVIDER_DUMP=<arg>
```

Arguments

<arg></arg>	Binary indicator
enable yes on 1	Print the list of all OFI providers and their attributes from an OFI library
disable no off 0	No action. This is the default value

Description

Set this environment variable to control the capability of printing information about all OFI providers and their attributes from an OFI library.

I_MPI_OFI_DRECV

Control the capability of the direct receive in the OFI fabric.

Syntax

```
I MPI OFI DRECV=<arg>
```

Arguments

<arg></arg>	Binary indicator
enable yes on 1	Enable direct receive. This is the default value
disable no off 0	Disable direct receive

Description

Use the direct receive capability to block MPI_Recv calls only. Before using the direct receive capability, ensure that you use it for single-threaded MPI applications and check if you have selected OFI as the network fabric by setting $I_MPI_FABRICS=ofi$.

I_MPI_OFI_LIBRARY_INTERNAL

Control the usage of libfabric* shipped with the Intel® MPI Library.

Syntax

```
I MPI OFI LIBRARY INTERNAL=<arg>
```

Arguments

<arg></arg>	Binary indicator
enable yes on 1	Use libfabric from the Intel MPI Library
disable no off 0	Do not use libfabric from the Intel MPI Library

Description

Set this environment variable to disable or enable usage of libfabric from the Intel MPI Library. The variable must be set before sourcing the vars.[c]sh script.

Example

```
$ export I_MPI_OFI_LIBRARY_INTERNAL=1
$ source <installdir> /bin/vars.sh
```

Setting this variable is equivalent to passing the -ofi internal option to the vars. [c] sh script.

For more information, refer to the Intel® MPI Library Developer Guide, section Libfabric* Support.

Environment Variables for Memory Policy Control

Intel® MPI Library supports non-uniform memory access (NUMA) nodes with high-bandwidth (HBW) memory (MCDRAM) on Intel® Xeon Phi™ processors (codenamed Knights Landing). Intel® MPI Library can attach memory of MPI processes to the memory of specific NUMA nodes. This section describes the environment variables for such memory placement control.

I_MPI_HBW_POLICY

Set the policy for MPI process memory placement for using HBW memory.

Syntax

I_MPI_HBW_POLICY=<user memory policy>[,<mpi memory policy>][,<win_allocate policy>]

In the syntax:

- <user memory policy> memory policy used to allocate the memory for user applications (required)
- <mpi memory policy> memory policy used to allocate the internal MPI memory (optional)
- <win_allocate policy> memory policy used to allocate memory for window segments for RMA operations (optional)

Each of the listed policies may have the values below:

Arguments

<value></value>	The memory allocation policy used.
hbw_preferred	Allocate the local HBW memory for each process. If the HBW memory is not available, allocate the local dynamic random access memory.
hbw_bind	Allocate only the local HBW memory for each process.
hbw_interleave	Allocate the HBW memory and dynamic random access memory on the local node in the round-robin manner.

Description

Use this environment variable to specify the policy for MPI process memory placement on a machine with HBW memory.

By default, Intel MPI Library allocates memory for a process in local DDR. The use of HBW memory becomes available only when you specify the I MPI HBW POLICY variable.

Examples

The following examples demonstrate different configurations of memory placement:

• I MPI HBW POLICY=hbw bind, hbw preferred, hbw bind

Only use the local HBW memory allocated in user applications and window segments for RMA operations. Use the local HBW memory internally allocated in Intel® MPI Library first. If the HBW memory is not available, use the local DDR internally allocated in Intel MPI Library.

• I MPI HBW POLICY=hbw bind, , hbw bind

Only use the local HBW memory allocated in user applications and window segments for RMA operations. Use the local DDR internally allocated in Intel MPI Library.

• I_MPI_HBW_POLICY=hbw_bind,hbw_preferred

Only use the local HBW memory allocated in user applications. Use the local HBW memory internally allocated in Intel MPI Library first. If the HBW memory is not available, use the local DDR internally allocated in Intel MPI Library. Use the local DDR allocated in window segments for RMA operations.

I_MPI_BIND_NUMA

Set the NUMA nodes for memory allocation.

Syntax

 ${\tt I_MPI_BIND_NUMA} = <\! value >$

Arguments

<value></value>	Specify the NUMA nodes for memory allocation.
localalloc	Allocate memory on the local node. This is the default value.
Node_1,,Node_k	Allocate memory according to <code>I_MPI_BIND_ORDER</code> on the specified NUMA
	nodes.

Description

Set this environment variable to specify the NUMA node set that is involved in the memory allocation procedure.

I_MPI_BIND_ORDER

Set this environment variable to define the memory allocation manner.

Syntax

I_MPI_BIND_ORDER=<value>

Arguments

<value></value>	Specify the allocation manner.
compact	Allocate memory for processes as close as possible (in terms of NUMA nodes), among the NUMA nodes specified in <code>I_MPI_BIND_NUMA</code> . This is the default value.
scatter	Allocate memory among the NUMA nodes specified in <code>I_MPI_BIND_NUMA</code> using the round-robin manner.

Description

Set this environment variable to define the memory allocation manner among the NUMA nodes specified in I MPI BIND NUMA. The variable has no effect without I MPI BIND NUMA set.

I_MPI_BIND_WIN_ALLOCATE

Set this environment variable to control memory allocation for window segments.

Syntax

I MPI BIND WIN ALLOCATE=<value>

Arguments

<value></value>	Specify the memory allocation behavior for window segments.
localalloc	Allocate memory on the local node. This is the default value.
hbw_preferred	Allocate the local HBW memory for each process. If the HBW memory is not available, allocate the local dynamic random access memory.
hbw_bind	Allocate only the local HBW memory for each process.
hbw_interleave	Allocate the HBW memory and dynamic random access memory on a local node in the round-robin manner.
<numa id="" node=""></numa>	Allocate memory on the given NUMA node.

Description

Set this environment variable to create window segments allocated in HBW memory with the help of the MPI Win allocate shared or MPI Win allocate functions.

MPI_Info

You can control memory allocation for window segments with the help of an MPI_Info object, which is passed as a parameter to the MPI_Win_allocate or MPI_Win_allocate_shared function. In an application, if you specify such an object with the numa_bind_policy key, window segments are allocated in accordance with the value for numa_bind_policy. Possible values are the same as for I MPI BIND WIN ALLOCATE.

A code fragment demonstrating the use of MPI Info:

```
MPI_Info info;
...
MPI_Info_create( &info );
MPI_Info_set( info, "numa_bind_policy", "hbw_preferred" );
...
MPI Win allocate shared( size, disp unit, info, comm, &baseptr, &win );
```

NOTE

When you specify the memory placement policy for window segments, Intel MPI Library recognizes the configurations according to the following priority:

- **1.** Setting of MPI Info.
- 2. Setting of I_MPI_HBW_POLICY, if you specified <win allocate policy>.
- **3.** Setting of <code>I_MPI_BIND_WIN_ALLOCATE</code>.

Environment Variables for Asynchronous Progress Control

NOTE

This feature is supported for the release_mt and debug_mt library configurations only. To specify the configuration, run the following command:

```
$ source <installdir>/bin/vars.sh release mt
```

I MPI ASYNC PROGRESS

Control the usage of progress threads.

Syntax

```
I MPI ASYNC PROGRESS=<arg>
```

Arguments

<arg></arg>	Binary indicator
disable no off 0	Disable asynchronous progress threads for each rank. This is the default value.
enable yes on 1	Enable asynchronous progress threads.

Description

Set this environment variable to enable asynchronous progress. If disabled, the <code>I_MPI_ASYNC_PROGRESS_*</code> knobs are ignored.

I MPI ASYNC PROGRESS THREADS

Control the number of asynchronous progress threads.

Syntax

I_MPI_ASYNC_PROGRESS_THREADS=<arg>

Arguments

<nthreads></nthreads>	Define the number of progress threads. The default value
	is 1.

Description

Set this environment variable to control the number of asynchronous progress threads for each rank.

I MPI ASYNC PROGRESS PIN

Control the asynchronous progress threads pinning.

Syntax

I MPI ASYNC PROGRESS PIN=<arg>

Arguments

<arg></arg>	Comma-separated list of logical processors
<cpu list=""></cpu>	Pin all progress threads of local processes to the listed
	CPUs. By default, N progress threads are pinned to the
	last N logical processors.

Description

Set this environment variable to control pinning for all progress threads of local processes.

Example

```
I_MPI_ASYNC_PROGRESS_THREADS=3
I_MPI_ASYNC_PROGRESS_PIN="0,1,2,3,4,5"
```

In case of three MPI processes per node, progress threads of the first process are pinned to 0, 1, second are pinned to 2, 3, and third are pinned to 4, 5.

I_MPI_ASYNC_PROGRESS_ID_KEY

Set the MPI info object key that is used to explicitly define the progress thread id for a communicator.

Syntax

```
I MPI ASYNC PROGRESS ID KEY=<arg>
```

Arguments

<key></key>	MPI info object key. The default value is thread_id.
-------------	--

Description

Set this environment variable to control the MPI info object key that is used to define the progress thread id for a communicator. The progress thread id is used for work distribution between progress threads. By default, communication goes over the first progress thread.

NOTE

Exclude selected processors for progress threads from pinning of computational threads to avoid oversubscription of cores.

For more information and examples, refer to the Intel® MPI Library Developer Guide, section Asynchronous Progress Control.

Environment Variables for Multi-EP

NOTE This feature is supported for the release_mt and debug_mt library configurations only. To specify the configuration, run the following command: $\$ source <install-dir>/bin/vars.sh release_mt

I_MPI_THREAD_SPLIT

Syntax

I MPI THREAD SPLIT=<value>

Arguments

Value	Binary Indicator
0 no off disable	Disable the MPI_THREAD_SPLIT model support. This is the default value.
1 yes on enable	Enable the MPI_THREAD_SPLIT model support.

Description

Use this environment variable to control the I MPI THREAD SPLIT programming model.

I_MPI_THREAD_RUNTIME

Syntax

I MPI THREAD RUNTIME=<value>

Arguments

Value	Thread Runtime
generic	Enable runtime support (for example, pthreads, TBB). This is the default value if OpenMP* cannot be detected at runtime.
openmp	Enable OpenMP runtime support. This is the default value if OpenMP is detected at runtime.

Description

Use this environment variable to control threading runtime support.

NOTE: MPI THREAD SPLIT model support is enabled.

I_MPI_THREAD_MAX

Syntax

I MPI THREAD MAX=<int>

Arguments

<int></int>	The maximum number of threads per rank. The default value is
	<pre>omp_get_max_threads() if I_MPI_THREAD_RUNTIME is set to openmp.</pre>
	The value is 1 otherwise

Description

Use this environment variable to set the maximum number of threads to be used in each process concurrently.

I_MPI_THREAD_ID_KEY

Syntax

I MPI THREAD ID KEY=<string>

Arguments

<string></string>	Define the MPI info object key. The default value is <pre>thread_id</pre>
-------------------	---

Description

Use this environment variable to set the MPI info object key that is used to explicitly define the logical thread $number\ thread\ id.$

Other Environment Variables

I_MPI_DEBUG

Print out debugging information when an MPI program starts running.

Syntax

I MPI DEBUG=<level>[,<flags>]

<level></level>	Indicate the level of debug information provided.
0	Output no debugging information. This is the default value.
1,2	Output libfabric* version and provider.
3	Output effective MPI rank, pid and node mapping table.
4	Output process pinning information.
5	Output environment variables specific to the Intel $^{\mbox{\scriptsize 0}}$ MPI Library.
> 5	Add extra levels of debug information.

<flag< th=""><th>Comma-separated list of debug flags</th></flag<>	Comma-separated list of debug flags
s>	
pid	Show process id for each debug message.
tid	Show thread id for each debug message for multithreaded library.
time	Show time for each debug message.
datet	Show time and date for each debug message.
ime	
host	Show host name for each debug message.
level	Show level for each debug message.
scope	Show scope for each debug message.
line	Show source line number for each debug message.
file	Show source file name for each debug message.
nofun	Do not show routine name.
С	

noran	Do not show rank.	ĺ
k		ĺ
nousr	Suppress warnings for improper use case (for example, incompatible combination of controls).	ĺ
warn		ĺ
flock	Synchronize debug output from different process or threads.	ĺ
nobuf	Do not use buffered I/O for debug output.	

Set this environment variable to print debugging information about the application.

```
NOTE Set the same < level> value for all ranks.
```

You can specify the output file name for debug information by setting the $I_MPI_DEBUG_OUTPUT$ environment variable.

Each printed line has the following format:

```
[<identifier>] <message>
```

where:

- <identifier> is the MPI process rank, by default. If you add the '+' sign in front of the <level> number, the <identifier> assumes the following format: rank#pid@hostname. Here, rank is the MPI process rank, pid is the UNIX* process ID, and hostname is the host name. If you add the '-' sign, <identifier> is not printed at all.
- <message> contains the debugging output.

The following examples demonstrate possible command lines with the corresponding output:

```
$ mpirun -n 1 -env I_MPI_DEBUG=2 ./a.out
...
[0] MPI startup(): shared memory data transfer mode
```

The following commands are equal and produce the same output:

```
$ mpirun -n 1 -env I_MPI_DEBUG=2,pid,host ./a.out
...
[0#1986@mpicluster001] MPI startup(): shared memory data transfer mode
```

NOTE Compiling with the $\neg g$ option adds a considerable amount of printed debug information.

I_MPI_DEBUG_OUTPUT

Set output file name for debug information.

Syntax

```
I_MPI_DEBUG_OUTPUT=<arg>
```

Argument	String Value
stdout	Output to stdout. This is the default value.
stderr	Output to stderr.
<file_name></file_name>	Specify the output file name for debug information (the maximum file name length is 256 symbols).

Set this environment variable if you want to split output of debug information from the output produced by an application. If you use format like \$r, \$p or \$h, rank, process ID or host name is added to the file name accordingly.

I_MPI_DEBUG_COREDUMP

Controls core dump files generation in case of failure during MPI application execution.

Syntax

I MPI DEBUG COREDUMP=<arg>

Arguments

Argument	Binary Indicator
enable yes on 1	Enable coredump files generation.
disable no off 0	Do not generate coredump files. Default value.

Description

Set this environment variable to enable coredump files dumping in case of termination caused by segmentation fault. Available for both release and debug builds.

I_MPI_STATS

Collect MPI statistics from your application using Application Performance Snapshot.

Syntax

I MPI STATS=<level>

Arguments

<level></level>	Indicate the level of statistics collected
1,2,3,4,5	Specify the level to indicate amount of MPI statistics to be collected by Application Performance Snapshot (APS).
	The full description of levels is available in the official APS documentation.

Description

Set this variable to collect MPI-related statistics from your MPI application using Application Performance Snapshot. The variable creates a new folder aps_result_<date>-<time> containing statistics data. To analyze the collected data, use the aps utility. For example:

```
$ export I_MPI_STATS=5
$ mpirun -n 2 ./myApp
$ aps-report aps_result_20171231_235959
```

I MPI STARTUP MODE

Select a mode for the Intel® MPI Library process startup algorithm.

Syntax

I MPI STARTUP MODE=<arg>

Argument	String Value	
	<u> </u>	

pmi_shm	Use shared memory to reduce the number of PMI calls. This mode is enabled by default.
pmi_shm_netmod	Use the netmod infrastructure for address exchange logic in addition to PMI and shared memory.

The pmi_shm and pmi_shm_netmod modes reduce the application startup time. The efficiency of the modes is more clearly observed with the higher -ppn value, while there is no improvement at all with -ppn 1.

I_MPI_PMI_LIBRARY

Specify the name to third party implementation of the PMI library.

Syntax

I MPI PMI LIBRARY=<name>

Arguments

<name></name>	Full name of the third party PMI library
---------------	--

Description

Set I_MPI_PMI_LIBRARY to specify the name of third party PMI library. When you set this environment variable, provide full name of the library with full path to it.

Currently supported PMI versions: PMI1, PMI2

I_MPI_PMI_VALUE_LENGTH_MAX

Control the length of the value buffer in PMI on the client side.

Syntax

I MPI PMI VALUE LENGTH MAX=<length>

Arguments

<length></length>	Define the value of the buffer length in bytes.
<n> > 0</n>	The default value is -1, which means do not override the value received from
	<pre>the PMI_KVS_Get_value_length_max() function.</pre>

Description

Set this environment variable to control the length of the value buffer in PMI on the client side. The length of the buffer will be the lesser of I MPI PMI VALUE LENGTH MAX and PMI KVS Get value length max().

I_MPI_OUTPUT_CHUNK_SIZE

Set the size of the stdout/stderr output buffer.

Syntax

I MPI OUTPUT CHUNK SIZE=<size>

Arguments

<size></size>	Define output chunk size in kilobytes
<n>> 0</n>	The default chunk size value is 1 KB

Description

Set this environment variable to increase the size of the buffer used to intercept the standard output and standard error streams from the processes. If the $\langle size \rangle$ value is not greater than zero, the environment variable setting is ignored and a warning message is displayed.

Use this setting for applications that create a significant amount of output from different processes. With the -ordered-output option of mpiexec.hydra, this setting helps to prevent the output from garbling.

NOTE Set the <code>I_MPI_OUTPUT_CHUNK_SIZE</code> environment variable in the shell environment before executing the <code>mpiexec.hydra/mpirun</code> command. Do not use the <code>-genv</code> or <code>-env</code> options for setting the <code><size></code> value. Those options are used only for passing environment variables to the MPI process environment.

I_MPI_REMOVED_VAR_WARNING

Print out a warning if a removed environment variable is set.

Syntax

I MPI REMOVED VAR WARNING=<arg>

Arguments

Argument	Binary Indicator
enable yes on 1	Print out the warning. This is the default value
disable no off 0	Do not print the warning

Description

Use this environment variable to print out a warning if a removed environment variable is set. Warnings are printed regardless of whether I MPI DEBUG is set.

I MPI VAR CHECK SPELLING

Print out a warning if an unknown environment variable is set.

Syntax

I_MPI_VAR_CHECK_SPELLING=<arg>

Arguments

Argument	Binary Indicator
enable yes on 1	Print out the warning. This is the default value
disable no off 0	Do not print the warning

Description

Use this environment variable to print out a warning if an unsupported environment variable is set. Warnings are printed in case of removed or misprinted environment variables.

I_MPI_LIBRARY_KIND

Specify the Intel® MPI Library configuration.

Syntax

I MPI LIBRARY KIND=<value>

Value	Binary Indicator
release	Multi-threaded optimized library (with the global lock). This is the default value
debug	Multi-threaded debug library (with the global lock)
release_mt	Multi-threaded optimized library (with per-object lock for the thread-split model)
debug_mt	Multi-threaded debug library (with per-object lock for the thread-split model)

Use this variable to set an argument for the vars.[c]shscript. This script establishes the Intel® MPI Library environment and enables you to specify the appropriate library configuration. To ensure that the desired configuration is set, check the LD LIBRARY PATH variable.

Example

```
$ export I MPI LIBRARY KIND=debug
```

Setting this variable is equivalent to passing an argument directly to the vars. [c] sh script:

Example

\$. <installdir>/bin/vars.sh release

I_MPI_PLATFORM

Select the intended optimization platform.

Syntax

I MPI PLATFORM=<platform>

Arguments

<pre><platform></platform></pre>	Intended optimization platform (string value)	
auto	Use only with heterogeneous runs to determine the appropriate platform across all nodes. May slow down MPI initialization time due to collective operation across all nodes.	
ivb	Optimize for the Intel® Xeon® Processors E3, E5, and E7 V2 series and other Intel® Architecture processors formerly code named Ivy Bridge.	
hsw	Optimize for the Intel Xeon Processors E3, E5, and E7 V3 series and other Intel® Architecture processors formerly code named Haswell.	
bdw	Optimize for the Intel Xeon Processors E3, E5, and E7 V4 series and other Intel Architecture processors formerly code named Broadwell.	
knl	Optimize for the Intel [®] Xeon Phi [™] processor and coprocessor formerly code named Knights Landing.	
skx	Optimize for the Intel Xeon Processors E3 V5 and Intel Xeon Scalable Family series, and other Intel Architecture processors formerly code named Skylake.	
clx	Optimize for the 2nd Generation Intel Xeon Scalable Processors, and other Intel® Architecture processors formerly code named Cascade Lake.	
clx-ap	Optimize for the 2nd Generation Intel Xeon Scalable Processors, and other Intel Architecture processors formerly code named Cascade Lake AP Note: The explicit $clx-ap$ setting is ignored if the actual platform is not Intel.	

Description

Set this environment variable to use the predefined platform settings. The default value is a local platform for each node.

The variable is available for both Intel and non-Intel microprocessors, but it may utilize additional optimizations for Intel microprocessors than it utilizes for non-Intel microprocessors.

NOTE The values auto[:min], auto:max, and auto:most may increase the MPI job startup time.

I_MPI_MALLOC

Control the Intel® MPI Library custom allocator of private memory.

Syntax

I MPI MALLOC=<arg>

Argument

Argument	Binary Indicator
Enable the Intel MPI Library custom allocator of private r	
	Use the Intel MPI custom allocator of private memory for MPI_Alloc_mem/MPI_Free_mem.
0	Disable the Intel MPI Library custom allocator of private memory.
	Use the system-provided memory allocator for MPI_Alloc_mem/MPI_Free_mem.

Description

Use this environment variable to enable or disable the Intel MPI Library custom allocator of private memory for MPI Alloc mem/MPI Free mem.

By default, <code>I_MPI_MALLOC</code> is enabled for release and debug Intel MPI Library configurations and disabled for release mt sand debug mt configurations.

NOTE If the platform is not supported by the Intel MPI Library custom allocator of private memory, a system-provided memory allocator is used and the I MPI MALLOC variable is ignored.

I_MPI_SHM_HEAP

Control the Intel® MPI Library custom allocator of shared memory.

Syntax

I MPI SHM HEAP=<arg>

Argument

Argument	Binary Indicator	
1	Use the Intel MPI custom allocator of shared memory for MPI_Alloc_mem/	
	MPI_Free_mem.	
0	Do not use the Intel MPI custom allocator of shared memory for	
	MPI_Alloc_mem/MPI_Free_mem.	

Description

Use this environment variable to enable or disable the Intel MPI Library custom allocator of shared memory for MPI Alloc mem/MPI Free mem.

By default, $I_MPI_SHM_HEAP$ is disabled. If enabled, it can improve performance of the shared memory transport because in that case it is possible to make only one memory copy operation instead of two copy-in/copy-out memory copy operations. If both $I_MPI_SHM_HEAP$ and I_MPI_MALLOC are enabled, the shared memory allocator is used first. The private memory allocator is used only when required volume of shared memory is not available.

Details

By default, the shared memory segment is allocated on tmpfs file system on the /dev/shm/ mount point. Starting from Linux kernel 4.7, it is possible to enable transparent huge pages on the shared memory. If Intel MPI Library shared memory heap is used, it is recommended to enable transparent huge pages on your system. To enable transparent huge pages on /dev/shm, please contact your system administrator or execute the following command:

```
sudo mount -o remount, huge=advise /dev/shm
```

In order to use another tmpfs mount point instead of /dev/shm/, use I_MPI_SHM_FILE_PREFIX_4K, I MPI SH M FILE PREFIX 2M, and I MPI SHM FILE PREFIX 1G.

NOTE If your application does not use MPI_Alloc_mem/MPI_Free_mem directly, you can override standard malloc/calloc/realloc/free procedures by preloading the libmpi_shm_heap_proxy.so library:

```
export LD_PRELOAD=$I_MPI_ROOT/lib/libmpi_shm_heap_proxy.so
export I_MPI_SHM_HEAP=1
```

In this case, the malloc/calloc/realloc is a proxy for MPI_Alloc_mem and free is a proxy for MPI Free mem.

NOTE

If the platform is not supported by the Intel MPI Library custom allocator of shared memory, the I MPI SHM HEAP variable is ignored.

I MPI SHM HEAP VSIZE

Change the size (per rank) of virtual shared memory available for the Intel MPI Library custom allocator of shared memory.

Syntax

I MPI SHM HEAP VSIZE=<size>

Argument

<size></size>	The size (per rank) of shared memory used in shared memory heap (in megabytes).
>0	If shared memory heap is enabled for MPI_Alloc_mem/MPI_Free_mem, the default value is 4096.

Description

Intel MPI Library custom allocator of shared memory works with fixed size virtual shared memory. The shared memory segment is allocated on MPI Init and cannot be enlarged later.

The I MPI SHM HEAP VSIZE=0 completely disables the Intel MPI Library shared memory allocator.

I MPI SHM HEAP CSIZE

Change the size (per rank) of shared memory cached in the Intel MPI Library custom allocator of shared memory.

Syntax

I MPI SHM HEAP CSIZE=<size>

Argument

<size></size>	The size (per rank) of shared memory used in Intel MPI Library shared memory allocator (in megabytes).
>0	It depends on the available shared memory size and number of ranks. Normally, the size is less than 256.

Description

Small values of I_MPI_SHM_HEAP_CSIZE may reduce overall shared memory consumption. Larger values of this variable may speed up MPI Alloc mem/MPI Free mem.

I_MPI_SHM_HEAP_OPT

Change the optimization mode of Intel MPI Library custom allocator of shared memory.

Syntax

I MPI SHM HEAP OPT=<mode>

Argument

Mode	Optimization Mode	
rank	In this mode, each rank has its own dedicated amount of shared memory. This	
	is the default value when <code>I_MPI_SHM_HEAP=1</code>	
numa	In this mode, all ranks from NUMA-node use the same amount of shared	
	memory.	

Description

It is recommended to use <code>I_MPI_SHM_HEAP_OPT=rank</code> when each rank uses the same amount of memory, and <code>I MPI SHM HEAP OPT=numa</code> when ranks use significantly different amounts of memory.

Usually, the I_MPI_SHM_HEAP_OPT=rank works faster than I_MPI_SHM_HEAP_OPT=numa but the numa optimization mode may consume smaller volume of shared memory.

I_MPI_WAIT_MODE

Control the Intel® MPI Library optimization for oversubscription mode.

Syntax

I_MPI_WAIT_MODE=<arg>

Arguments

Argument	Binary Indicator
0	Optimize MPI application to work in the normal mode (1 rank on 1 CPU). This is the default value if the number of processes on a computation node is less than or equal to the number of CPUs on the node.
1	Optimize MPI application to work in the oversubscription mode (multiple ranks on 1 CPU). This is the default value if the number of processes on a computation node is greater than the number of CPUs on the node.

Description

It is recommended to use this variable in the oversubscription mode.

I_MPI_THREAD_YIELD

Control the Intel® MPI Library thread yield customization during MPI busy wait time.

Syntax

I_MPI_THREAD_YIELD=<arg>

Arguments

Argument	Binary indicator
0	Do nothing for thread yield during the busy wait (spin wait). This is the default value when $I_MPI_WAIT_MODE=0$
1	Do the pause processor instruction for <code>I_MPI_PAUSE_COUNT</code> during the busy wait.
2	Do the <pre>shied_yield()</pre> system call for thread yield during the busy wait.
	This is the default value when <pre>I_MPI_WAIT_MODE=1</pre>
3	Do the usleep() system call for I_MPI_THREAD_SLEEP number of microseconds for thread yield during the busy wait.

Description

I_MPI_THREAD_YIELD=0 or I_MPI_THREAD_YIELD=1 in the normal mode and I_MPI_THREAD_YIELD=2 or I MPI THREAD YIELD=3 in the oversubscription mode.

I_MPI_PAUSE_COUNT

Control the Intel® MPI Library pause count for the thread yield customization during MPI busy wait time.

Syntax

I MPI PAUSE COUNT=<arg>

Argument

Argument	Description Pause count for thread yield customization during MPI busy wait time.	
>=0		
	The default value is 0. Normally, the value is less than 100.	

Description

This variable is applicable when <code>I_MPI_THREAD_YIELD=1</code>. Small values of <code>I_MPI_PAUSE_COUNT</code> may increase performance, while larger values may reduce energy consumption.

I_MPI_THREAD_SLEEP

Control the Intel® MPI Library thread sleep microseconds timeout for thread yield customization while MPI busy wait progress.

Syntax

I MPI THREAD SLEEP=<arg>

Argument	Description
>=0	Thread sleep microseconds timeout. The default value is 0. Normally, the value
	is less than 100.

This variable is applicable when <code>I_MPI_THREAD_YIELD=3</code>. Small values of <code>I_MPI_PAUSE_COUNT</code> may increase performance in the normal mode, while larger values may increase performance in the oversubscription mode

I_MPI_EXTRA_FILESYSTEM

Control native support for parallel file systems.

Syntax

I MPI EXTRA FILESYSTEM=<arg>

Argument

Argument	Binary Indicator	
enable yes on 1	Enable native support for parallel file systems.	
disable no off 0	Disable native support for parallel file systems. This is the default value.	

Description

Use this environment variable to enable or disable native support for parallel file systems.

I_MPI_EXTRA_FILESYSTEM_FORCE

Syntax

I MPI EXTRA FILESYSTEM FORCE=<ufs|nfs|gpfs|panfs|lustre>

Description

Force filesystem recognition logic.

Miscellaneous

3

This topic provides the following information:

• Java* Bindings for MPI-2 Routines describes the Java language support by Intel® MPI Library.

Java* Bindings for MPI-2 Routines

Intel® MPI Library provides an experimental feature to enable support for Java* MPI applications. Intel MPI Library provides Java bindings for a subset of MPI-2 routines.

You can find all supported MPI routines in the table below. All the classes below belong to the mpi package.

NOTE

- For static methods, parameters fully correspond to the ones of C routines.
- For non-static methods, the object that calls the method corresponds to the OUT parameter of the original C routine.

Java* Bindings for MPI-2 Routines

Java Class	Public Fields and Methods	Original C Routine
MPI	<pre>static int Init(String[] args)</pre>	MPI_Init
	<pre>static void Finalize()</pre>	MPI_Finalize
	<pre>static double wTime()</pre>	MPI_Wtime
	<pre>static void abort(Comm comm, int errorCode)</pre>	MPI_Abort
	<pre>String getProcessorName()</pre>	MPI_Get_processor_name
Aint	<pre>static void getExtent(Datatype dt, Aint lb, Aint extent)</pre>	MPI_Type_get_extent
	<pre>static void getTrueExtent(Datatype dt, Aint true_lb, Aint true_extent)</pre>	MPI_Type_get_true_extent
	<pre>static void getAddress(long location, Aint address)</pre>	MPI_Get_address
	<pre>static void getContents(Datatype dt, int maxIntegers, int maxAddresses, int maxDatatypes, int[] integers, Aint[] addresses, Datatype[] datatypes)</pre>	MPI_Type_get_contents
Collective	static void allToAll(Object sendbuf, int sendcount, Datatype sendtype, Object recvbuf, int recvcount, Datatype recvtype, Comm comm)	MPI_Alltoall

Java Class	Public Fields and Methods	Original C Routine
	static void reduce(Object sendbuf, Object recvbuf, int count, Datatype type, Op op, int root, Comm comm)	MPI_Reduce
	<pre>static void bcast(Object buffer, int count, Datatype type, int root, Comm comm)</pre>	MPI_Bcast
	<pre>static void gather(Object sendBuffer, int sendCount, Datatype sendType, Object recvBuffer, int recvCount, Datatype recvType, int root, Comm comm)</pre>	MPI_Gather
	static void gatherv(Object sendBuffer, int sendCount, Datatype sendType, Object recvBuffer, Object recvCount, Object displs, Datatype recvType, int root, Comm comm)	MPI_Gatherv
	<pre>static void allGather(Object sendBuffer, int sendCount, Datatype sendType, Object recvBuffer, int recvCount, Datatype recvType, Comm comm)</pre>	MPI_Allgather
	static void allGatherv(Object sendBuffer, int sendCount, Datatype sendType, Object recvBuffer, Object recvCount, Object displs, Datatype recvType, Comm comm)	MPI_Allgatherv
	static void allReduce(Object sendbuf, Object recvbuf, int count, Datatype type, Op op, Comm comm)	MPI_Allreduce
	static void allToAllv(Object sendbuf, Object sendCount, Object sdispls, Datatype sendType, Object recvbuf, Object recvCount, Object rdispls, Datatype recvType, Comm comm)	MPI_Alltoallv
	static void reduceScatter(Object sendbuf, Object recvbuf, Object recvcounts, Datatype type, Op op, Comm comm)	MPI_Reduce_scatter
	static void scatter(Object sendBuffer, int sendCount, Datatype sendType, Object	MPI_Scatter

Java Class	Public Fields and Methods	Original C Routine
	<pre>recvBuffer, int recvCount, Datatype recvType, int root, Comm comm)</pre>	
	static void scatterv(Object sendBuffer, Object sendCount, Object displs, Datatype sendType, Object recvBuffer, int recvCount, Datatype recvType, int	MPI_Scatterv
	<pre>root, Comm comm) static void barrier(Comm</pre>	MPI Barrier
	comm)	
Comm	Static field: Comm WORLD	MPI COMM WORLD
	Static field: Comm SELF	MPI COMM SELF
	int getSize()	MPI Comm size
	int getRank()	MPI Comm rank
	Comm create(Group group)	MPI Comm create
	static Comm create(Comm comm, Group group)	MPI_Comm_create
	Comm dup()	MPI Comm dup
	Comm split(int color, int key)	MPI_Comm_split
Group	Static field: int MPI PROC NULL	MPI PROC NULL
	Static field: int MPI_IDENT	MPI_IDENT
	Static field: int MPI_CONGRUENT	MPI_CONGRUENT
	Static field: int MPI_SIMILAR	MPI_SIMILAR
	Static field: int MPI_UNEQUAL	MPI_UNEQUAL
	Static field: Group WORLD	MPI_GROUP_WORLD
	<pre>void group(Comm comm)</pre>	MPI_Comm_group
	<pre>int getSize()</pre>	MPI_Group_size
	<pre>int getRank()</pre>	MPI_Group_rank
	<pre>int MPI_Group_translate_ranks(i nt[] ranks1, Group group2, int[] ranks2)</pre>	MPI_Group_translate_ranks
	<pre>static int MPI_Group_translate_ranks(G roup group1, int[] ranks1, Group group2, int[] ranks2)</pre>	MPI_Group_translate_ranks
	<pre>int MPI_Group_compare(Group group2)</pre>	MPI_Group_compare
	<pre>int MPI_Group_union(Group group1, Group group2)</pre>	MPI_Group_union
	<pre>int MPI_Group_intersection(Grou p group1, Group group2)</pre>	MPI_Group_intersection
	<pre>int MPI_Group_difference(Group group1, Group group2)</pre>	MPI_Group_difference
	<pre>int MPI_Group_incl(Group group, int n, int[] ranks)</pre>	MPI_Group_incl
	<pre>int MPI_Group_excl(Group group, int n, int[] ranks)</pre>	MPI_Group_excl

Java Class	Public Fields and Methods	Original C Routine
Datatype	Static field: Datatype NULL	MPI_DATATYPE_NULL
	Static field: Datatype BYTE	MPI UINT8 T
	Static field: Datatype CHAR	MPI CHAR
	Static field: Datatype SHORT	MPI INT16 T
	Static field: Datatype BOOLEAN	MPI UINT8 T
	Static field: Datatype INT	MPI INT32 T
	Static field: Datatype LONG	MPI INT64 T
	Static field: Datatype FLOAT	MPI FLOAT
	Static field: Datatype DOUBLE	MPI DOUBLE
	Static field: Datatype PACKED	MPI PACKED
	Static field: Datatype INT2	MPI 2INT
	Static field: Datatype SHORT INT	MPI SHORT INT
	Static field: Datatype LONG INT	MPI LONG INT
	Static field: Datatype FLOAT INT	MPI FLOAT INT
	Static field: Datatype	MPI DOUBLE INT
	DOUBLE_INT	
	Static field: Datatype FLOAT_COMPLEX	MPI_C_FLOAT_COMPLEX
	Static field: Datatype DOUBLE_COMPLEX	MPI_C_DOUBLE_COMPLEX
	<pre>void contiguous(int count, Datatype type)</pre>	MPI_Type_contiguous
	<pre>void commit()</pre>	MPI_Type_commit
	<pre>int getTypeSize()</pre>	MPI_Type_size
	<pre>void free()</pre>	MPI_Type_free
	<pre>void vector(int count, int blockLength, int stride, Datatype baseType)</pre>	MPI_Type_vector
	<pre>void hvector(int count, int blockLength, int stride, Datatype oldType)</pre>	MPI_Type_create_hvector
	<pre>void indexed(int count, int[] blockLength, int[] displacement, Datatype oldType)</pre>	MPI_Type_indexed
	<pre>void hindexed(int count, int[] blockLength, Aint[] displacement, Datatype oldType)</pre>	MPI_Type_create_hindexed
	<pre>void struct(int count, int[] blockLength, Aint[] displacement, Datatype[] oldTypes)</pre>	MPI_Type_struct
Ор	Static field: Op MPI_OP_NULL	MPI OP NULL
-r	Static field: Op MPI MAX	MPI MAX
	Static field: Op MPI MIN	MPI MIN
	Static field: Op MPI SUM	MPI SUM
	Static field: Op MPI PROD	MPI PROD
	Static field: Op MPI LAND	MPI LAND
	Static field: Op MPI_BAND	_
	Static field: On MPT BAND	MPI BAND

Java Class	Public Fields and Methods	Original C Routine
	Static field: Op MPI_BOR	MPI_BOR
	Static field: Op MPI LXOR	MPI LXOR
	Static field: Op MPI BXOR	MPI BXOR
	Static field: Op MPI MINLOC	MPI MINLOC
	Static field: Op MPI MAXLOC	MPI MAXLOC
	Op(UserFunction uf)	_
	void	-
	<pre>setUserFunction(UserFunctio n userFunction)</pre>	
	<pre>void createOP(boolean commute)</pre>	MPI_Op_Create
UserFunction (abstract)	<pre>UserFunction(Datatype type, int length)</pre>	-
	<pre>void setInoutvec(ByteBuffer inoutvec)</pre>	-
	<pre>void setInvec(ByteBuffer invec)</pre>	-
	abstract void call(int type, int length)	-
PTP	<pre>static void send(Buffer buffer, int count, Datatype type, int dest, int tag, Comm comm)</pre>	MPI_Send
	<pre>static void send(<java array=""> buffer, int count, Datatype type, int dest, int tag, Comm comm)</java></pre>	MPI_Send
	static Status recv(Buffer buf, int count, Datatype type, int source, int tag, Comm comm)	MPI_Recv
	,	MPI_Recv
	static Request isend(Buffer buffer, int count, Datatype type, int dest, int tag, Comm comm)	MPI_Isend
	<pre>static Request isend(<java array=""> buffer, int count, Datatype type, int dest, int tag, Comm comm)</java></pre>	MPI_Isend
	static Request irecv(Buffer buf, int count, Datatype type, int source, int tag, Comm comm)	MPI_Irecv
	static Request irecv(<java array=""> buf, int count, Datatype type, int source, int tag, Comm comm)</java>	MPI_Irecv
	static Status sendRecv(Buffer sendbuf, int sendcount, Datatype	MPI_Sendrecv

Java Class	Public Fields and Methods	Original C Routine
	sendtype, int senddest, int sendtag, Buffer recvbuf, int recvcount, Datatype recvtype, int recvsource, int recvtag, Comm comm)	
Request	Status Wait()	MPI_Wait
	<pre>static Status[] waitAll(int count, Request[] reqs)</pre>	MPI_Waitall
	<pre>static Status waitAny(int count, Request[] reqs, int[] index)</pre>	MPI_Waitany
	<pre>static Status[] waitSome(int count, Request[] reqs, int[] outcount, int[] indexes)</pre>	MPI_Waitsome
	boolean test(Status status)	MPI_Test

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