SGI MPI and SGI SHMEM™ User Guide
New Features in This Manual

This revision adds the following information:

• Compatibility information.
• Miscellaneous technical and editorial corrections.
# Record of Revision

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<th>Version</th>
<th>Description</th>
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<td></td>
<td>Original Printing. This manual documents the Message Passing Toolkit implementation of the Message Passing Interface (MPI).</td>
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<td>Supports the MPT 1.11 release.</td>
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<td>April 2010</td>
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<td>February 2014</td>
<td>Supports the SGI Performance Suite 1.7 release, the MPT 2.09 release, and the MPI 1.7 release. Clarifies SLURM support.</td>
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025 June 2014
Supports the SGI Performance Suite 1.8 release, the SGI MPT 2.10 release, and the SGI MPI 1.8 release. This is the last revision of this documentation with the title *Message Passing Toolkit (MPT) User Guide*.

026 May 2015
Supports the SGI Performance Suite 1.10 release, the SGI MPT 2.12 release, and the SGI MPI 1.10 release. This documentation is now called the *SGI MPI and SGI SHMEM User Guide*.

027 November 2015
Supports the SGI Performance Suite 1.11 release, the SGI MPT 2.13 release, and the SGI MPI 1.11 release.

028 May 2016
Supports the SGI Performance Suite 1.12 release, the SGI MPT 2.14 release, and the SGI MPI 1.12 release.

029 May 2016
Supports the SGI Performance Suite 1.12 release, the SGI MPT 2.14 release, and the SGI MPI 1.12 release and adds third-party compatibility information.
## Contents

**About This Guide** ............................................. xix  
Compatibility Information ...................................... xx  
Related SGI Publications ....................................... xxi  
Related Publications From Other Sources .................. xxii  
Obtaining Publications ........................................ xxii  
Conventions ....................................................... xxiii  
Reader Comments ............................................... xxiii  

**1. Configuring the SGI Message Passing Toolkit (MPT)** ........ 1  
About Configuring SGI MPT ..................................... 1  
Configuring SGI MPT on an SGI UV Computer System (Single System Image) ........................................... 2  
  Verifying Prerequisites ....................................... 2  
  (Optional) Installing the SGI MPT Software Into a Nondefault Working Directory ............................. 3  
  Adjusting File Resource Limits ............................ 5  
  Completing the Configuration ............................ 7  
Configuring SGI MPT on an SGI UV Computer System (Partitioned) ....................................................... 7  
  Verifying Prerequisites ..................................... 8  
  Configuring the OpenFabrics Enterprise Distribution (OFED) Software ............................................ 9  
  Adjusting File Resource Limits .......................... 11  
  Creating a Directory and Removing the Current Software .............................................................. 12  
  (Optional) Configuring the MUNGE Security Software ................................................................. 14  
  Updating Other Partitions or Continuing the Configuration ......................................................... 15  
  Configuring Array Services .................................. 15  
  Enabling Cross-partition NUMAlink MPI Communication and Restarting Services ........................ 18
## Contents

<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Enabling Cross-partition Communication and Restarting Services (RHEL)</td>
<td>18</td>
</tr>
<tr>
<td>Enabling Cross-partition Communication and Restarting Services (SLES)</td>
<td>19</td>
</tr>
<tr>
<td>Completing the Configuration</td>
<td>20</td>
</tr>
<tr>
<td><strong>2. Getting Started</strong></td>
<td>21</td>
</tr>
<tr>
<td>About Running MPI Applications</td>
<td>21</td>
</tr>
<tr>
<td>Loading the MPI Software Module and Specifying the Library Path</td>
<td>21</td>
</tr>
<tr>
<td>Compiling and Linking the MPI Program</td>
<td>23</td>
</tr>
<tr>
<td>Compiling With the Wrapper Compilers</td>
<td>23</td>
</tr>
<tr>
<td>Compiling With the GNU or Intel Compilers</td>
<td>24</td>
</tr>
<tr>
<td>Launching the MPI Application</td>
<td>25</td>
</tr>
<tr>
<td>Using a Workload Manager to Launch an MPI Application</td>
<td>25</td>
</tr>
<tr>
<td>PBS Professional</td>
<td>25</td>
</tr>
<tr>
<td>Torque</td>
<td>26</td>
</tr>
<tr>
<td>Simple Linux Utility for Resource Management (SLURM)</td>
<td>27</td>
</tr>
<tr>
<td>Using the <code>mpirun</code> Command to Launch an MPI Application</td>
<td>27</td>
</tr>
<tr>
<td>Launching a Single Program on the Local Host</td>
<td>27</td>
</tr>
<tr>
<td>Launching a Multiple Program, Multiple Data (MPMD) Application on the Local Host</td>
<td>28</td>
</tr>
<tr>
<td>Launching a Distributed Application</td>
<td>28</td>
</tr>
<tr>
<td>Launching an Application by Using MPI Spawn Functions</td>
<td>29</td>
</tr>
<tr>
<td>Compiling and Running SHMEM Applications</td>
<td>29</td>
</tr>
<tr>
<td>Using Huge Pages</td>
<td>30</td>
</tr>
<tr>
<td>Using SGI MPI in an SELinux Environment (RHEL Platforms Only)</td>
<td>32</td>
</tr>
<tr>
<td><strong>3. Programming With SGI MPI</strong></td>
<td>33</td>
</tr>
<tr>
<td>About Programming With SGI MPI</td>
<td>33</td>
</tr>
<tr>
<td>Job Termination and Error Handling</td>
<td>33</td>
</tr>
</tbody>
</table>
SGI MPI and SGI SHMEM™ User Guide

MPI_Abort ........................................... 34
Error Handling ........................................ 34
MPI_Finalize and Connect Processes .................. 34
Signals ............................................... 35
Buffering .......................................... 35
Multithreaded Programming ............................ 36
Interoperability with the SHMEM programming model . 37
Miscellaneous SGI MPI Features ....................... 37
Programming Optimizations ............................ 38
Using MPI Point-to-Point Communication Routines 38
Using MPI Collective Communication Routines 39
Using MPI_Pack/MPI_Unpack ........................... 39
Avoiding Derived Data Types ........................... 40
Avoiding Wild Cards .................................. 40
Avoiding Message Buffering — Single Copy Methods 40
Managing Memory Placement ......................... 41
Additional Programming Model Considerations .......... 41

4. Debugging MPI Applications ........................ 43
MPI Routine Argument Checking ........................ 43
Using the TotalView Debugger with MPI Programs .......... 43
Using idb and gdb with MPI Programs .................. 44
Using the DDT Debugger with MPI Programs .......... 44
Using Valgrind With MPI Programs ........................ 45

5. Using PerfBoost ........................................ 47
About PerfBoost ........................................ 47
Using PerfBoost ......................................... 47
Contents

MPI Supported Functions .................................................. 48

6. Berkeley Lab Checkpoint/Restart ........................................ 51
   About Berkeley Lab Checkpoint/Restart ............................... 51
   BLCR Installation ......................................................... 51
   Using BLCR with SGI MPT ............................................... 52

7. Run-time Tuning ............................................................. 53
   About Run-time Tuning ..................................................... 53
   Reducing Run-time Variability ......................................... 54
   Tuning MPI Buffer Resources .......................................... 55
   Avoiding Message Buffering – Enabling Single Copy ................. 56
      Buffering and MPI_Send ............................................... 56
      Using the XPMEM Driver for Single Copy Optimization .......... 56
   Memory Placement and Policies ........................................ 57
      MPI_DSM_CPULIST ..................................................... 57
      MPI_DSM_DISTRIBUTE ............................................... 58
      MPI_DSM_VERBOSE .................................................... 59
      Using dplace .......................................................... 59
   Tuning MPI/OpenMP Hybrid Codes ..................................... 59
   Tuning Running Applications Across Multiple Hosts ................. 61
   Tuning for Running Applications over the InfiniBand Interconnect ...................................................... 63
   MPI on SGI UV Systems .................................................. 65
      General Considerations ............................................... 66
      Performance Problems and Corrective Actions ..................... 66
      Other ccNUMA Performance Considerations ......................... 67
   Suspending MPI Jobs ...................................................... 68
8. MPI Performance Profiling ............................................. 71
  About MPI Performance Profiling ................................... 71
  Using perfcatch(1) .................................................. 72
    The perfcatch(1) Command ....................................... 72
    MPI_PROFILING_STATS Results File Example .................. 73
    Environment Variables Used With perfcatch(1) .............. 76
  Writing Your Own Profiling Interface ............................ 77
  Using Third-party Profilers ....................................... 78
  MPI Internal Statistics ............................................ 78

9. Troubleshooting and Frequently Asked Questions ................. 81
  What are some things I can try to figure out why mpirun is failing? .................. 81
  My code runs correctly until it reaches MPI_Finalize() and then it hangs. ............ 83
  My hybrid code (using OpenMP) stalls on the mpirun command. ......................... 83
  I keep getting error messages about MPI_REQUEST_MAX being too small. ............... 83
  I am not seeing stdout and/or stderr output from my MPI application. ............... 84
  How can I get the SGI Message Passing Toolkit (MPT) software to install on my machine? .................. 84
  Where can I find more information about the SHMEM programming model? ............... 84
  The ps(1) command says my memory use (SIZE) is higher than expected. ............... 84
  What does MPI: could not run executable mean? ................................ 85
  How do I combine MPI with insert favorite tool here? ................................ 85
  Why do I see “stack traceback” information when my MPI job aborts? .................. 86

10. Array Services ...................................................... 87
  About Array Services ................................................ 87
  Retrieving the Array Services Release Notes ................................ 88
  Managing Local Processes .......................................... 89
    Monitoring Local Processes and System Usage .................... 89
<table>
<thead>
<tr>
<th>Contents</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scheduling and Killing Local Processes</td>
<td>89</td>
</tr>
<tr>
<td>Summary of Local Process Management Commands</td>
<td>90</td>
</tr>
<tr>
<td>Using Array Services Commands</td>
<td>90</td>
</tr>
<tr>
<td>About Array Sessions</td>
<td>91</td>
</tr>
<tr>
<td>About Names of Arrays and Nodes</td>
<td>91</td>
</tr>
<tr>
<td>About Authentication Keys</td>
<td>91</td>
</tr>
<tr>
<td>Array Services Commands</td>
<td>91</td>
</tr>
<tr>
<td>Specifying a Single Node</td>
<td>93</td>
</tr>
<tr>
<td>Common Environment Variables</td>
<td>93</td>
</tr>
<tr>
<td>Obtaining Information About the Array</td>
<td>94</td>
</tr>
<tr>
<td>Learning Array Names</td>
<td>94</td>
</tr>
<tr>
<td>Learning Node Names</td>
<td>95</td>
</tr>
<tr>
<td>Learning Node Features</td>
<td>95</td>
</tr>
<tr>
<td>Learning User Names and Workload</td>
<td>96</td>
</tr>
<tr>
<td>Learning User Names</td>
<td>96</td>
</tr>
<tr>
<td>Learning Workload</td>
<td>96</td>
</tr>
<tr>
<td>Additional Array Configuration Information</td>
<td>97</td>
</tr>
<tr>
<td>Security Considerations for Standard Array Services</td>
<td>97</td>
</tr>
<tr>
<td>About the Uses of the Configuration Files</td>
<td>98</td>
</tr>
<tr>
<td>About Configuration File Format and Contents</td>
<td>99</td>
</tr>
<tr>
<td>Loading Configuration Data</td>
<td>100</td>
</tr>
<tr>
<td>About Substitution Syntax</td>
<td>101</td>
</tr>
<tr>
<td>Testing Configuration Changes</td>
<td>101</td>
</tr>
<tr>
<td>Specifying Arrayname and Machine Names</td>
<td>102</td>
</tr>
<tr>
<td>Specifying IP Addresses and Ports</td>
<td>102</td>
</tr>
<tr>
<td>Specifying Additional Attributes</td>
<td>103</td>
</tr>
</tbody>
</table>
Configuring Array Commands ........................................... 103
  Operation of Array Commands ........................................ 104
  Summary of Command Definition Syntax .............................. 104
  Configuring Local Options ........................................... 107
  Designing New Array Commands ...................................... 108

11. Using the SGI MPT Plugin for Nagios .............................. 111
  About the SGI MPT Plugin for Nagios ................................. 111
  Installing the SGI MPT Nagios Plugin on the Admin Node ........ 112
  (Optional) Installing the SGI MPT Nagios Plugin on a Rack Leader Controller (RLC) Node 115
  Viewing SGI MPT Messages From Within Nagios and Clearing the Messages ...................... 116
  (Optional) Modifying the Notification Email ........................ 119

Appendix A. Guidelines for Using SGI MPT on a Virtual Machine Within an SGI UV Computer System ........................................ 121
  About SGI MPT on a Virtual Machine ............................... 121
  Installing Software Within the Virtual Machine (VM) .............. 121
  Adjusting SGI UV Virtual Machine System Settings ................. 122
  Running SGI MPI Programs From Within a Virtual Machine (VM) ........................................ 124

Appendix B. Configuring Array Services Manually .................. 125
  About Configuring Array Services Manually ........................ 125
  Configuring Array Services on Multiple Partitions or Hosts ........ 125

Index ................................................................. 129
Tables

**Table 1-1** Array Configuration Resources ............................................. 16
**Table 3-1** Outline of Improper Dependence on Buffering ............................ 36
**Table 7-1** Available Interconnects and the Inquiry Order for Available Interconnects ..... 61
**Table 10-1** Information Sources: Local Process Management ...................... 90
**Table 10-2** Common Array Services Commands ........................................ 90
**Table 10-3** Array Services Command Option Summary .............................. 92
**Table 10-4** Array Services Environment Variables .................................... 94
**Table 10-5** Subentries of a COMMAND Definition .................................... 105
**Table 10-6** Substitutions Used in a COMMAND Definition .......................... 106
**Table 10-7** Options of the COMMAND Definition ..................................... 106
**Table 10-8** Subentries of the LOCAL Entry ............................................. 107
About This Guide

The Message Passing Interface (MPI) standard supports C and Fortran programs with a library and supporting commands. MPI operates through a technique known as message passing, which is the use of library calls to request data delivery from one process to another or between groups of processes. MPI also supports parallel file I/O and remote memory access (RMA).

The SGI MPI software supports the MPI standard. SGI MPI facilitates parallel programming on large systems and on computer system clusters. This publication describes SGI MPI 1.12, which supports the MPI 3.1 standard. SGI MPI includes significant features that make it the preferred implementation for use on SGI hardware. The following are some of these features:

- Data transfer optimizations for NUMAlink, where available, including single-copy data transfer.
- Multirail InfiniBand support, which takes full advantage of the multiple InfiniBand fabrics available on SGI® ICETM systems.
- Optimized MPI remote memory access (RMA) one-sided commands.
- Interoperability with the SHMEM (LIBSMA) programming model.

SGI MPI also supports the OpenSHMEM standard. The OpenSHMEM standard describes a low-latency library that supports RMA on symmetric memory in parallel environments. The OpenSHMEM programming model is a partitioned global address space (PGAS) programming model that presents distributed processes with symmetric arrays that are accessible via PUT and GET operations from other processes. This publication describes SGI SHMEM, which supports OpenSHMEM version 1.3. The SGI SHMEM programming model is the basis for the OpenSHMEM™ programming model specification that is being developed by the Open Source Software Solutions multivendor working group.

SGI’s support for MPI and OpenSHMEM is built on top of the SGI Message Passing Toolkit (MPT). SGI MPT is a high-performance communications middleware software product that runs on SGI’s shared memory and cluster supercomputers. On some of these machines, SGI MPT uses SGI Array Services to launch applications. SGI MPT is optimized for all SGI hardware platforms. This document describes SGI MPT 2.14.
# Compatibility Information

The following table describes compatibility between SGI MPI 1.12 and other software products.

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<th>Technology</th>
<th>Notes</th>
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<td>Red Hat Enterprise Linux (RHEL) operating system</td>
<td>RHEL 6.X and RHEL 7.X</td>
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<td>SLES operating system</td>
<td>SLES 12 SPX and SLES 11 SPX</td>
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<tr>
<td>CentOS operating system</td>
<td>CentOS 7.X</td>
</tr>
<tr>
<td>Fortran 2008</td>
<td>Supports Fortran 2008 for Intel compilers only.</td>
</tr>
<tr>
<td>SGI compute platforms</td>
<td>Supports all SGI compute platforms:</td>
</tr>
<tr>
<td></td>
<td>• SGI UV systems</td>
</tr>
<tr>
<td></td>
<td>• SGI ICE systems</td>
</tr>
<tr>
<td></td>
<td>• SGI Rackable systems</td>
</tr>
<tr>
<td>Multi-rail InfiniBand (IB)</td>
<td>No support for MPI spawn</td>
</tr>
<tr>
<td>Multi-rail Intel Omni-Path Architecture (OPA) TCP/IP communication</td>
<td></td>
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<tr>
<td>Mellanox Fabric Collective Accelerator (FCA) 3.x / HCOLL</td>
<td>Requires Mellanox Open Fabrics Enterprise Distribution (OFED).</td>
</tr>
<tr>
<td>NVIDIA GPUDirect remote direct memory access (RDMA) over IB</td>
<td>No support for MPI RMA passive windows.</td>
</tr>
<tr>
<td>Checkpoint-restart (CPR), supported through Berkeley Lab checkpoint restart (BLCR).</td>
<td>Supports jobs running over shared memory, IB, TCP/IP, and the SGI UV global resource unit (GRU). No support for CPR when using the following:</td>
</tr>
<tr>
<td></td>
<td>• OpenSHMEM</td>
</tr>
<tr>
<td></td>
<td>• MPI remote memory access (RMA) passive windows</td>
</tr>
<tr>
<td></td>
<td>• MPI Spawn</td>
</tr>
<tr>
<td></td>
<td>• Process management interface (PMI), which is commonly used by SLURM</td>
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</tbody>
</table>
### Third-party debugging and profiling tools:

- Allinea DDT
- RogueWave TotalView
- Tuning and Analysis Utilities (TAU)
- Vampir

Contact SGI for information about additional debugging and profiling tools.

### Third-party workload managers:

- Altair PBS Professional
- SLURM
- UNIVA Grid Engine
- IBM LSF
- Moab / TORQUE

Supported when running under the simple linux utility for resource management (SLURM).

### Related SGI Publications

The SGI Foundation Software release notes and the SGI Performance Suite release notes contain information about the specific software packages provided in those products. The release notes also list SGI publications that provide information about the products. The release notes are available in the following locations:

- **Online at the SGI customer portal.** After you log into the SGI customer portal, you can access the release notes. The SGI Foundation Software release notes are posted to the following website:
  
  https://support1-sgi.custhelp.com/app/answers/detail/a_id/4983

  The SGI Performance Suite release notes are posted to the following website:
  
  https://support1-sgi.custhelp.com/app/answers/detail/a_id/6093

  **Note:** You must sign into the SGI customer portal, at https://support.sgi.com/login, in order for the preceding links to work.

- **On the product media.** The release notes reside in a text file in the /docs directory on the product media. For example, /docs/SGI-MPI-1.x-readme.txt.
• On the system. After installation, the release notes and other product documentation reside in the /usr/share/doc/packages/product directory.

The MPInside Reference Guide describes SGI’s MPInside MPI profiling tool.

SGI creates hardware manuals that are specific to each product line. The hardware documentation typically includes a system architecture overview and describes the major components. It also provides the standard procedures for powering on and powering off the system, basic troubleshooting information, and important safety and regulatory specifications.

Related Publications From Other Sources

Information about MPI is available from a variety of sources. For information about the MPI standard, see the following:

• The Message Passing Interface Forum’s website, which is as follows:
  http://www.mpi-forum.org/


• Journal articles in the following publications:

Obtaining Publications

All SGI publications are available on the SGI customer portal at http://support.sgi.com. Select the following:

Support by Product > productname > Documentation
If you do not find what you are looking for, search for document-title keywords by selecting Search Knowledgebase and using the category Documentation.

You can view man pages by typing man title on a command line.

Conventions

The following conventions are used throughout this document:

<table>
<thead>
<tr>
<th>Convention</th>
<th>Meaning</th>
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<tbody>
<tr>
<td>command</td>
<td>This fixed-space font denotes literal items such as commands, files, routines, path names, signals, messages, and programming language structures.</td>
</tr>
<tr>
<td>manpage(x)</td>
<td>Man page section identifiers appear in parentheses after man page names.</td>
</tr>
<tr>
<td>variable</td>
<td>Italic typeface denotes variable entries and words or concepts being defined.</td>
</tr>
<tr>
<td>user input</td>
<td>This bold, fixed-space font denotes literal items that the user enters in interactive sessions. (Output is shown in nonbold, fixed-space font.)</td>
</tr>
<tr>
<td>[ ]</td>
<td>Brackets enclose optional portions of a command or directive line.</td>
</tr>
<tr>
<td>...</td>
<td>Ellipses indicate that a preceding element can be repeated.</td>
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</tbody>
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Reader Comments

If you have comments about the technical accuracy, content, or organization of this publication, contact SGI. Be sure to include the title and document number of the publication with your comments. (Online, the document number is located in the front matter of the publication. In printed publications, the document number is located at the bottom of each page.)

You can contact SGI in either of the following ways:

- Send e-mail to the following address:
• Contact your customer service representative and ask that an incident be filed in the SGI incident tracking system:
  http://www.sgi.com/support/supportcenters.html

SGI values your comments and will respond to them promptly.
Chapter 1

Configuring the SGI Message Passing Toolkit (MPT)

This chapter includes the following topics:

• "About Configuring SGI MPT" on page 1
• "Configuring SGI MPT on an SGI UV Computer System (Single System Image)" on page 2
• "Configuring SGI MPT on an SGI UV Computer System (Partitioned)" on page 7

About Configuring SGI MPT

When you installed SGI Performance Suite, you also installed SGI MPT. Before you can run any SGI MPI programs, however, you need to configure the SGI MPT software. The procedures in this chapter explain how to configure SGI MPT.

SGI computers often host several released versions of SGI MPT. This environment provides users with the flexibility they need to develop and run MPI programs. The configuration instructions in this chapter explain how to accommodate these multiple versions if your site needs to have multiple versions installed.

The configuration procedure differs, depending on your platform, as follows:

• On a standalone SGI UV computer system, the configuration procedure differs depending on whether your system is partitioned or not, as follows:
  – If you have an SGI UV system that is configured as a single system image (SSI), complete the following procedure:
    "Configuring SGI MPT on an SGI UV Computer System (Single System Image)" on page 2
  – If you have an SGI UV system that is configured into two or more partitions, complete the following procedure:
    "Configuring SGI MPT on an SGI UV Computer System (Partitioned)" on page 7
• On an SGI cluster computing system, such as an SGI® ICE™ cluster or an SGI Rackable™ cluster, the configuration procedure includes image-management steps
that this chapter does not address. For information about how to configure SGI MPT on an SGI cluster computer, see the following:

SGI Management Center Installation and Configuration Guide for Clusters

Configuring SGI MPT on an SGI UV Computer System (Single System Image)

The information in the following procedures explains how to configure SGI MPT on a large, single SGI UV SSI:

• "Verifying Prerequisites" on page 2
• "(Optional) Installing the SGI MPT Software Into a Nondefault Working Directory" on page 3
• "Adjusting File Resource Limits" on page 5
• "Completing the Configuration" on page 7

Verifying Prerequisites

The following procedure explains how to verify the SGI MPT software’s installation prerequisites.

Procedure 1-1 To verify prerequisites

1. As the root user, log into the SGI UV computer.

2. (Conditional) Reboot the computer.

   Perform this step if the SGI UV computer was not rebooted after SGI Performance Suite was installed.

   If you do not know whether the computer has been rebooted, reboot at this time.

3. Verify that you have one of the following operating system software packages installed and configured:

   • Red Hat Enterprise Linux (RHEL) 6 or 7
   • SLES 11 or 12
You can type the following command to verify your operating system version:

```
# cat /etc/issue
```

4. Type a series of `cat(1)` commands to verify that the following required products from the SGI Performance Suite 1.12 release are installed:

- SGI Accelerate
- SGI MPI

For example:

```
# cat /etcsgi-accelerate-release
SGI Performance Suite 1.12, Build xxxxxxx.sles11sp4-xxxxxxxxxx
# cat /etc/sgi-mpi-release
SGI MPI 1.12, Build xxxxxxx.sles11sp4-xxxxxxxxxx
```

5. Proceed to one of the following:

- "(Optional) Installing the SGI MPT Software Into a Nondefault Working Directory" on page 3, which explains how to configure SGI MPT in a way that lets you maintain more than one released version of the software on your SGI UV computer system.

- "Adjusting File Resource Limits" on page 5, which assumes you want the SGI MPT software to remain in the default installation directory.

**(Optional) Installing the SGI MPT Software Into a Nondefault Working Directory**

Perform the procedure in this topic if you want to install SGI MPT into a custom, nondefault working directory. You might want to perform the procedure in this topic if, for example, you have a nondefault filesystem.

The RPM utility enables you to create, install, and manage relocatable packages. You can install a matched set of SGI MPT RPMs in either a default location or an alternate location. The default location for installing the SGI MPT RPM is `/opt/sgi/mpt/mpt-2.rel_level`. To install the SGI MPT RPM in an alternate location, use the `--relocate` parameter to the `rpm` command. The `--relocate` parameter specifies an alternate base directory for the SGI MPT software installation.

Either `/opt/sgi/mpt/mpt-2.rel_level` or both `/opt/sgi/mpt/mpt-2.rel_level` and `/usr/share/modules/modulefiles/mpt` can be relocated. The post installation
script reconfigures the module file for the new location as long as the \textit{oldpath} precisely matches the description in the RPM info.

The general format for the \texttt{rpm} command is as follows:

\begin{verbatim}
rpm --relocate oldpath=newpath
\end{verbatim}

- For \textit{oldpath}, specify the SGI MPT software’s current location.
  If you install the SGI MPT software in an alternate location, the \texttt{rpm} command’s \textit{oldpath} argument must precisely match the relocation listed in the RPM for the environment module automatic modification feature to be correct.

- For \textit{newpath}, specify the location to which you want to install the SGI MPT software.

\textbf{Procedure 1-2} To install the SGI MPT software in an alternate location

1. Plan how to avoid problems related to uninstalled RPM dependencies.

   The following are two approaches:

   - Option 1: If you install from a system that does not run MPT jobs, it might be appropriate to use the \texttt{--nodeps} parameter on the \texttt{rpm(8)} command line.
     This parameter directs the \texttt{rpm(8)} command to ignore dependencies.

   - Option 2: If you install from a system or cluster nodes upon which MPT jobs need to run, type the following package manager commands on each cluster node or cluster node image to locally install the needed prerequisites on all the cluster nodes:
     - On SLES platforms, type the following command:
       \begin{verbatim}
       # zypper install cpuset-utils arraysvcs xpmem libbitmask
       \end{verbatim}
     - On RHEL platforms, type the following command:
       \begin{verbatim}
       # yum install cpuset-utils arraysvcs xpmem libbitmask
       \end{verbatim}

2. Use the \texttt{rpm} command to specify an alternate location for the SGI MPT software bundle.

   Example 1. The following example shows how to install SGI MPT in
   \texttt{/usr/local/sgi/mpt/mpt-2.14} rather than in \texttt{/opt}, which is the default:

   \begin{verbatim}
   # rpm -i --relocate /opt/sgi/mpt/mpt-2.14=/usr/local/sgi/mpt/mpt-2.14 \ sgi-mpt-*.*86_64.rpm
   \end{verbatim}
Example 2: The following RHEL example shows how to install the modules, in addition to the total SGI MPT software bundle, to 
/usr/local/sgi/mpt/mpt-2.14 and /usr/local/mod/mpt:

```
# rpm -i --relocate /opt/sgi/mpt/mpt-2.14=/usr/local/sgi/mpt/mpt-2.14 \
--relocate /usr/share/Modules/modulefiles/mpt=/usr/local/mod/mpt \
sgi-mpt-*.x86_64.rpm
```

In the preceding RHEL example, note that the Modules directory in the argument to the second --relocate parameter begins with an uppercase letter.

Example 3. The following SLES example shows how to install the modules, in addition to the total SGI MPT software bundle, to 
/usr/local/sgi/mpt/mpt-2.14 and /usr/local/mod/mpt:

```
# rpm -i --relocate /opt/sgi/mpt/mpt-2.14=/usr/local/sgi/mpt/mpt-2.14 \
--relocate /usr/share/modules/modulefiles/mpt=/usr/local/mod/mpt \
sgi-mpt-*.x86_64.rpm
```

In the preceding SLES example, note that the modules directory in the argument to the second --relocate parameter begins with a lowercase letter.

Example 4:

The following example rpm command output shows the available relocations:

```
# rpm -qpi sgi-mpt-2.14-sgi*.x86_64.rpm
... Relocations: /opt/sgi/mpt/mpt-2.14 /usr/share/modules/modulefiles/mpt
```

**Note:** In the preceding output, the example shows only the significant message at the end of the output string.

3. Proceed to the following:

"Adjusting File Resource Limits" on page 5

For more information about using the rpm command, see the rpm man page.

**Adjusting File Resource Limits**

The following procedure explains how to increase resource limits on the number of open files and enforce new security policies.
Procedure 1-3 To adjust file resource limits

1. Type the following command to retrieve the number of cores on this computer:

   
   ```bash
   # cat /proc/cpuinfo | grep processor | wc -l
   ```

   In the preceding line, the last character is a lowercase L, not the number 1.
   
   This `cat(1)` command returns the number of cores on the SGI UV computer system.

2. Use a text editor to open file `/etc/security/limits.conf`.

3. Add the following line to file `/etc/security/limits.conf`:

   ```plaintext
   * hard nofile limit
   ```

   For `limit`, specify an open file limit, for the number of MPI processes per host, based on the following guidelines:

<table>
<thead>
<tr>
<th>Processes/host</th>
<th>limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fewer than 512</td>
<td>3000</td>
</tr>
<tr>
<td>Up to 1024</td>
<td>6000</td>
</tr>
<tr>
<td>Up to 2048</td>
<td>8192 (default)</td>
</tr>
<tr>
<td>4096 or more</td>
<td>21000</td>
</tr>
</tbody>
</table>

   MPI jobs require a large number of file descriptors, and on larger systems, you might need to increase the system-wide limit on the number of open files. The default value for the file-limit resource is 8192. For example, the following line is suitable for 512 MPI processes per host:

   ```plaintext
   * hard nofile 3000
   ```

4. Add the following line to file `/etc/security/limits.conf`:

   ```plaintext
   * hard memlock unlimited
   ```

   The preceding line increases the resource limit for locked memory.

5. Save and close file `/etc/security/limits.conf`.

6. Use a text editor to open file `/etc/pam.d/login`, which is the Linux pluggable authentication module (PAM) configuration file.
7. Add the following line to file /etc/pam.d/login:

   session required /lib/security/pam_limits.so

8. Save and close the file.

9. (Conditional) Update other authentication configuration files as needed.
   Perform this step if your site allows other login methods, such as ssh, rlogin, and so on.

10. Proceed to the following:
    "Completing the Configuration" on page 7

**Completing the Configuration**

The following procedure explains how to complete the SGI MPT configuration.

**Procedure 1-4 To complete the SGI MPT configuration**

1. Run a test MPI program to make sure that the new software is working as expected.

2. (Conditional) Inform your user community of the location of the new SGI MPT release on this computer.
   Perform this step if you moved the SGI MPT software to a nondefault location.
   In this procedure’s examples, the module files are located in the following directories:
   • On RHEL platforms:
     /opt/mpt/mpt-2.14/usr/share/Modules/modulefiles/mpt/mpt-2.14
   • On SLES platforms:
     /opt/mpt/mpt-2.14/usr/share/modules/modulefiles/mpt/mpt-2.14

**Configuring SGI MPT on an SGI UV Computer System (Partitioned)**

You can configure SGI MPT on an SGI UV computer system that is divided into two or more partitions. Generally, you configure each partition individually, and then you configure the partitions into an array. If you have several partitions, you can use only
some of them for the SGI MPT array; you do not have to configure all the partitions into an array.

The information in the following procedures explains how to configure SGI MPT on a partitioned SGI UV server:

- "Verifying Prerequisites" on page 8
- "Configuring the OpenFabrics Enterprise Distribution (OFED) Software" on page 9
- "Adjusting File Resource Limits" on page 11
- "Creating a Directory and Removing the Current Software" on page 12
- "(Optional) Configuring the MUNGE Security Software" on page 14
- "Updating Other Partitions or Continuing the Configuration" on page 15
- "Configuring Array Services" on page 15
- "Enabling Cross-partition NUMAlink MPI Communication and Restarting Services" on page 18
- "Completing the Configuration" on page 20

Verifying Prerequisites

The following procedure explains how to verify the SGI MPT software’s configuration prerequisites.

Procedure 1-5 To verify prerequisites

1. (Conditional) Make sure that an NFS share is available.

   An NFS share is needed only if you plan to move the SGI MPT installation to a nondefault location on two or more partitions of an SGI UV computer.

2. As the root user, log into one of the partitions on the partitioned SGI UV computer.

3. (Conditional) Reboot the partition.

   Perform this step if the SGI UV computer was not rebooted after SGI Performance Suite was installed.

   If you do not know whether the computer has been rebooted, reboot at this time.
4. Verify that you have one of the following operating system software packages installed and configured:
   • Red Hat Enterprise Linux (RHEL) 6 or 7
   • SLES 11 or 12

   You can type the following command to verify your operating system version:

   `# cat /etc/issue`

5. Type a series of `cat(1)` commands to verify that the following products from the SGI Performance Suite are installed:

   • SGI Accelerate
   • SGI MPI

   For example:

   `# cat /etc/sgi-accelerate-release`

   SGI Performance Suite 1.12, Build xxxxxxx.sles11sp4-xxxxxxxxxx

   `# cat /etc/sgi-mpi-release`

   SGI MPI 1.12, Build xxxxxxx.sles11sp4-xxxxxxxxxx

6. Proceed to the following:

   "Configuring the OpenFabrics Enterprise Distribution (OFED) Software" on page 9

**Configuring the OpenFabrics Enterprise Distribution (OFED) Software**

All SGI UV computers are equipped with NUMAlink technology. Some SGI UV computers are also equipped with InfiniBand hardware, which uses OFED software. The procedure in this topic explains how to test for the presence of InfiniBand hardware and how to specify the number of queue pairs (QPs) for the OFED software.

If you are installing a kernel-based virtual machine (KVM), be aware that neither SGI, nor RHEL, nor SLES support InfiniBand hardware within a KVM.

The following procedure explains how to adjust the `log_num_qp` parameter.
1: Configuring the SGI Message Passing Toolkit (MPT)

Procedure 1-6 To specify the log_num_qp parameter

1. Type the following command to determine whether this partition is equipped with InfiniBand hardware:

   # lspci | grep Mellanox

   Note whether the command returns information similar to the following, which informs you of the presence of InfiniBand hardware:

   03:00.0 Network controller: Mellanox Technologies MT27500 Family [ConnectX-3]

   If the lspci command returns nothing, this partition is not connected to InfiniBand hardware. You do not need to perform the rest of this procedure. Proceed to:

   "Adjusting File Resource Limits" on page 11

2. Type one of the following commands to determine whether the OFED software is installed on this partition:

   • On SLES platforms, type the following:

     # zypper info -t pattern ofed

   • On RHEL platforms, type the following:

     # yum grouplist "Infiniband Support"

   The operating system packages include OFED by default.

3. Use a text editor to open file /etc/modprobe.d/libmlx4.conf.

4. Add a line similar to the following to file /etc/modprobe.d/libmlx4.conf:

   options mlx4_core log_num_qp=21

   The default maximum number of queue pairs is $2^{18}$ (262144).

   The log_num_qp parameter defines the log2 of the number of queue pairs (QPs). This step specifies the maximum number of queue pairs (QPs) for SHMEM applications. If the log_num_qp parameter is set to a number that is too low, the system generates the following message:

   MPT Warning: IB failed to create a QP

5. Save and close the file.
6. Proceed to the following:

"Adjusting File Resource Limits" on page 11

Adjusting File Resource Limits

The following procedure explains how to increase resource limits on the number of open files and how to enforce new security policies.

Procedure 1-7 To adjust file resource limits

1. Type the following command to retrieve the number of cores on this computer:

   ```
   # cat /proc/cpuinfo | grep processor | wc -l
   ```

   In the preceding line, the last character is a lowercase L, not the number 1.

   This `cat(1)` command returns the number of cores on the SGI UV computer system.

2. Use a text editor to open file `/etc/security/limits.conf`.

3. Add the following line to file `/etc/security/limits.conf`:

   ```
   * hard nofile limit
   ```

   For `limit`, specify an open file limit, for the number of MPI processes per host, based on the following guidelines:

<table>
<thead>
<tr>
<th>Processes/host</th>
<th>limit</th>
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<td>Up to 2048</td>
<td>8192 (default)</td>
</tr>
<tr>
<td>4096 or more</td>
<td>21000</td>
</tr>
</tbody>
</table>

   MPI jobs require a large number of file descriptors, and on larger systems, you might need to increase the system-wide limit on the number of open files. The default value for the file-limit resource is 8192. For example, the following line is suitable for 512 MPI processes per host:

   ```
   * hard nofile 3000
   ```
4. Add the following line to file /etc/security/limits.conf:
   *

   hard memlock unlimited

   The preceding line increases the resource limit for locked memory.

5. Save and close file /etc/security/limits.conf.

6. Use a text editor to open file /etc/pam.d/login, which is the Linux pluggable
   authentication module (PAM) configuration file.

7. Add the following line to file /etc/pam.d/login:

   session required /lib/security/pam_limits.so

8. Save and close the file.

9. (Conditional) Update other authentication configuration files as needed.

   Perform this step if your site allows other login methods, such as ssh, rlogin,
   and so on.

10. Proceed to the following:

    "Creating a Directory and Removing the Current Software" on page 12

---

**Creating a Directory and Removing the Current Software**

The following procedure explains how to create an NFS-mounted directory and
remove the SGI MPT software that currently resides on each partition.

**Procedure 1-8** To create a directory and remove the existing software

1. Familiarize yourself with the current SGI MPT working directory structure, and
   create a directory for the SGI MPT software you want to configure at this time.

   By default, the product installs into the /opt/sgi/mpt/mpt-2.14 directory.
   Make a plan for the nondefault structure at this time. In a partitioned
   environment, you install SGI MPT into a central NFS-mounted location.

   For example, use the mkdir(1) command to create the following alternate
directory:

   # mkdir -p /nfsmount/sgimpi/mpt-2.14
This documentation uses directory `/nfsmount/sgimpi/mpt-2.14` as an example nondefault working directory, configures SGI MPT 2.14 in that directory, and uses that example directory in the remaining steps of this configuration procedure.

2. Type the following command, and verify that all SGI MPT packages are in the default installation directory at this time:

```
# rpm -qa | grep sgi-mpt
```

Scan the `rpm` command output, and make sure that the following three SGI MPT packages appear:

- `sgi-mpt-shmem-2.14-sgi714r6.sles11sp4`
- `sgi-mpt-2.14-sgi714r6.sles11sp4`
- `sgi-mpt-fs-2.14-sgi714r6.sles11sp4`

3. Use a series of `rpm` commands to remove the SGI MPT packages from the default installation directory.

Your goal is to remove only the following packages:

- `sgi-mpt-shmem-release_number`
- `sgi-mpt-release_number`
- `sgi-mpt-fs-release_number`

The `rpm` command you need to use has the following format:

```
rpm -e --nodeps package_name
```

For `package_name`, specify the name of the package in the default directory at this time.

For example:

```
# rpm -e --nodeps sgi-mpt-shmem-2.14-sgi714r6.sles11sp4
# rpm -e --nodeps sgi-mpt-2.14-sgi714r6.sles11sp4
# rpm -e --nodeps sgi-mpt-fs-2.14-sgi714r6.sles11sp4
```

4. Proceed to one of the following:

- "(Optional) Configuring the MUNGE Security Software" on page 14
- "Updating Other Partitions or Continuing the Configuration" on page 15
(Optional) Configuring the MUNGE Security Software

Perform the procedure in this topic if you want to configure the MUNGE security software.

Array Services provides authentication services, but MUNGE provides additional authentication and security for Array Services operations. If you want to configure MUNGE, you need to configure it on each partition that you want to include in the array.

Procedure 1-9 To configure MUNGE

1. Verify that the partition is connected to a good time source, such as an NTP server.
   
   MUNGE depends on time synchronization across all nodes in the array.

2. Type one of the following commands to start the MUNGE installation, and respond to the installation prompts:
   
   • On Red Hat Enterprise Linux platforms, type the following command:
     
     # yum install munge
   
   • On SUSE Linux Enterprise Server platforms, type the following command:
     
     # zypper install munge

   For more information about how to install MUNGE, see the SGI MPI release notes.

3. Type the following command to restart MUNGE:
   
   # service munge restart

4. Type the following command to verify the existence of a MUNGE key on the partition:
   
   # md5sum /etc/munge/munge.key

5. (Conditional) Copy one partition’s MUNGE key to all of the partitions.

   Perform this step if this is the last partition that you need to configure.

   Immediately after you install MUNGE, each partition should have a unique key. When you run the partitions as an array, however, each partition needs to have the same key. After you have MUNGE installed on all the partitions that you want to include in the array, select one partition, and copy that partition’s MUNGE key to file /etc/munge/munge.key on each of the other partitions.
6. Proceed to the following:

"Updating Other Partitions or Continuing the Configuration" on page 15

Updating Other Partitions or Continuing the Configuration

At this point, at least one of the partitions on your SGI UV computer is configured correctly for SGI MPT.

The following procedure explains how to proceed.

Procedure 1-10 To assess progress

1. (Conditional) Configure additional partitions.

   Make sure that you completed all the preceding procedures on all of the partitions that you want to include in the array before you continue with the procedures that follow.

   If you want to include additional partitions in the array, repeat the following procedures on the additional partitions:
   • "Verifying Prerequisites" on page 8
   • "Configuring the OpenFabrics Enterprise Distribution (OFED) Software" on page 9
   • "Adjusting File Resource Limits" on page 11
   • "Creating a Directory and Removing the Current Software" on page 12
   • "(Optional) Configuring the MUNGE Security Software" on page 14

2. Proceed to the following:

   "Configuring Array Services" on page 15

Configuring Array Services

SGI MPI depends on Array Services for several capabilities. During the configuration, your goal is to specify the partitions that you want to include in the array and to distribute the configuration files to each partition.

Table 1-1 on page 16 lists the documentation resources that contain additional configuration information.
Table 1-1 Array Configuration Resources

<table>
<thead>
<tr>
<th>Topic</th>
<th>Documentation Resource</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advanced configuration information</td>
<td>&quot;Additional Array Configuration Information&quot; on page 97</td>
</tr>
<tr>
<td>Array Services overview</td>
<td>array_services(5)</td>
</tr>
<tr>
<td>Configuration file format</td>
<td>arrayd.conf(4) /usr/lib/array/arrayd.conf.template</td>
</tr>
<tr>
<td>Configuration file validator</td>
<td>ascheck(1)</td>
</tr>
<tr>
<td>Array Services configuration utility</td>
<td>arrayconfig(8)</td>
</tr>
</tbody>
</table>

The procedure in this topic uses the arrayconfig(8) command to specify SGI UV partitions for an array and to update the Array Services configuration files on each host.

Procedure 1-11 To configure Array Services for multiple partitions

1. (Optional) Synchronize and distribute secure shell (SSH) keys to each partition you want to include in the array.

   If you have SSH keys configured, you can complete work on one partition and log into the next without typing passwords each time. When you configure Array Services, it might be convenient for you if SSH is configured in each partition.

2. Plan the authentication method you want the Array Services software to use.

   Your choices are as follows:
   - munge. Specify munge if you configured the MUNGE software in the following procedure:
     "(Optional) Configuring the MUNGE Security Software" on page 14
   - none. Disables all authentication.
   - noreMOTE. Disallows requests from remote systems.
   - simple (default). Generates hostname/key pairs by using the OpenSSL rand command, 64-bit values (if available), or by using $RANDOM Bash facilities.
For more information about the authentication levels, see arrayd.auth(5).

3. Log in as root to the partition to which you expect users to log in when they want to run SGI MPI jobs.

Run the arrayconfig(1M) command from the partition to which you expect users to run their SGI MPI jobs.

4. Use the arrayconfig(1M) command to specify the partitions that you want to include in the array.

The arrayconfig command configures the /etc/array/arrayd.conf and /etc/array/arrayd.auth files on each partition.

Type the arrayconfig(1M) command in the following format:

/usr/sbin/arrayconfig -a arrayname -A method -D -m hostname1 hostname2 ...

For arrayname, type a name for the array. For example, sgicluster. The default is default.

For method, type one of the following authentication methods: munge, none, noremote, or simple (default). For information about the authentication methods, see the arrayd.auth(4) man page.

For each hostname, specify the hostnames of the partitions upon which you installed the SGI Message Passing Toolkit (MPT) software. That is, for hostname1, hostname2, and so on, specify the hostnames of the partitions that you want to include in the array.

5. (Optional) Reset the default user account or the default array port.

By default, the Array Services installation and configuration process sets the following defaults in the /usr/lib/array/arrayd.conf configuration file:

- A default user account of arraysvcs.

Array Services requires that a user account exist on all hosts in the array for the purpose of running certain Array Services commands. If you create a different account, make sure to update the arrayd.conf file and set the user account permissions correctly on all hosts.

- A default port number of 5434.
The `/etc/services` file contains a line that defines the `arrayd` service and port number as follows:

```bash
sgi-arrayd 5434/tcp # SGI Array Services daemon
```

You can set any value for the port number, but all systems mentioned in the `arrayd.conf` file must use the same value.

6. Proceed to the following:

"Enabling Cross-partition NUMAlink MPI Communication and Restarting Services" on page 18

**Note:** If you have trouble with the Array Services configuration, examine the Array Services manual configuration procedure in the following topic:

Appendix B, "Configuring Array Services Manually"

---

**Enabling Cross-partition NUMAlink MPI Communication and Restarting Services**

When you configure a large SGI UV system into two or more NUMAlink-connected partitions, the partitions act as separate, clustered hosts. The hardware supports efficient and flexible global memory access for cross-partition communication on such systems, but to enable this access, you need to load special kernel modules. If you do not enable cross-partition NUMAlink MPI communication at this time, users might receive the following message when they run an application:

```
MPT ERROR from do_cross_gets/xpmem_get, rc = -1, errno = 22
```

Depending on your operating system, perform one of the following procedures to ensure that the kernel modules load every time the system boots:

- "Enabling Cross-partition Communication and Restarting Services (RHEL)" on page 18
- "Enabling Cross-partition Communication and Restarting Services (SLES)" on page 19

**Enabling Cross-partition Communication and Restarting Services (RHEL)**

The following procedure explains how to load the kernel modules on one partition that hosts a RHEL operating system.
Procedure 1-12 To load the kernel modules at boot

1. As the root user, log into one of the partitions upon which you installed the SGI MPT software.

2. Type the following command:

```
# echo "modprobe xpc" >> /etc/sysconfig/modules/sgi-propack.modules
```

3. Save and close the file.

4. Type one of the following command sequences:

```
# reboot -f
```

Or

```
# modprobe xpc
# modprobe xpmem
# /etc/init.d/procset restart
# /etc/init.d/array restart
```

5. Repeat the preceding steps on the other partitions in the array.

6. Proceed to the following:

"Completing the Configuration" on page 20

Enabling Cross-partition Communication and Restarting Services (SLES)

The following procedure explains how to load the kernel modules on one partition that hosts a SLES operating system.

Procedure 1-13 To load the kernel modules at boot

1. As the root user, log into one of the partitions upon which you installed the SGI MPT software.

2. Use a text editor to open file /etc/sysconfig/kernel.

3. Within file /etc/sysconfig/kernel, search for the line that begins with

   `MODULES_LOADED_ON_BOOT`

4. Add `xpc` to the list of modules that are loaded at boot time.

5. Save and close the file.
6. Type one of the following command sequences:

```
# reboot -f
```

Or
```
# modprobe xpc
# modprobe xpmem
# /etc/init.d/procset restart
# /etc/init.d/array restart
```

7. Repeat the preceding steps on the other partitions in the array.

8. Proceed to the following:

"Completing the Configuration" on page 20

### Completing the Configuration

The following procedure explains how to complete the SGI MPT configuration.

**Procedure 1-14** To complete the SGI MPT configuration

1. Run a test MPI program to make sure that the new software is working as expected.

2. (Conditional) Inform your user community of the location of the new SGI MPT release on this computer.

Perform this step if you moved the SGI MPT software to a nondefault location.

In this procedure’s examples, the module files are located in the following directories:

- **On RHEL platforms:**
  ```
  /nfsmount/sgimpi/mpt-2.14/usr/share/Modules/modulefiles/mpt/2.14
  ```

- **On SLES platforms:**
  ```
  /nfsmount/sgimpi/mpt-2.14/usr/share/modules/modulefiles/mpt/2.14
  ```
About Running MPI Applications

This chapter provides procedures for building MPI applications. It provides examples of the use of the `mpirun(1)` command to launch MPI jobs. It also provides procedures for building and running SHMEM applications.

The process of running MPI applications consists of the following procedures:

- "Loading the MPI Software Module and Specifying the Library Path" on page 21
- "Compiling and Linking the MPI Program" on page 23
- "Launching the MPI Application" on page 25

Loading the MPI Software Module and Specifying the Library Path

You need to ensure that programs can find the SGI MPT library routines when the programs run.

The default locations for the include files, the `.so` files, the `.a` files, and the `mpirun` command are pulled in automatically. To ensure that the `mpt` software module is loaded, you can load site-specific library modules, or you can specify the library path on the command line before you run the program.
The following procedure explains how to specify the path to the MPI libmpi.so library.

**Procedure 2-1** To determine the library path

1. (Optional) Set the library path in the mpt module file.
   Complete this step if your site uses module files.
   Sample module files reside in the following locations:
   - /opt/sgi/mpt/mpt-mpt_rel/doc
   - /usr/share/modules/modulefiles/mpt/mpt_rel
   To load the SGI MPT module, type the following command:
   ```
   module load mpt
   ```

2. Determine the directory into which the SGI MPT software is installed.
   ```
   ldd a.out
   libmpi.so => /tmp/usr/lib/libmpi.so (0x40014000)
   libc.so.6 => /lib/libc.so.6 (0x402ac000)
   libdl.so.2 => /lib/libdl.so.2 (0x4039a000)
   /lib/ld-linux.so.2 => /lib/ld-linux.so.2 (0x40000000)
   ```
   Line 1 in the preceding output shows the library path correctly as /tmp/usr/lib/libmpi.so. If you do not specify the correct library path, the SGI MPT software searches incorrectly for the libraries in the default location of /usr/lib/libmpi.so.

3. Type the following command to set the library path:
   ```
   setenv LD_LIBRARY_PATH /library_path/usr/lib
   ```
   For library_path, type the path to the directory in which the SGI MPT software is installed.
   Example 1. The following command uses information from the previous step to set the library path to /tmp/usr/lib:
   ```
   setenv LD_LIBRARY_PATH /tmp/usr/lib
   ```
Example 2. The following command assumes that the libraries reside in /data/nfs/lib, which might be the case if you installed SGI MPT in an NFS-mounted file system:

```
% setenv LD_LIBRARY_PATH /data/nfs/lib
```

**Compiling and Linking the MPI Program**

You can use one of the MPI wrapper compiler commands to run your program, or you can call the compiler directly. The following topics explain these two alternatives:

- "Compiling With the Wrapper Compilers" on page 23
- "Compiling With the GNU or Intel Compilers" on page 24

**Compiling With the Wrapper Compilers**

The MPI wrapper compilers automatically incorporate the compiling and linking functions into the compiler command. If possible, use one of the following wrapper compiler commands to run your program:

- `mpif08 -I /install_path/usr/include file.f -L lib_path/usr/lib`
- `mpif90 -I /install_path/usr/include file.f -L lib_path/usr/lib`
- `mpicxx -I /install_path/usr/include file.c -L lib_path/usr/lib`
- `mpicc -I /install_path/usr/include file.c -L lib_path/usr/lib`

The variables in the preceding commands are as follows:

- For `install_path`, type the path to the directory in which the SGI MPT software is installed.
- For `file`, type the name of your C or Fortran program file name.
- For `lib_path`, type the path to the library files.

For example:

```
% mpicc -I /tmp/usr/include simple1_mpi.c -L /tmp/usr/lib
```
Compiling With the GNU or Intel Compilers

This topic explains how to run an MPI program if you need to call the GNU or Intel compilers directly. When the SGI MPT RPM is installed as default, the commands to build an MPI-based application using the .so files are as follows:

- To compile using GNU compilers, choose one of the following commands:

  `g++ -o myprog myprog.C -lmpi++ -lmpi`
  `gcc -o myprog myprog.c -lmpi`

- To compile programs with the Intel compilers, choose one of the following commands:

  `icc -o myprog myprog.c -lmpi`  # C - version 8
  `mpif08 simple1_mpi.f`  # Fortran 2008 wrapper compiler
  `mpif90 simple1_mpi.f`  # Fortran 90 wrapper compiler
  `ifort -o myprog myprog.f -lmpi`  # Fortran - version 8
  `mpicc -o myprog myprog.c`  # MPI C wrapper compiler
  `mpicxx -o myprog myprog.C`  # MPI C++ wrapper compiler

**Note:** Use the Intel compiler to compile Fortran 90 programs.

- To compile Fortran programs with the Intel compiler and enable compile-time checking of MPI subroutine calls, insert a `USE MPI` statement near the beginning of each subprogram to be checked. Also, use the following command:

  `ifort -I/usr/include -o myprog myprog.f -lmpi`  # version 8

**Note:** The preceding command assumes a default installation. If your site has more than one version of SGI MPT installed, or if your site installed MPT into a nondefault location, contact your system administrator to verify the location of the module files. For a nondefault installation location, replace `/usr/include` with the name of the relocated directory.

- The special case of using the Open64 compiler in combination with hybrid MPI/OpenMP applications requires separate compilation and link command lines. The Open64 version of the OpenMP library requires the use of the `~openmp`
option on the command line for compiling, but it interferes with proper linking of MPI libraries. Use the following sequence:

```
% opencc -o myprog.o -openmp -c myprog.c
% opencc -o myprog myprog.o -lopenmp -lmpi
```

### Launching the MPI Application

You can use either a workload manager or the `mpirun` command to launch an MPI application.

The following topics explain these alternatives:

- "Using a Workload Manager to Launch an MPI Application" on page 25
- "Using the `mpirun` Command to Launch an MPI Application" on page 27

### Using a Workload Manager to Launch an MPI Application

When an MPI job is run from a workload manager like PBS Professional, Torque, or Load Sharing Facility (LSF), it needs to start on the cluster nodes and CPUs that have been allocated to the job. For multi-node MPI jobs, the command that you use to start this type of job requires you to communicate the node and CPU selection information to the workload manager. SGI MPT includes one of these commands, `mpiexec_mpt(1)`, and the PBS Professional workload manager includes another such command, `mpiexec(1)`. The following topics describe how to start MPI jobs with specific workload managers:

- "PBS Professional" on page 25
- "Torque" on page 26
- "Simple Linux Utility for Resource Management (SLURM)" on page 27

### PBS Professional

You can run MPI applications from job scripts that you submit through workload managers such as the PBS Professional workload manager.

Process and thread pinning onto CPUs is especially important on cache-coherent non-uniform memory access (ccNUMA) systems such as the SGI UV system series. Process pinning is performed automatically if PBS Professional is set up to run each application in a set of dedicated cpusets. In these cases, PBS Professional sets the
PBS_CPUSET_DEDICATED environment variable to the value YES. This has the same
effect as setting MPI_DSM_DISTRIBUTE=ON. Process and thread pinning are also
performed in all cases if omplace(1) is used.

Example 1. To run an MPI application with 512 processes, include the following in
the directive file:

```bash
#PBS -l select=512:ncpus=1
mpiexec_mpt ./a.out
```

Example 2. To run an MPI application with 512 Processes and four OpenMP threads
per process, include the following in the directive file:

```bash
#PBS -l select=512:ncpus=4
mpiexec_mpt omplace -nt 4 ./a.out
```

Some third-party debuggers support the mpiexec_mpt(1) command. The
mpiexec_mpt(1) command includes a -tv option for use with TotalView and
includes a -ddt option for use with DDT. For more information, see Chapter 4,
"Debugging MPI Applications" on page 43.

PBS Professional includes an mpiexec(1) command that enables you to run SGI MPI
applications. PBS Professional’s command does not support the same set of extended
options that the SGI mpiexec_mpt(1) supports.

For more information about the PBS Professional workload manager, see the
following website:


**Torque**

When running Torque, SGI recommends that you use the following mpiexec_mpt(1)
command to launch SGI MPT MPI jobs.

```bash
mpiexec_mpt [ -n P ] ./a.out
```

The P argument is optional. By default, the program runs with the original number of
processes specified on the job initialization in Torque. To use P, specify is the total
number of MPI processes in the application. This syntax applies whether running on
a single host or a clustered system.

For more information, see the mpiexec_mpt(1) man page. The mpiexec_mpt
command has a -tv option for use by SGI MPT when running the TotalView
Debugger with a workload manager like Torque. For more information about using
the mpiexec_mpt command -tv option, see "Using the TotalView Debugger with MPI Programs" on page 43.

Simple Linux Utility for Resource Management (SLURM)

SGI MPI is adapted for use with the SLURM workload manager. If you want to use SGI MPI with SLURM, use the SLURM pmi2 MPI plug-in. SGI MPI 1.8 or later requires SLURM 2.6 or later.

For general information about SLURM, see the following website:
http://slurm.schedmd.com

For more information about how to use MPI with SLURM, see the following website:
http://slurm.schedmd.com/mpi_guide.html

Using the mpirun Command to Launch an MPI Application

The mpirun(1) command starts an MPI application. For a complete specification of the command line syntax, see the mpirun(1) man page.

The following topics explain how to use the mpirun command to launch a variety of applications:

• "Launching a Single Program on the Local Host" on page 27
• "Launching a Multiple Program, Multiple Data (MPMD) Application on the Local Host" on page 28
• "Launching a Distributed Application" on page 28
• "Launching an Application by Using MPI Spawn Functions" on page 29

Launching a Single Program on the Local Host

To run an application on the local host, enter the mpirun command with the -np argument. Your entry must include the number of processes to run and the name of the MPI executable file.

Example 1. The following command starts three instances of the mtest application, which is passed an argument list (arguments are optional):

% mpirun -np 3 mtest 1000 "arg2"
Launching a Multiple Program, Multiple Data (MPMD) Application on the Local Host

You are not required to use a different host in each entry that you specify on the \texttt{mpirun} command. You can start a job that has multiple executable files on the same host.

Example 1. The following command runs one copy of \texttt{prog1} and five copies of \texttt{prog2} on the local host, and both executable files use shared memory:

\% \texttt{mpirun -np 1 prog1 : -np 5 prog2}

Launching a Distributed Application

You can use the \texttt{mpirun} command to start a program that consists of any number of executable files and processes, and you can distribute the program to any number of hosts. A host is usually a single machine, but it can be any accessible computer running the Array Services software. For a list of the available nodes on systems running Array Services software, type the following command:

\% \texttt{ainfo machines}

You can list multiple entries on the \texttt{mpirun} command line. Each entry contains an MPI executable file and a combination of hosts and process counts for running it. This gives you the ability to start different executable files on the same or different hosts as part of the same MPI application.

The examples show various ways to start an application that consists of multiple MPI executable files on multiple hosts.

Example 1. The following command runs ten instances of the \texttt{a.out} file on \texttt{host_a}:

\% \texttt{mpirun host_a -np 10 a.out}

Example 2. The following command launches ten instances of \texttt{fred} on each of three hosts. \texttt{fred} has two input arguments.

\% \texttt{mpirun host_a, host_b, host_c -np 10 fred arg1 arg2}

Example 3. The following command launches ten instances of \texttt{fred}, with different numbers of instances on each processor:

\% \texttt{mpirun host_a -np 2, host_b -np 3, host_c -np 5 fred arg1 arg2}
Example 4. The following command launches an MPI application on different hosts with different numbers of processes and executable files:

```
% mpirun host_a 6 a.out : host_b -np 26 b.out
```

### Launching an Application by Using MPI Spawn Functions

To use the MPI process creation functions `MPI_Comm_spawn` or `MPI_Comm_spawn_multiple`, use the `-up` option on the `mpirun` command to specify the universe size.

Example 1. The following command starts three instances of the `mtest` MPI application in a universe of size 10:

```
% mpirun -up 10 -np 3 mtest
```

By using one of the preceding MPI spawn functions, `mtest` can start up to seven more MPI processes.

When running MPI applications across multiple hosts that use the `MPI_Comm_spawn` or `MPI_Comm_spawn_multiple` functions, you might need to explicitly specify the partitions on which additional MPI processes can be launched. For more information, see the `mpirun(1)` man page.

### Compiling and Running SHMEM Applications

The following procedure explains how to compile and run SHMEM programs in general terms.

**Procedure 2-2 Compiling and Running SHMEM applications**

1. Use one of the SHMEM wrapper compiler commands to run your program or call the compiler directly.

   To use the wrapper compiler, use one of the following commands:

   - `oshcc`
   - `oshCC`
   - `oshfort`

   To compile the SHMEM program directly, use GNU compiler or Intel compiler commands.
2. Getting Started

- To compile SHMEM programs with a GNU compiler, choose one of the following commands:
  - g++ compute.C -lsma -lmpi
  - gcc compute.c -lsma -lmpi
- To compile SHMEM programs with an Intel compiler, choose one of the following commands:
  - icc compute.C -lsma -lmpi
  - icc compute.c -lsma -lmpi
  - ifort compute.f -lsma -lmpi

2. Use the mpirun command to launch the SHMEM application.

To request the desired number of processes to launch, set the -np option on the mpirun command. The NPES variable has no effect on SHMEM programs.

The SHMEM programming model supports both single-host SHMEM applications and SHMEM applications that span multiple partitions. To launch an SHMEM application on more than one partition, use multiple-host syntax on the mpirun syntax, as follows:

```
% mpirun hostA, hostB -np 16 ./shmem_app_name
```

For more information, see the intro_shmem(3) man page.

### Using Huge Pages

Huge pages optimize MPI application performance. The MPI_HUGEPAGE_HEAP_SPACE environment variable defines the minimum amount of heap space each MPI process can allocate using huge pages. If set to a positive number, libmpi verifies that enough hugetlbfs overcommit resources are available at program start-up to satisfy that amount on all MPI processes. The heap uses all available hugetlbfs space, even beyond the specified minimum amount. A value of 0 disables this check and disables the allocation of heap variables on huge pages. Values can be followed by K, M, G, or T to denote scaling by 1024, 1024², 1024³, or 1024⁴, respectively.

For information about the MPI_HUGEPAGE_HEAP_SPACE environment variable, see the mpi(1) man page.
The following steps explain how to configure system settings for huge pages.

**Procedure 2-3** To configure system settings for huge pages

1. Type the following command to make sure that the current SGI MPT software release module is installed:

   ```
   sys:~ # module load mpt
   ```

2. Log in as the root user, and type the following command to configure the system settings for huge pages:

   ```
   sys:~ # mpt_hugepage_config -u
   Updating system configuration
   System config file: /proc/sys/vm/nr_overcommit_hugepages
   Huge Pages Allowed: 28974 pages (56 GB) 90% of memory
   Huge Page Size: 2048 KB
   Huge TLB FS Directory: /etc/mpt/hugepage_mpt
   ```

3. Type the following command to retrieve the current system configuration:

   ```
   sys:~ # mpt_hugepage_config -v
   Reading current system configuration
   System config file: /proc/sys/vm/nr_overcommit_hugepages
   Huge Pages Allowed: 28974 pages (56 GB) 90% of memory
   Huge Page Size: 2048 KB
   Huge TLB FS Directory: /etc/mpt/hugepage_mpt (exists)
   ```

4. When running your SGI MPT program, make sure the `MPI_HUGEPAGE_HEAP_SPACE` environment variable is set to 1.

   This activates the new libmpi huge page heap. Memory allocated by calls to the `malloc` function are allocated on huge pages. This makes single-copy MPI sends much more efficient when using the SGI UV global reference unit (GRU) for MPI messaging.

5. Log in as the root user, and type the following command to clear the system configuration settings:

   ```
   sys:~ # mpt_hugepage_config -e
   Removing MPT huge page configuration
   ```
6. To verify that the SGI MPT huge page configuration has been cleared, type the following command to retrieve the system configuration again:

```
uv44-sys:~ # mpt_hugepage_config -v
Reading current system configuration

System config file: /proc/sys/vm/nr_overcommit_hugepages
Huge Pages Allowed: 0 pages (0 KB) 0% of memory
Huge Page Size: 2048 KB
Huge TLB FS Directory: /etc/mpt/hugepage_mpt (does not exist)
```

For more information about how to configure huge pages for MPI applications, see the `mpt_hugepage_config(1)` man page.

### Using SGI MPI in an SELinux Environment (RHEL Platforms Only)

SGI supports Security-Enhanced Linux (SELinux) for single-host runs on SGI computer systems that run the Red Hat Enterprise Linux (RHEL) operating system. The following guidelines pertain to using SELinux:

- SELinux for is configured. For configuration information, see the following manual:
  
  *SGI UV System Software Installation and Configuration Guide*

- The `MPI_USE_ARRAY` environment variable is set as follows:

  `MPI_USE_ARRAY=false`

  When set to `false`, Array Services is disabled. For more information about this environment variable, see the `MPI(1)` man page.

For more information about how to run SGI MPI with security software, contact SGI technical support.
Chapter 3

Programming With SGI MPI

This chapter includes the following topics:
• "About Programming With SGI MPI" on page 33
• "Job Termination and Error Handling" on page 33
• "Signals" on page 35
• "Buffering" on page 35
• "Multithreaded Programming" on page 36
• "Interoperability with the SHMEM programming model" on page 37
• "Miscellaneous SGI MPI Features" on page 37
• "Programming Optimizations" on page 38
• "Additional Programming Model Considerations" on page 41

About Programming With SGI MPI

Portability is one of the main advantages MPI has over vendor-specific message passing software. Nonetheless, the MPI Standard offers sufficient flexibility for general variations in vendor implementations. In addition, there are often vendor-specific programming recommendations for optimal use of the MPI library. This chapter’s topics explain how to develop or port MPI applications to SGI systems.

Job Termination and Error Handling

This section describes the behavior of the SGI MPI implementation upon normal job termination. Error handling and characteristics of abnormal job termination are also described.

This section includes the following topics:
• "MPI_Abort" on page 34
• "Error Handling" on page 34
• "MPI_Finalize and Connect Processes" on page 34

**MPI_Abort**

In the SGI MPI implementation, a call to `MPI_Abort` has the following effect:

- The MPI job terminates, regardless of the communicator argument used.
- The error code value is returned as the exit status of the `mpirun` command.
- A stack traceback is displayed that shows where the program called `MPI_Abort`.

**Error Handling**

The MPI Standard describes MPI error handling. Although almost all MPI functions return an error status, an error handler is invoked before returning from the function. If the function has an associated communicator, the error handler associated with that communicator is invoked. Otherwise, the error handler associated with `MPI_COMM_WORLD` is invoked.

The SGI MPI implementation provides the following predefined error handlers:

- `MPI_ERRORS_ARE_FATAL`. When called, causes the program to abort on all executing processes. This has the same effect as if `MPI_Abort` were called by the process that invoked the handler.
- `MPI_ERRORS_RETURN`. This handler has no effect.

By default, the `MPI_ERRORS_ARE_FATAL` error handler is associated with `MPI_COMM_WORLD` and any communicators derived from it. Hence, to handle the error statuses returned from MPI calls, it is necessary to associate either the `MPI_ERRORS_RETURN` handler or another user-defined handler with `MPI_COMM_WORLD` near the beginning of the application.

**MPI_Finalize and Connect Processes**

In the SGI implementation of MPI, all pending communications involving an MPI process must be complete before the process calls `MPI_Finalize`. If there are any pending `send` or `recv` requests that are unmatched or not completed, the application hangs in `MPI_Finalize`. For more details, see the MPI Standard.
If the application uses the MPI remote memory access (RMA) spawn functionality described in the MPI RMA standard, there are additional considerations. In the SGI implementation, all MPI processes are connected. The MPI RMA standard defines what is meant by connected processes. When the MPI RMA spawn functionality is used, `MPI_Finalize` is collective over all connected processes. Thus all MPI processes, both launched on the command line, or subsequently spawned, synchronize in `MPI_Finalize`.

**Signals**

In the SGI implementation, MPI processes are UNIX processes. As such, the general rule regarding signal handling applies as it would to ordinary UNIX processes.

In addition, the `SIGURG` and `SIGUSR1` signals can be propagated from the `mpirun` process to the other processes in the MPI job, whether they belong to the same process group on a single host or are running across multiple hosts in a cluster. To use this feature, the MPI program must have a signal handler that catches `SIGURG` or `SIGUSR1`. When the `SIGURG` or `SIGUSR1` signals are sent to the `mpirun` process ID, the `mpirun` process catches the signal and propagates it to all MPI processes.

**Buffering**

Most MPI implementations use buffering for overall performance reasons, and some programs depend on it. However, you should not assume that there is any message buffering between processes because the MPI Standard does not mandate a buffering strategy. Table 3-1 on page 36 illustrates a simple sequence of MPI operations that cannot work unless messages are buffered. If sent messages are not buffered, each process hangs in the initial call, waiting for an `MPI_Recv` call to take the message.

Because most MPI implementations buffer messages to some degree, a program like this does not usually hang. The `MPI_Send` calls return after putting the messages into buffer space, and the `MPI_Recv` calls get the messages. Nevertheless, program logic like this is not valid according to the MPI Standard. Programs that require this sequence of MPI calls should employ one of the buffer MPI send calls, `MPI_Bsend` or `MPI_Ibsend`.
Table 3-1 Outline of Improper Dependence on Buffering

<table>
<thead>
<tr>
<th>Process 1</th>
<th>Process 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_Send(2,....)</td>
<td>MPI_Send(1,....)</td>
</tr>
<tr>
<td>MPI_Recv(2,....)</td>
<td>MPI_Recv(1,....)</td>
</tr>
</tbody>
</table>

By default, the SGI implementation of MPI uses buffering under most circumstances. Short messages (64 or fewer bytes) are always buffered. Longer messages are also buffered, although under certain circumstances, buffering can be avoided. For performance reasons, it is sometimes desirable to avoid buffering. For further information on unbuffered message delivery, see "Programming Optimizations" on page 38.

Multithreaded Programming

SGI MPI supports a hybrid programming model, in which MPI handles one level of parallelism in an application and POSIX threads or OpenMP processes are used to handle another level. When mixing OpenMP with MPI, for performance reasons, it is better to consider invoking MPI functions only outside parallel regions or only from within master regions. When used in this manner, it is not necessary to initialize MPI for thread safety. You can use MPI_Init to initialize MPI. However, to safely invoke MPI functions from any OpenMP process or when using Posix threads, MPI must be initialized with MPI_Init_thread.

When using MPI_Thread_init() with the threading level MPI_THREAD_MULTIPLE, link your program as follows:

- If you use the compiler wrappers for MPI or SHMEM, use the -mt option on the command line.
- If you want to call the compilers directly, use the -lmpi_mt parameter instead of the -lmpi parameter on the compiler command line.

For more information about compiling and linking MPI programs, see the mpi(1) man page.
Interoperability with the SHMEM programming model

You can mix SHMEM and MPI message passing in the same program. The application must be linked with both the SHMEM and MPI libraries.

Start with an MPI program that calls MPI_Init (or MPI_Init_thread()) and MPI_Finalize. Next, add SHMEM calls, and be aware that the PE numbers are equal to the MPI rank numbers in MPI_COMM_WORLD.

If your program uses both SHMEM and MPI, make sure your program includes calls to the shmem_init() and shmem_finalize() library routines. This practice is similar to how you include calls to MPI_Init() (or MPI_Init_thread()) and MPI_Finalize.

When running the application across a cluster using SHMEM and SHMEM functions, some processes might not be able to communicate with other processes. You can use the shmem_pe_accessible and shmem_addr_accessible functions to determine whether a SHMEM call can be used to access data residing in another process. Because the SHMEM model functions only with respect to MPI_COMM_WORLD, these functions cannot be used to exchange data between MPI processes that are connected via MPI intercommunicators returned from MPI spawn-related functions.

For more information about the SHMEM programming model, see the intro_shmem(3) man page.

Miscellaneous SGI MPI Features

The following other characteristics of the SGI MPI implementation might interest you:

• stdin/stdout/stderr.

In this implementation, stdin is enabled for only the process that is rank 0 in the first MPI_COMM_WORLD. Such processes do not need to be located on the same host as mpirun. The stdout and stderr results are enabled for all MPI processes in the job, whether started by mpirun or started by one of the MPI spawn functions.

• MPI_Get_processor_name

The MPI_Get_processor_name function returns the Internet host name of the computer upon which the MPI process that started this subroutine is running.
Programming Optimizations

You might need to modify your MPI application to use the SGI MPI optimization features.

The following topics describe how to use the optimized features of SGI’s MPI implementation:

• "Using MPI Point-to-Point Communication Routines" on page 38
• "Using MPI Collective Communication Routines" on page 39
• "Using MPI_Pack/MPI_Unpack" on page 39
• "Avoiding Derived Data Types" on page 40
• "Avoiding Wild Cards" on page 40
• "Avoiding Message Buffering — Single Copy Methods" on page 40
• "Managing Memory Placement” on page 41

Using MPI Point-to-Point Communication Routines

MPI provides a number of different routines for point-to-point communication. The most efficient ones in terms of latency and bandwidth are the blocking and nonblocking send/receive functions, which are as follows:

• MPI_Send
• MPI_Isend
• MPI_Recv
• MPI_Irecv

Unless required for application semantics, avoid the synchronous send calls, which are as follows:

• MPI_Ssend
• MPI_Issend

Also avoid the buffered send calls, which double the amount of memory copying on the sender side. These calls are as follows:
• MPI_Bsend
• MPI_Ibsend

This implementation treats the ready-send routines, MPI_Rsend and MPI_Irsend, as standard MPI_Send and MPI_Isend routines. Persistent requests do not offer any performance advantage over standard requests in this implementation.

Using MPI Collective Communication Routines

The MPI collective calls are frequently layered on top of the point-to-point primitive calls. For small process counts, this can be reasonably effective. However, for higher process counts of 32 processes or more, or for clusters, this approach can be less efficient. For this reason, a number of the MPI library collective operations have been optimized to use more complex algorithms.

SGI’s MPI collectives have been optimized for use with clusters. In these cases, steps are taken to reduce the number of messages using the relatively slower interconnect between hosts.

Some of the collective operations have been optimized for use with shared memory. On SGI UV systems, barriers and reductions have been optimized to use the SGI GRU hardware accelerator. The MPI_Alltoall routines also use special techniques to avoid message buffering when using shared memory. For more information, see "Avoiding Message Buffering — Single Copy Methods" on page 40.

Note: Collectives are optimized across partitions by using the XPMEM driver which is explained in Chapter 7, "Run-time Tuning". The collectives (except MPI_BARRIER) try to use single-copy by default for large transfers unless MPI_DEFAULT_SINGLE_COPY_OFF is specified.

Using MPI_Pack/MPI_Unpack

While MPI_Pack and MPI_Unpack are useful for porting parallel virtual machine (PVM) codes to MPI, they essentially double the amount of data to be copied by both the sender and receiver. Generally, either restructure your data or use derived data types to avoid using these functions. Note, however, that use of derived data types can lead to decreased performance in certain cases.
Avoiding Derived Data Types

Avoid derived data types when possible. In the SGI implementation, using derived data types does not generally lead to performance gains. Using derived data types might disable certain types of optimizations, for example, unbuffered or single copy data transfer.

Avoiding Wild Cards

The use of wild cards (MPI_ANY_SOURCE, MPI_ANY_TAG) involves searching multiple queues for messages. While this is not significant for small process counts, for large process counts, the cost increases quickly.

Avoiding Message Buffering — Single Copy Methods

One of the most significant optimizations for bandwidth-sensitive applications in the MPI library is single-copy optimization, which avoids using shared memory buffers. However, as discussed in "Buffering" on page 35, some incorrectly coded applications might hang because of buffering assumptions. For this reason, this optimization is not enabled by default for MPI_Send, but you can use the MPI_BUFFER_MAX environment variable to enable this optimization at runtime. The following guidelines show how to increase the opportunity for use of the unbuffered pathway:

- The MPI data type on the send side must be a contiguous type.
- The sender and receiver MPI processes must reside on the same host. In the case of a partitioned system, the processes can reside on any of the partitions.
- The sender data must be globally accessible by the receiver. The SGI MPI implementation allows data allocated from the static region (common blocks), the private heap, and the stack region to be globally accessible. In addition, memory allocated via the MPI_Alloc_mem function or the SHMEM symmetric heap accessed via the shmalloc or shmalloc functions is globally accessible.

Certain run-time environment variables must be set to enable the unbuffered, single-copy method. For information about how to set the run-time environment, see "Avoiding Message Buffering – Enabling Single Copy" on page 56.
Managing Memory Placement

SGI UV series systems have a ccNUMA memory architecture. For single-process and small multiprocess applications, this architecture behaves similarly to flat memory architectures. For more highly parallel applications, memory placement becomes important. MPI takes placement into consideration when it lays out shared memory data structures and the individual MPI processes’ address spaces. Generally, you should not try to manage memory placement explicitly. To control the placement of the application at run time, however, see Chapter 7, "Run-time Tuning" on page 53.

Additional Programming Model Considerations

A number of additional programming options might be worth consideration when developing MPI applications for SGI systems. For example, using the SHMEM programming model can improve performance in the latency-sensitive sections of an application. Usually, this requires replacing MPI send/recv calls with shmem_put/shmem_get and shmem_barrier calls. The SHMEM programming model can deliver significantly lower latencies for short messages than traditional MPI calls. As an alternative to shmem_get/shmem_put calls, you might consider the MPI remote memory accesss (RMA) MPI_Put/ MPI_Get functions. These provide almost the same performance as the SHMEM calls, while providing a greater degree of portability.

Alternately, you might consider exploiting the shared memory architecture of SGI systems by handling one or more levels of parallelism with OpenMP, with the coarser grained levels of parallelism being handled by MPI. Also, there are special ccNUMA placement considerations to be aware of when running hybrid MPI/OpenMP applications. For further information, see Chapter 7, "Run-time Tuning" on page 53.
Debugging MPI Applications

This chapter includes the following topics:

- "MPI Routine Argument Checking" on page 43
- "Using the TotalView Debugger with MPI Programs" on page 43
- "Using idb and gdb with MPI Programs" on page 44
- "Using the DDT Debugger with MPI Programs" on page 44
- "Using Valgrind With MPI Programs" on page 45

MPI Routine Argument Checking

Debugging MPI applications can be more challenging than debugging sequential applications. By default, the SGI MPI implementation does not check the arguments to some performance-critical MPI routines, such as most of the point-to-point and collective communication routines. You can force MPI to always check the input arguments to MPI functions by setting the `MPI_CHECK_ARGS` environment variable. However, setting this variable might result in some degradation in application performance, so it is not recommended that it be set except when debugging.

Using the TotalView Debugger with MPI Programs

The SGI Message Passing Toolkit (MPT) `mpiexec_mpt(1)` command has a `-tv` option for use by SGI MPT with the TotalView Debugger. Note that the PBS Professional `mpiexec(1)` command does not support the `-tv` option. TotalView does not operate with MPI processes started via the `MPI_Comm_spawn` or `MPI_Comm_spawn_multiple` functions.

Example 1. To run an SGI MPT MPI job with TotalView without a workload manager, type the following:

```
% totalview mpirun -a -np 4 a.out
```
Example 2. To run an SGI MPT MPI job with the TotalView Debugger with a workload manager, such as PBS Professional or Torque, type the following:

```
% mpiexec_mpt -tv -np 4 a.out
```

### Using idb and gdb with MPI Programs

Because the idb and gdb debuggers are designed for sequential, non-parallel applications, they are generally not well suited for use in MPI program debugging and development. However, the use of the `MPI_SLAVE_DEBUG_ATTACH` environment variable makes these debuggers more usable.

If you set the `MPI_SLAVE_DEBUG_ATTACH` environment variable to a global rank number, the MPI process sleeps briefly in startup while you use idb or gdb to attach to the process. A message is printed to the screen, telling you how to use idb or gdb to attach to the process.

Similarly, if you want to debug the MPI daemon, setting `MPI_DAEMON_DEBUG_ATTACH` sleeps the daemon briefly while you attach to it.

### Using the DDT Debugger with MPI Programs

Allinea Software’s DDT product is a parallel debugger that supports SGI MPT. You can run DDT in either interactive (online) or batch (offline) mode. In batch mode, DDT can create a text or HTML report that tracks variable values and shows the location of any errors. DDT records the data for a program’s variables across all processes, and DDT logs values in the HTML output files as sparkline charts.

For information about how to configure Allinea for use with MPI on SGI systems, use the instructions in the Allinea user guide that is posted to the following website:


Example 1. The following command starts DDT in interactive (online) mode:

```
# ddt -np 4 a.out
```

Example 2. The following command generates a debugging report in HTML format:

```
# ddt -offline my-log.html -np 4 a.out
```
Example 3. Assume that you want to trace variables \( x \), \( y \), and \( my\_arr(x,y) \) in parallel across all processes. The following command directs DDT to record the values of \( x \), \( y \), and \( my\_arr(x,y) \) each time it encounters line 147:

```
# ddt -offline my-log.html -trace-at "my-file.f:147,x,y,my_arr(x,y)" -np 4 a.out
```

Example 4. You can specify batch (offline) DDT commands from within a queue submission script. Instead of specifying `mpiexec_mpt -np 4 a.out`, specify the following:

```
# ddt -noqueue -offline my-log.html -trace-at "my-file.f:147,x,y,my_arr(x,y)" -np 4 a.out
```

### Using Valgrind With MPI Programs

Valgrind is a tool that can profile your program and can automatically detect memory management and threading bugs.

Valgrind is not compatible with the memory mapping functionality in SGI MPT. When SGI MPT detects that Valgrind is in use, SGI MPT automatically enables the `MPI_MEMMAP_OFF` environment variable, which disables SGI MPT’s own memory mapping.
This chapter includes the following topics:

- "About PerfBoost" on page 47
- "Using PerfBoost" on page 47
- "MPI Supported Functions" on page 48

**About PerfBoost**

SGI PerfBoost uses a wrapper library to run applications compiled against other MPI implementations under the SGI Message Passing Toolkit (MPT) product on SGI platforms. This chapter describes how to use PerfBoost software.

**Note:** PerfBoost does not support the MPI C++ API.

**Using PerfBoost**

The following procedure explains how to use PerfBoost with an SGI MPI program.

**Procedure 5-1** To use PerfBoost

1. Load the `perfboost` environment module.

   The module include the `PERFBOOST_VERBOSE` environment variable.

   If you set the `PERFBOOST_VERBOSE` environment variable, it enables a message when PerfBoost activates and also when the MPI application is completed through the `MPI_Finalize()` function. This message indicates that the PerfBoost library is active and also indicates when the MPI application completes through the `libperfboost` wrapper library.

   The MPI environment variables that are documented in the `MPI(1)` man page are available to PerfBoost. MPI environment variables that are not used by SGI MPT are currently not supported.
5: Using PerfBoost

Note: Some applications redirect stderr. In this case, the verbose messages might not appear in the application output.

2. Type a command that inserts the `perfboost` command in front of the executable name along with the choice of MPI implementation to emulate.

   In other words, run the executable file with the SGI MPT `mpiexec_mpt(1)` or the `mpirun(1)` command.

   The following are MPI implementations and corresponding command line options:

<table>
<thead>
<tr>
<th>Implementation</th>
<th>Command Line Option</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platform MPI 7.1+</td>
<td>-pmpi</td>
</tr>
<tr>
<td>HP-MPI</td>
<td>-pmpi</td>
</tr>
<tr>
<td>Intel MPI</td>
<td>-impi</td>
</tr>
<tr>
<td>OpenMPI</td>
<td>-ompi</td>
</tr>
<tr>
<td>MPICH1</td>
<td>-mpich</td>
</tr>
<tr>
<td>MPICH2, version 2 and later</td>
<td>-impi</td>
</tr>
<tr>
<td>MVAPICH2, version 2 and later</td>
<td>-impi</td>
</tr>
</tbody>
</table>

   The following are some examples that use `perfboost`:

   ```
   % module load mpt
   % module load perfboost
   % mpirun -np 32 perfboost -impi a.out arg1
   % mpiexec_mpt perfboost -pmpi b.out arg1
   % mpirun host1 32, host2 64 perfboost -impi c.out arg1 arg2
   ```

**MPI Supported Functions**

SGI PerfBoost supports the commonly used elements of the C and Fortran MPI APIs. If a function is not supported, the job aborts and issues an error message. The message shows the name of the missing function. You can contact the SGI Customer
Support Center at the following website to schedule a missing function to be added to PerfBoost:

https://support.sgi.com/caselist
This chapter includes the following topics:

- "About Berkeley Lab Checkpoint/Restart’ on page 51
- "BLCR Installation’ on page 51
- "Using BLCR with SGI MPT’ on page 52

About Berkeley Lab Checkpoint/Restart

The SGI Message Passing Toolkit (MPT) supports the Berkeley Lab Checkpoint/Restart (BLCR) checkpoint/restart. This checkpoint/restart implementation allows applications to periodically save a copy of their state. Applications can resume from that point if the application crashes or if the job is aborted to free resources for higher-priority jobs.

The following are the implementation’s limitations:

- BLCR does not checkpoint the state of any data files that the application might be using.
- When using checkpoint/restart, MPI does not support certain features, including spawning and one-sided MPI.
- InfiniBand XRC queue pairs are not supported.
- Checkpoint files are often very large and require significant disk bandwidth to create in a timely manner.

For more information on BLCR, see http://crd.lbl.gov/departments/computer-science/CLaSS/research/BLCR/.

BLCR Installation

To use checkpoint/restart with SGI MPT, BLCR must first be installed.

Procedure 6-1 To install BLCR

1. Log in as the root user.
2. Install the blcr-, blcr-libs-, and blcr-kmp- RPMs.

BLCR uses a kernel module that must be built against the specific kernel that the operating system is running. If the kernel module fails to load, you need to rebuild and reinstall. Install the blcr- source RPM. In the blcr.spec file, set the kernel variable to the name of the current kernel, then rebuild and install the new set of RPMs.

3. Type the following command to enable BLCR:

   # chkconfig blcr on

Using BLCR with SGI MPT

To enable checkpoint/restart within SGI MPT, you need to pass the –cpr option to mpirun or mpiexec_mpt. For example:

   % mpirun –cpr hostA, hostB –np 8 ./a.out

To checkpoint a job, run the mpt_checkpoint command on the same host upon which mpirun is running. Make sure to pass the mpt_checkpoint command the PID of mpirun and the name with which you want to prefix all the checkpoint files. For example:

   % mpt_checkpoint –p 12345 –f my_checkpoint

The preceding example command creates a my_checkpoint.cps metadata file and a number of my_checkpoint.*.cpd files.

To restart the job, pass the name of the .cps file to mpirun. For example:

   % mpirun –cpr hostC, hostD –np 8 mpt_restart my_checkpoint.cps

You can restart the job on a different set of hosts, but the number of hosts must be the same. In addition, each host must have the same number of ranks as the corresponding host in the original run of the job.
Run-time Tuning

This chapter includes the following topics:

- "About Run-time Tuning" on page 53
- "Reducing Run-time Variability" on page 54
- "Tuning MPI Buffer Resources" on page 55
- "Avoiding Message Buffering – Enabling Single Copy" on page 56
- "Memory Placement and Policies" on page 57
- "Tuning MPI/OpenMP Hybrid Codes" on page 59
- "Tuning Running Applications Across Multiple Hosts" on page 61
- "Tuning for Running Applications over the InfiniBand Interconnect" on page 63
- "MPI on SGI UV Systems" on page 65
- "Suspending MPI Jobs" on page 68

About Run-time Tuning

This chapter describes the ways in which a user can tune the run-time environment to improve the performance of an MPI message passing application on SGI computers. None of these ways involve application code changes.

The run-time tuning topics are as follows:

- "Reducing Run-time Variability" on page 54
- "Tuning MPI Buffer Resources" on page 55
- "Avoiding Message Buffering – Enabling Single Copy" on page 56
- "Memory Placement and Policies" on page 57
- "Tuning MPI/OpenMP Hybrid Codes" on page 59
- "Tuning Running Applications Across Multiple Hosts" on page 61
Reducing Run-time Variability

One of the most common problems with optimizing message passing codes on large, shared-memory computers is to achieve reproducible timings from run to run. To reduce run-time variability, you can take the following precautions:

- Do not oversubscribe the system. In other words, do not request more CPUs than are available, and do not request more memory than is available. Oversubscribing causes the system to wait unnecessarily for resources to become available, leads to variations in the results, and leads to less than optimal performance.

- Avoid interference from other system activity. The Linux kernel uses more memory on node 0 than on other nodes. Node 0 is also known as the kernel node. If your application uses almost all of the available memory per processor, the memory for processes assigned to the kernel node can unintentionally spill over to nonlocal memory. By keeping user applications off of the kernel node, you can avoid this effect.

By restricting system daemons to run on the kernel node, you can also deliver an additional percentage of each application CPU to the user program.

- Avoid interference with other applications. If necessary, use cpusets to address this problem. The cpuset software enables you to partition a large, distributed memory host in a fashion that minimizes interactions between jobs running concurrently on the system. For more information about cpusets, see the following:

  SGI Cpuset Software Guide

- On a quiet, dedicated system, you can use the dplace(1) tool or the MPI_DSM_CPULIST environment variable to improve run-time performance repeatability. These approaches are not suited to shared, nondedicated systems.

- Use a workload manager such as Platform LSF from IBM or PBS Professional from Altair Engineering, Inc. These workload managers use cpusets to avoid oversubscribing the system and to avoid possible interference between applications.
Tuning MPI Buffer Resources

By default, the SGI MPI implementation buffers messages that are longer than 64 bytes. The system buffers these longer messages in a series of 16 KB buffers. Messages that exceed 64 bytes are handled as follows:

- If the message is 128 K in length or shorter, the sender MPI process buffers the entire message.

  In this case, the sender MPI process delivers a message header, also called a control message, to a mailbox. When an MPI call is made, the MPI receiver polls the mailbox. If the receiver finds a matching receive request for the sender’s control message, the receiver copies the data out of the buffers into the application buffer indicated in the receive request. The receiver then sends a message header back to the sender process, indicating that the buffers are available for reuse.

- If the message is longer than 128 K, the software breaks the message into chunks that are 128 K in length.

  The smaller chunks allow the sender and receiver to overlap the copying of data in a pipelined fashion. Because there are a finite number of buffers, this can constrain overall application performance for certain communication patterns. You can use the MPI_BUFS_PER_PROC shell variable to adjust the number of buffers available for each process, and you can use the MPI statistics counters to determine if the demand for buffering is high.

  Generally, you can avoid excessive numbers of retries for buffers if you increase the number of buffers. However, when you increase the number of buffers, you consume more memory, and you might increase the probability for cache pollution. Cache pollution is the excessive filling of the cache with message buffers. Cache pollution can degrade performance during the compute phase of a message passing application.

For information about statistics counters, see "MPI Internal Statistics" on page 78.

For information about buffering considerations when running an MPI job across multiple hosts, see "Tuning Running Applications Across Multiple Hosts" on page 61.

For information about the programming implications of message buffering, see "Buffering" on page 35.
Avoiding Message Buffering – Enabling Single Copy

It is possible to avoid the need to buffer messages for message transfers between MPI processes within the same host or message transfers that use devices that support remote direct memory access (RDMA), such as InfiniBand.

The following topics provide more information about buffering:
- "Buffering and MPI_Send" on page 56
- "Using the XPMEM Driver for Single Copy Optimization" on page 56

Buffering and MPI_Send

Many MPI applications are written to assume infinite buffering, so message buffering is not enabled by default for MPI_Send. For MPI_Isend, MPI_Sendrecv, and most collectives, this optimization is enabled by default for large message sizes. To disable this default, single-copy feature used for the collectives, use the MPI_DEFAULT_SINGLE_COPY_OFF environment variable.

Using the XPMEM Driver for Single Copy Optimization

MPI uses the XPMEM driver to support single-copy message transfers between two processes within the same host or across partitions.

Enabling single-copy transfers can increase performance because this technique improves MPI's bandwidth. On the other hand, single-copy transfers can introduce additional synchronization points, which can reduce application performance.

The MPI_BUFFER_MAX environment variable specifies the threshold for message lengths. Its value should be set to the message length, in bytes, beyond which you want MPI to use the single-copy method. In general, a value of 2000 or higher is beneficial for many applications.

During job startup, MPI uses the XPMEM driver, via the xpmem kernel module, to map memory from one MPI process to another. The mapped areas include the static (BSS) region, the private heap, the stack region, and (optionally) the symmetric heap region of each process.

Memory mapping allows each process to directly access memory from the address space of another process. This technique allows MPI to support single-copy transfers for contiguous data types from any of these mapped regions. For these transfers,
whether between processes residing on the same host or across partitions, MPI uses the \texttt{bcopy} process to copy the data. The \texttt{bcopy} process also transfers data between two different executable files on the same host or between two different executable files across partitions. For data residing outside of a mapped region (a /dev/zero region, for example), MPI uses a buffering technique to transfer the data.

Memory mapping is enabled by default. To disable it, set the \texttt{MPI\_MEMMAP\_OFF} environment variable. Memory mapping must be enabled to allow single-copy transfers, MPI remote memory access (RMA) one-sided communication, support for the SHMEM model, and certain collective optimizations.

\section*{Memory Placement and Policies}

The MPI library takes advantage of NUMA placement functions that are available. Usually, the default placement is adequate. However, you can set one or more environment variables to modify the default behavior.

For a complete list of the environment variables that control memory placement, see the \texttt{MPI(1)} man page.

The following topics contain information on environment variables and tools that enable you to tune memory placement:

- "\texttt{MPI\_DSM\_CPULIST}" on page 57
- "\texttt{MPI\_DSM\_DISTRIBUTE}" on page 58
- "\texttt{MPI\_DSM\_VERBOSE}" on page 59
- "Using dplace" on page 59

\section*{\texttt{MPI\_DSM\_CPULIST}}

The \texttt{MPI\_DSM\_CPULIST} environment variable allows you to select the processors to use for an MPI application. At times, specifying a list of processors on which to run a job can be the best means to insure highly reproducible timings, particularly when running on a dedicated system.

The setting is an ordered list that uses commas (,) and hyphens (-) to specify a mapping of MPI processes to CPUs. If running across multiple hosts, separate the per-host components of the CPU list with a colon (:). When using a hyphen-delineated list, you can specify CPU striding by specifying \texttt{/stride\_distance} after the list.
For example:

<table>
<thead>
<tr>
<th>Value</th>
<th>CPU Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>8, 16, 32</td>
<td>Place three MPI processes on CPUs 8, 16, and 32.</td>
</tr>
<tr>
<td>32, 16, 8</td>
<td>Place the MPI process rank zero on CPU 32, one on 16, and two on CPU 8.</td>
</tr>
<tr>
<td>8–15/2</td>
<td>Place the MPI processes 0 through 3 strided on CPUs 8, 10, 12, and 14.</td>
</tr>
<tr>
<td>8–15, 32–39</td>
<td>Place the MPI processes 0 through 7 on CPUs 8 to 15. Place the MPI processes 8 through 15 on CPUs 32 to 39.</td>
</tr>
<tr>
<td>39–32, 8–15</td>
<td>Place the MPI processes 0 through 7 on CPUs 32 to 39. Place the MPI processes 8 through 15 on CPUs 8 to 15.</td>
</tr>
<tr>
<td>8–15:16–23</td>
<td>Place the MPI processes 0 through 7 on the first host on CPUs 8 through 15. Place MPI processes 8 through 15 on CPUs 16 to 23 on the second host.</td>
</tr>
</tbody>
</table>

Note that the process rank is the `MPI_COMM_WORLD` rank. The interpretation of the CPU values specified in the `MPI_DSM_CPULIST` depends on whether the MPI job is being run within a cpuset, as follows:

- If the job is run outside of a cpuset, the CPUs specify `cpunum` values beginning with 0 and up to the number of CPUs in the system, minus one.

- If the job is run within a cpuset, the default behavior is to interpret the CPU values as relative processor numbers within the cpuset.

The number of processors specified should equal the number of MPI processes that are used to run the application. The number of colon-delineated parts of the list must equal the number of hosts used for the MPI job. If an error occurs in processing the CPU list, the default placement policy is used.

**MPI_DSM_DISTRIBUTE**

The `MPI_DSM_DISTRIBUTE` environment variable ensures that each MPI process gets a physical CPU and memory on the node to which it was assigned.

`MPI_DSM_DISTRIBUTE` assigns MPI ranks, as follows:
• On systems that do not include InfiniBand interconnect, **MPI_DSM_DISTRIBUTE** assigns MPI ranks starting at logical CPU 0 and incrementing until all ranks have been placed.

• On systems that include InfiniBand interconnect, if the job spans hosts, **MPI_DSM_DISTRIBUTE** assigns MPI ranks starting with the CPU that is closest to the first InfiniBand host channel adapter (HCA).

If you set both **MPI_DSM_DISTRIBUTE** and **MPI_DSM_CPULIST**, **MPI_DSM_CPULIST** overrides **MPI_DSM_DISTRIBUTE**.

**MPI_DSM_VERBOSE**

Setting the **MPI_DSM_VERBOSE** environment variable directs MPI to display a synopsis of the NUMA and host placement options being used at run time.

**Using dplace**

The **dplace** tool offers another way to specify the placement of MPI processes within a distributed memory host. The **dplace** tool and MPI interoperate, and allow MPI to better manage placement of certain shared memory data structures.

For information about **dplace** with MPI, see the following:

• The **dplace**(1) man page.

• The *Linux Application Tuning Guide*.

**Tuning MPI/OpenMP Hybrid Codes**

A hybrid MPI/OpenMP application is one in which each MPI process itself is a parallel threaded program. These programs often exploit the OpenMP parallelism at the loop level while also implementing a higher-level parallel algorithm that uses MPI.

Many parallel applications perform better if the MPI processes and the threads within them are pinned to particular processors for the duration of their execution. For ccNUMA systems, this pinning ensures that all local, non-shared memory is allocated on the same memory node as the processor referencing the memory. For all systems, pinning can ensure that some or all of the OpenMP threads stay on processors that share a bus or perhaps a processor cache, which can speed up thread synchronization.
The SGI Message Passing Toolkit (MPT) provides the `omplace(1)` command to help with the placement of OpenMP threads within an MPI program. The `omplace(1)` command causes the threads in a hybrid MPI/OpenMP job to be placed on unique CPUs within the containing cpuset. For example, the threads in a 2-process MPI program with 2 threads per process would be placed as follows:

- Rank 0, thread 0 on CPU 0
- Rank 0, thread 1 on CPU 1
- Rank 1, thread 0 on CPU 2
- Rank 1, thread 1 on CPU 3

The CPU placement is performed by dynamically generating a `dplace(1)` placement file and invoking `dplace`.

For more information, see the following:

- The `omplace(1)` man page.
- The `dplace(1)` man page and the *Linux Application Tuning Guide for SGI X86-64 Based Systems*. Both contain information about `dplace(1)`.
- The *SGI Cpuset Software Guide*.

**Example 7-1 Running a Hybrid MPI/OpenMP Application**

The following command line runs a hybrid MPI/OpenMP application with eight MPI processes that are two-way threaded on two hosts:

```bash
mpirun host1,host2 -np 4 omplace -nt 2 ./a.out
```

- When using the PBS workload manager to schedule the hybrid MPI/OpenMP job, use the following resource allocation specification:
  ```
  #PBS -l select=8:ncpus=2
  ```

- In addition, use the following `mpiexec` command:
  ```
  mpiexec -n 8 omplace -nt 2 ./a.out
  ```

For more information about running SGI MPT programs with PBS, see the following: “Using a Workload Manager to Launch an MPI Application” on page 25.
Tuning Running Applications Across Multiple Hosts

When you run an MPI application across a cluster of hosts, you can use the environment variables in this topic to improve application performance across these hosts.

Table 7-1 on page 61 shows the interconnect types and the run-time environment settings and configurations that you can use to improve performance.

Table 7-1 Available Interconnects and the Inquiry Order for Available Interconnects

<table>
<thead>
<tr>
<th>Interconnect Type</th>
<th>Default Order of Selection</th>
<th>Environment Variable Required</th>
</tr>
</thead>
<tbody>
<tr>
<td>XPMEM</td>
<td>1</td>
<td>MPI_USE_XPMEM</td>
</tr>
<tr>
<td>Intel Omni-Path Architecture</td>
<td>2</td>
<td>MPI_USE_OPA</td>
</tr>
<tr>
<td>InfiniBand</td>
<td>3</td>
<td>MPI_USE_IB</td>
</tr>
<tr>
<td>InfiniBand Unreliable Datagram</td>
<td>4</td>
<td>MPI_USE_UD</td>
</tr>
<tr>
<td>TCP/IP</td>
<td>5</td>
<td>MPI_USE_TCP</td>
</tr>
</tbody>
</table>

Table 7-1 on page 61 shows the different types of interconnects that systems can employ as the multihost interconnect. When launched as a distributed application, MPI probes for these interconnects at job startup. For information about how to launch a distributed application, see "Using the mpirun Command to Launch an MPI Application" on page 27.

When MPI detects a high-performance interconnect, MPI attempts to use this interconnect, if it is available, on every host being used by the MPI job. If the interconnect is not available for use on every host, the library attempts to use the next slower interconnect until this connectivity requirement is met. Table 7-1 on page 61 specifies the order in which MPI probes for available interconnects.

The third column of Table 7-1 on page 61 indicates the environment variable you can set to pick a particular interconnect other than the default. In general, to insure the best application performance, allow MPI to pick the fastest available interconnect.
When using the TCP/IP interconnect, unless specified otherwise, MPI uses the default IP adapter for each host. To use a nondefault adapter, enter the adapter-specific host name on the `mpirun` command line.

The following environment variables enable you to tune your application for multiple hosts:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_IB_RAILS</td>
<td>When this variable is set to 1 and the MPI library uses the InfiniBand driver as the inter-host interconnect, SGI MPT sends InfiniBand traffic over the first fabric that it detects. When this variable is set to 2, the library tries to use multiple, available, separate, InfiniBand fabrics and splits the traffic across them. If the separate InfiniBand fabrics do not have unique subnet IDs, then the <code>rail-config</code> utility is required. It must be run by the system administrator to enable the library to correctly use the separate fabrics. The default is 1 on all SGI UV systems.</td>
</tr>
<tr>
<td>MPI_IB_SINGLE_COPY_BUFFER_MAX</td>
<td>If MPI transfers data over InfiniBand and if the size of the cumulative data is greater than this value, then MPI attempts to send the data directly between the processes’s buffers and not through intermediate buffers inside the MPI library. The default is 32767.</td>
</tr>
<tr>
<td>MPI_USE_IB</td>
<td>When set, the MPI library uses the InfiniBand driver as the interconnect when running across multiple hosts or running with multiple binaries. SGI MPT requires the OFED software stack when the InfiniBand interconnect is used. If InfiniBand is used, the <code>MPI_COREDUMP</code> environment variable is forced to <code>INHIBIT</code>, to comply with the InfiniBand driver restriction that no <code>fork()</code> actions occur after InfiniBand resources have been allocated.</td>
</tr>
</tbody>
</table>
The default is `false`.

For more information on these environment variables, see the `ENVIRONMENT VARIABLES` section of the `mpi(1)` man page.

**Tuning for Running Applications over the InfiniBand Interconnect**

When running an MPI application across a cluster of hosts using the InfiniBand interconnect, there are run-time environment variables that you can set to to improve application performance. The following are these variables:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MPI_COLL_IB_OFFLOAD</strong></td>
<td>Enables or disables the Mellanox fabric collectives accelerator (FCA) offload. If FCA offload is configured on your cluster, set <code>MPI_COLL_IB_OFFLOAD=true</code>. You might also need to set Mellanox’s <code>fca_ib_dev_name</code> and <code>fca_ib_port_num</code> environment variables to the name and port of the host channel adapter (HCA) you want to use. For example, <code>fca_ib_dev_name=mlx4_0</code> and <code>fca_ib_port_num=1</code>. The default is <code>MPI_COLL_IB_OFFLOAD=false</code>.</td>
</tr>
<tr>
<td><strong>MPI_CONNECTIONS_THRESHOLD</strong></td>
<td>For very large MPI jobs, the time and resource cost to create a connection between every pair of ranks at job start time can be prodigious. When the number of ranks is set to at least this value, the MPI library creates InfiniBand connections on a demand basis. The default is 1025 ranks.</td>
</tr>
<tr>
<td><strong>MPI_IB_FAILOVER</strong></td>
<td>When the MPI library uses InfiniBand fabric and this variable is set, if an InfiniBand transmission error occurs, SGI MPT tries to restart the connection to the other rank a certain number of times. The <code>MPI_IB_FAILOVER</code> variable specifies the number of times SGI MPT tries to restart the connection. SGI MPT can handle a number of errors of this type between any pair of ranks equal to the value of this variable. The default is 32 times.</td>
</tr>
</tbody>
</table>
MPI_IB_PAYLOAD

When the MPI library uses InfiniBand fabric, it allocates memory for each message header that it uses for InfiniBand. If the size of data to be sent is not greater than this amount minus 64 bytes for the actual header, the data is inlined with the header. If the size is greater than this value, then the message is sent through remote direct memory access (RDMA) operations. The default is 16512 bytes.

MPI_IB_RNR_TIMER

When a packet arrives at an InfiniBand host channel adaptor (HCA) and there are no remaining receive buffers for it, the receiving HCA sends a negative acknowledgement (NAK) to the requestor. The requesting HCA tries again after some period of time, and this variable controls the delay time.

If you set a value higher than the default, performance can degrade in some circumstances. The higher value, however, is likely to improve fabric health significantly during high congestion. For precise translations of this value to delay times, see Table 45 of the official InfiniBand specification. The default is 14.

MPI_IB_TIMEOUT

When an InfiniBand card sends a packet, it waits some amount of time for an ACK packet to be returned by the receiving InfiniBand card. If it does not receive one, it sends the packet again. This variable controls the wait period. The time spent is equal to $4.096 \times 2^n$, where $n$ is specified by the MPI_IB_TIMEOUT variable. By default, the variable is set to 18, and the time spent is expressed in microseconds.

MPI_NUM_MEMORY_REGIONS

For zero-copy sends over the InfiniBand interconnect, SGI MPT keeps a cache of application data buffers registered for these transfers. This environment variable controls the size of the cache. If the application rarely reuses data buffers, it may make sense to set this value to 0 to avoid cache trashing. By default, this variable is set to 1024 (1K). The possible range is from 0 to 8192 (8K).
MPI_NUM_QUICKS

Controls the number of other ranks that a rank can receive from over InfiniBand using a short message fast path. This is 8 by default and can be any value between 0 and 32.

MPI on SGI UV Systems

Note: This section does not apply to SGI UV 30 systems, SGI UV 10 systems, or SGI UV 20 systems.

The SGI® UV™ series systems are scalable, nonuniform memory access (NUMA) systems that support a single Linux image of thousands of processors distributed over many sockets and many SGI UV hub application-specific integrated circuits (ASICs). The SGI UV hub is the heart of the SGI UV system compute blade. Each processor is a hyperthread on a particular core within a particular socket. Typically, each SGI UV hub connects to two sockets. All communication between the sockets and the SGI UV hub uses Intel QuickPath Interconnect (QPI) channels. The following information pertains to specific SGI UV systems:

- On SGI UV 3000 systems and SGI UV 300 systems, the SGI UV hub board assembly has an SGI UV hub ASIC with two identical hubs. Each hub supports one 9.6 GT/s QPI channel to a processor socket. On SGI UV 3000 systems and the SGI UV 300 systems, the hub has eight NUMAlink 7 ports that connect with the NUMAlink 7 interconnect fabric.

- On SGI UV 2000 systems, the SGI UV hub board assembly has an SGI UV hub ASIC with two identical hubs. Each hub supports one 8.0 GT/s QPI channel to a processor socket. The SGI UV 2000 series hub has eight NUMAlink 6 ports that connect with the NUMAlink 6 interconnect fabric.

- The SGI UV 1000 system’s hub has four NUMAlink 5 ports that connect with the NUMAlink 5 interconnect fabric.

The SGI UV hub acts as a crossbar between the processors, local SDRAM memory, and the network interface. The hub ASIC enables any processor in the single-system image (SSI) to access the memory of all processors in the SSI.

When MPI communicates between processes, two transfer methods are possible on an SGI UV system:
• By use of shared memory
• By use of the global reference unit (GRU), part of the SGI UV hub ASIC

MPI chooses the method depending on internal heuristics, the type of MPI communication that is involved, and some user-tunable variables. When using the GRU to transfer data and messages, the MPI library uses the GRU resources it allocates via the GRU resource allocator, which divides up the available GRU resources. It fairly allocates buffer space and control blocks between the logical processors being used by the MPI job.

For more information about the SGI UV hub, SGI UV compute blades, QPI, and NUMAlink 5, or NUMAlink 6, see your SGI hardware documentation.

The following topics contain more information about using MPI on SGI UV systems:
• “General Considerations” on page 66
• "Performance Problems and Corrective Actions" on page 66
• "Other ccNUMA Performance Considerations" on page 67

General Considerations

To run an MPI job optimally on an SGI UV system, it is best to pin MPI processes to CPUs and isolate multiple MPI jobs onto different sets of sockets and hubs. To accomplish this, you can configure a workload manager to create a cpuset for every MPI job. MPI pins its processes to the sequential list of logical processors within the containing cpuset by default, but you can control and alter the pinning pattern using the following:

• MPI_DSM_CPULIST. For more information, see "MPI_DSM_CPULIST" on page 57.
• omplace(1)
• dplace(1)

Performance Problems and Corrective Actions

The MPI library chooses buffer sizes and communication algorithms in an attempt to deliver the best performance to a wide variety of MPI applications automatically. The following list of performance problems can be remedied:
• Odd HyperThreads are idle.

Most high performance computing MPI programs run best using only one HyperThread per core. When an SGI UV system has multiple HyperThreads per core, logical CPUs are numbered such that odd HyperThreads are the high half of the logical CPU numbers. Therefore, the task of scheduling only on the even HyperThreads can be accomplished by scheduling MPI jobs as if only half the full number exist, leaving the high logical CPUs idle. You can use the `cpumap(1)` command to determine if cores have multiple HyperThreads on your SGI UV system. The output shows the following:

- The number of physical and logical processors.
- Whether HyperThreading is on or off.
- The way in which shared processors are paired. This information appears towards the bottom of the command’s output.

If an MPI job uses only half of the available logical CPUs, set `GRU_RESOURCE_FACTOR` to 2 so that the MPI processes can use all the available GRU resources on a hub rather than reserving some of them for the idle HyperThreads. For more information about GRU resource tuning, see `gru_resource(3)`.

• MPI large message bandwidth is inappropriate.

Some programs transfer large messages via the `MPI_Send` function. To use unbuffered, single-copy transport in these cases, set `MPI_BUFFER_MAX=0`. For more information, see `MPI(1)`.

• MPI small or near messages are very frequent.

For small fabric hop counts, shared memory message delivery is faster than using GRU messages. To deliver all messages within an SGI UV host via shared memory, set `MPI_SHARED_NEIGHBORHOOD=HOST`. For more information, see `MPI(1)`.

Other ccNUMA Performance Considerations

MPI application processes typically perform better if their local memory is allocated on the socket assigned to execute the process. This cannot happen if memory on that socket is exhausted, either by the application itself or by other system consumption (for example, by file buffer cache).
You can use the `nodeinfo(1)` command to view memory consumption on the nodes assigned to your job, and you can use the `bcfree(1)` command to clear out excessive file buffer cache. PBS Professional workload manager installations can be configured to issue `bcfree(1)` commands in the job prologue.

For more information, see the PBS Professional documentation and `bcfree(1)`.

**Suspending MPI Jobs**

Internally, the MPI software from SGI uses the XPMEM kernel module to provide direct access to data on remote partitions and to provide single-copy operations to local data. The XPMEM kernel module prevents any pages used by these operations from paging. If an administrator needs to temporarily suspend an MPI application to allow other applications to run, they can unpin these pages so they can be swapped out and made available for other applications.

Each process of an MPI application that is using the XPMEM kernel module has a `/proc/xpmem/pid` file associated with it. File `/proc/xpmem/pid` includes the number of pages owned by this process that are prevented from paging by XPMEM. You can display the content of this file. For example:

```
# cat /proc/xpmem/5562
pages pinned by XPMEM: 17
```

The following procedure explains how to unpin the pages for use by other processes.

**Procedure 7-1** To unpin pages

1. Log in as the system administrator.
2. Suspend all the processes in the application.
3. Use the `echo(1)` command to unpin the pages.

   You can echoing any value into the `/proc/xpmem/pid` file.

   For `pid`, specify the process ID.

   The echo command does not return until that process’s pages are unpinned.

   For example:

   ```
   # echo 1 > /proc/xpmem/5562
   ```
When the MPI application is resumed, the XPMEM kernel module prevents the pages from paging as they are referenced by the application.
Chapter 8

MPI Performance Profiling

This chapter includes the following topics:

- "About MPI Performance Profiling" on page 71
- "Using perfcatch(1)" on page 72
- "Writing Your Own Profiling Interface" on page 77
- "Using Third-party Profilers" on page 78
- "MPI Internal Statistics" on page 78

About MPI Performance Profiling

Performance profiling occurs when you run your MPI program or SHMEM program with a tool that can aggregate run time statistics. Profiling tools gather statistics that show the amount of time that your program spends in MPI, the number of messages sent, or the number of bytes sent. SGI includes profiling support in the libmpi.so library. When you use a profiling tool, the tool automatically replaces all MPI_Xxx prototypes and function names with PMPI_Xxx entry points.

This chapter describes the use of profiling tools to obtain performance information. Compared to the performance analysis of sequential applications, characterizing the performance of parallel applications can be challenging. Often it is most effective to first focus on improving the performance of MPI applications at the single process level.

It may also be important to understand the message traffic generated by an application. A number of tools can be used to analyze this aspect of a message passing application’s performance, including SGI’s MPInside and various third-party products.

The following topics contain more information about profiling:

- **MPInside Reference Guide.** This manual explains how to use the MPInside profiling tool.
- "Using perfcatch(1)" on page 72
- "Writing Your Own Profiling Interface" on page 77
Using perfcatch(1)

You can use SGI’s perfcatch utility to profile the performance of an MPI program or SHMEM program. The perfcatch utility runs the MPI program with the wrapper library, libmpi.so, and writes MPI call profiling information information to MPI_PROFILING_STATS.

The following topics contain more information about perfcatch(1):

• "The perfcatch(1) Command" on page 72
• "MPI_PROFILING_STATS Results File Example" on page 73
• "Environment Variables Used With perfcatch(1)" on page 76

The perfcatch(1) Command

The following format shows how to use the perfcatch command:

```
mpirun [ mpi_params ] perfcatch [ -i ] cmd [ args ]
```

By default, perfcatch assumes an SGI MPT program. The perfcatch utility accepts the following options:

- `mpi_params` Optional. Specifies the MPI parameters needed to launch the program.
- `-i` Specifies to use Intel MPI.
- `cmd` Specifies the name of the executable program. For example, a.out.
- `args` Optional. Specifies additional command line arguments.

To use perfcatch with an SGI Message Passing Toolkit MPI program, insert the perfcatch command in front of the executable file name, as the following examples show:

• `mpirun -np 64 perfcatch a.out arg1`
• `mpirun host1 32, host2 64 perfcatch a.out arg1`
To use perfcatch with Intel MPI, add the -i option, as follows:

```bash
mpiexec -np 64 perfcatch -i a.out arg1
```

For more information, see the `perfcatch(1)` man page.

**MPI_PROFILING_STATS Results File Example**

The `perfcatch(1)` utility’s output file is called `MPI_PROFILING_STATS`. Upon program completion, the `MPI_PROFILING_STATS` file resides in the current working directory of the MPI process with rank 0.

This output file includes a summary statistics section followed by a rank-by-rank profiling information section. The summary statistics section reports some overall statistics. These statistics include the percent time each rank spent in MPI functions and the MPI process that spent the least and the most time in MPI functions. Similar reports are made about system time usage.

In the rank-by-rank profiling information, there is a list of every profiled MPI function called by a particular MPI process. The report includes the number of calls and the total time consumed by these calls. Some functions report additional information, such as average data counts and communication peer lists.

The following is an example `MPI_PROFILING_STATS` results file:
PERFCATCHER version 22
(C) Copyright SGI. This library may only be used
on SGI hardware platforms. See LICENSE file for
details.

MPI program profiling information
Job profile recorded Wed Jan 17 13:05:24 2007
Program command line: /home/estes01/michel/sastest/mpi_hello_linux
Total MPI processes 2

Total MPI job time, avg per rank 0.0054768 sec
Profiled job time, avg per rank 0.0054768 sec
Percent job time profiled, avg per rank 100%

Total user time, avg per rank 0.001 sec
Percent user time, avg per rank 18.2588%
Total system time, avg per rank 0.0045 sec
Percent system time, avg per rank 82.1648%

Time in all profiled MPI routines, avg per rank 5.75004e-07 sec
Percent time in profiled MPI routines, avg per rank 0.0104989%

Rank-by-Rank Summary Statistics

Rank-by-Rank: Percent in Profiled MPI routines

<table>
<thead>
<tr>
<th>Rank</th>
<th>Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.0112245%</td>
</tr>
<tr>
<td>1</td>
<td>0.00968502%</td>
</tr>
</tbody>
</table>

Least: Rank 1 0.00968502%
Most: Rank 0 0.0112245%
Load Imbalance: 0.000771%

Rank-by-Rank: User Time

<table>
<thead>
<tr>
<th>Rank</th>
<th>Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>17.2683%</td>
</tr>
<tr>
<td>1</td>
<td>19.3699%</td>
</tr>
</tbody>
</table>

Least: Rank 0 17.2683%
Most: Rank 1 19.3699%

Rank-by-Rank: System Time

<table>
<thead>
<tr>
<th>Rank</th>
<th>Percent</th>
</tr>
</thead>
</table>

74
0:86.3416%  1:77.4796%
Least:  Rank 1  77.4796%
Most:  Rank 0  86.3416%

Notes
-----
Wtime resolution is 5e-08 sec

Rank-by-Rank MPI Profiling Results
-----------------------------------
Activity on process rank 0

Single-copy checking was not enabled.
comm_rank  calls:  1  time: 6.50005e-07 s  6.50005e-07 s/call

Activity on process rank 1

Single-copy checking was not enabled.
comm_rank  calls:  1  time: 5.00004e-07 s  5.00004e-07 s/call

recv profile
            cnt/sec for all remote ranks
  local  ANY_SOURCE  0  1
  rank

recv wait for data profile
            cnt/sec for all remote ranks
  local  0  1
  rank

recv wait for data profile
8: MPI Performance Profiling

<table>
<thead>
<tr>
<th>cnt/sec for all remote ranks</th>
</tr>
</thead>
<tbody>
<tr>
<td>local</td>
</tr>
<tr>
<td>rank</td>
</tr>
</tbody>
</table>

send profile

<table>
<thead>
<tr>
<th>cnt/sec for all destination ranks</th>
</tr>
</thead>
<tbody>
<tr>
<td>src</td>
</tr>
<tr>
<td>rank</td>
</tr>
</tbody>
</table>

ssend profile

<table>
<thead>
<tr>
<th>cnt/sec for all destination ranks</th>
</tr>
</thead>
<tbody>
<tr>
<td>src</td>
</tr>
<tr>
<td>rank</td>
</tr>
</tbody>
</table>

ibsend profile

<table>
<thead>
<tr>
<th>cnt/sec for all destination ranks</th>
</tr>
</thead>
<tbody>
<tr>
<td>src</td>
</tr>
<tr>
<td>rank</td>
</tr>
</tbody>
</table>

**Environment Variables Used With** perfcatch(1)

The MPI performance-profiling environment variables are as follows:

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI_PROFILE_AT_INIT</td>
<td>Activates MPI profiling immediately, that is, at the start of MPI program execution. To use this environment variable, set it to any value. For example, set MPI_PROFILE_AT_INIT to 1.</td>
</tr>
</tbody>
</table>
MPI_PROFILING_STATS_FILE  Specifies the perfcatch output file. This is the file to which MPI profiling results are written. By default, the profiler writes to MPI_PROFILING_STATS.

Writing Your Own Profiling Interface

You can write your own profiler by using the MPI standard PMPI_* calls. In addition, either within your own profiling library or within the application itself, you can use the MPI_Wtime function call to time specific calls or sections of your code.

The following example output is for a single rank of a program that was run on 128 processors using a user-created profiling library that performs call counts and timings of common MPI calls. Notice that for this rank, most of the MPI time is spent in MPI_Waitall and MPI_Allreduce.

Total job time 2.203333e+02 sec
Total MPI processes 128
Wtime resolution is 8.000000e-07 sec

activity on process rank 0
comm_rank calls 1  time 8.800002e-06
get_count calls 0  time 0.000000e+00
ibsend calls 0  time 0.000000e+00
probe calls 0  time 0.000000e+00
recv calls 0  time 0.000000e+00 avg datacnt 0  waits 0  wait time 0.000000e+00
irecv calls 22039  time 9.76185e-01  datacnt 23474032 avg datacnt 1065
send calls 0  time 0.000000e+00
ssend calls 0  time 0.000000e+00
isend calls 22039  time 2.950286e+00 datacnt 23474032 avg datacnt 1065
wait calls 0  time 0.000000e+00 avg datacnt 0
waitall calls 11045  time 7.73805e+01  # of Reqs 44078  avg data cnt 137944
barrier calls 680  time 5.133110e+00
alltoall calls 0  time 0.0e+00  avg datacnt 0
alltoallv calls 0  time 0.000000e+00
reduce calls 0  time 0.000000e+00
allreduce calls 4658  time 2.072872e+01
bcast calls 680  time 6.915840e-02
gather calls 0  time 0.000000e+00
gatherv calls 0  time 0.000000e+00
scatter calls 0  time 0.000000e+00
scatterv calls 0  time 0.000000e+00

activity on process rank 1
...

Using Third-party Profilers

You can use third-party profiling tools with SGI MPI. The following are examples of tools to consider:

- The TAU Performance System profiler from the University of Oregon. This software is a portable profiling and tracing toolkit for performance analysis of parallel programs written in Fortran, C, C++, UPC, Java, and Python.

- The Allinea MAP profiler. The Allinea MAP profiler is part of the Allinea Forge toolkit

MPI Internal Statistics

MPI keeps track of certain resource utilization statistics. You can use these statistics to determine potential performance problems caused by a lack of MPI message buffers or other MPI internal resources.

To display MPI internal statistics, use the MPI_STATS environment variable or the -stats option on the mpirun command. MPI internal statistics are always being gathered, so displaying them does not cause significant additional overhead. In addition, one can sample the MPI statistics counters from within an application, allowing for finely grained measurements.

If the MPI_STATS_FILE environment variable is set, when the program completes, the system writes internal statistics to the file specified by this variable.

These statistics can be very useful in optimizing codes in the following ways:

- To determine if there are enough internal buffers and if processes are waiting (retries) to acquire them

- To determine if single copy optimization is being used for point-to-point or collective calls
For additional information on how to use the MPI statistics counters to help tune the run-time environment for an MPI application, see Chapter 7, "Run-time Tuning" on page 53.
Chapter 9

Troubleshooting and Frequently Asked Questions

This chapter provides answers to some common problems that users encounter when they start to use SGI MPI and provides answers to other frequently asked questions. It covers the following topics:

- "What are some things I can try to figure out why mpirun is failing? " on page 81
- "My code runs correctly until it reaches MPI_Finalize() and then it hangs." on page 83
- "My hybrid code (using OpenMP) stalls on the mpirun command." on page 83
- "I keep getting error messages about MPI_REQUEST_MAX being too small." on page 83
- "I am not seeing stdout and/or stderr output from my MPI application." on page 84
- "How can I get the SGI Message Passing Toolkit (MPT) software to install on my machine?" on page 84
- "Where can I find more information about the SHMEM programming model? " on page 84
- "The ps(1) command says my memory use (SIZE) is higher than expected. " on page 84
- "What does MPI: could not run executable mean?" on page 85
- "How do I combine MPI with insert favorite tool here?" on page 85
- "Why do I see “stack traceback” information when my MPI job aborts?" on page 86

**What are some things I can try to figure out why mpirun is failing?**

Here are some things to investigate:

- Look in /var/log/messages for any suspicious errors or warnings. For example, if your application tries to pull in a library that it cannot find, a message should appear here. Only the root user can view this file.
• Be sure that you did not misspell the name of your application.

• To find dynamic link errors, try to run your program without mpirun. You will get the “mpirun must be used to launch all MPI applications” message, along with any dynamic link errors that might not be displayed when the program is started with mpirun.

As a last resort, setting the environment variable LD_DEBUG to all will display a set of messages for each symbol that rld resolves. This produces a lot of output, but should help you find the cause of the link error.

• Be sure that you are setting your remote directory properly. By default, mpirun attempts to place your processes on all machines into the directory that has the same name as $PWD. This should be the common case, but sometimes different functionality is required. For more information, see the section on $MPI_DIR and/or the -dir option in the mpirun man page.

• If you are using a relative pathname for your application, be sure that it appears in $PATH. In particular, mpirun will not look in '.' for your application unless '.' appears in $PATH.

• Run /usr/sbin/ascheck to verify that your array is configured correctly.

• Use the mpirun -verbose option to verify that you are running the version of MPI that you think you are running.

• Be very careful when setting MPI environment variables from within your .cshrc or .login files, because these will override any settings that you might later set from within your shell (due to the fact that MPI creates the equivalent of a fresh login session for every job). The safe way to set things up is to test for the existence of $MPI_ENVIRONMENT in your scripts and set the other MPI environment variables only if it is undefined.

• If you are running under a Kerberos environment, you may experience unpredictable results because currently, mpirun is unable to pass tokens. For example, in some cases, if you use telnet to connect to a host and then try to run mpirun on that host, it fails. But if you instead use rsh to connect to the host, mpirun succeeds. (This might be because telnet is kerberized but rsh is not.) At any rate, if you are running under such conditions, you will definitely want to talk to the local administrators about the proper way to launch MPI jobs.

• Look in /tmp/.arraysvcs on all machines you are using. In some cases, you might find an errlog file that may be helpful.
• You can increase the verbosity of the Array Services daemon (arrayd) using the -v option to generate more debugging information. For more information, see the arrayd(8) man page.

• Check error messages in /var/run/arraysvcs.

My code runs correctly until it reaches MPI_Finalize() and then it hangs.

This is almost always caused by send or recv requests that are either unmatched or not completed. An unmatched request is any blocking send for which a corresponding recv is never posted. An incomplete request is any nonblocking send or recv request that was never freed by a call to MPI_Test(), MPI_Wait(), or MPI_Request_free().

Common examples are applications that call MPI_Isend() and then use internal means to determine when it is safe to reuse the send buffer. These applications never call MPI_Wait(). You can fix such codes easily by inserting a call to MPI_Request_free() immediately after all such isend operations, or by adding a call to MPI_Wait() at a later place in the code, prior to the point at which the send buffer must be reused.

My hybrid code (using OpenMP) stalls on the mpirun command.

If your application was compiled with the Open64 compiler, make sure you follow the instructions about using the Open64 compiler in combination with MPI/OpenMP applications described in "Compiling and Linking the MPI Program" on page 23.

I keep getting error messages about MPI_REQUEST_MAX being too small.

There are two types of cases in which the MPI library reports an error concerning MPI_REQUEST_MAX. The error reported by the MPI library distinguishes these.

MPI has run out of unexpected request entries; the current allocation level is: XXXXXX

The program is sending so many unexpected large messages (greater than 64 bytes) to a process that internal limits in the MPI library have been exceeded. The options here
Troubleshooting and Frequently Asked Questions

are to increase the number of allowable requests via the MPI_REQUEST_MAX shell variable, or to modify the application.

MPI has run out of request entries;
the current allocation level is: MPI_REQUEST_MAX = XXXXX

You might have an application problem. You almost certainly are calling
MPI_Isend() or MPI_Irecv() and not completing or freeing your request objects.
You need to use MPI_Request_free(), as described in the previous section.

I am not seeing stdout and/or stderr output from my MPI application.

All stdout and stderr is line-buffered, which means that mpirun does not print
any partial lines of output. This sometimes causes problems for codes that prompt
the user for input parameters but do not end their prompts with a newline character.
The only solution for this is to append a newline character to each prompt.

You can set the MPI_UNBUFFERED_STDIO environment variable to disable
line-buffering. For more information, see the MPI(1) and mpirun(1) man pages.

How can I get the SGI Message Passing Toolkit (MPT) software to install on
my machine?

SGI MPT RPMs are included in the SGI Performance Suite releases. In addition, you
can obtain SGI MPT RPMs from the SGI customer portal at the following URL:
https://support.sgi.com

Where can I find more information about the SHMEM programming model?

See the intro_shmem(3) man page.

The ps(1) command says my memory use (SIZE) is higher than expected.

At MPI job start-up, MPI calls the SHMEM library to cross-map all user static memory
on all MPI processes to provide optimization opportunities. The result is large virtual
memory usage. The ps(1) command’s SIZE statistic is telling you the amount of
virtual address space being used, not the amount of memory being consumed. Even if all of the pages that you could reference were faulted in, most of the virtual address regions point to multiply-mapped (shared) data regions, and even in that case, actual per-process memory usage would be far lower than that indicated by SIZE.

**What does MPI: could not run executable mean?**

This message means that something happened while `mpirun` was trying to launch your application, which caused it to fail before all of the MPI processes were able to handshake with it.

The `mpirun` command directs `arrayd` to launch a master process on each host and listens on a socket for those masters to connect back to it. Since the masters are children of `arrayd`, `arrayd` traps SIGCHLD and passes that signal back to `mpirun` whenever one of the masters terminates. If `mpirun` receives a signal before it has established connections with every host in the job, it knows that something has gone wrong.

**How do I combine MPI with insert favorite tool here?**

In general, the rule to follow is to run `mpirun` on your tool and then the tool on your application. Do not try to run the tool on `mpirun`. Also, because of the way that `mpirun` sets up stdio, seeing the output from your tool might require a bit of effort. The most ideal case is when the tool directly supports an option to redirect its output to a file. In general, this is the recommended way to mix tools with `mpirun`. Of course, not all tools (for example, dplace) support such an option. However, it is usually possible to make it work by wrapping a shell script around the tool and having the script do the redirection, as in the following example:

```bash
> cat myscript
#!/bin/sh

setenv MPI_DSM_OFF

dplace -verbose a.out 2> outfile
> mpirun -np 4 myscript

hello world from process 0
```
hello world from process 1
hello world from process 2
hello world from process 3

> cat outfile
there are now 1 threads
Setting up policies and initial thread.
Migration is off.
Data placement policy is PlacementDefault.
Creating data PM.
Data pagesize is 16k.
Setting data PM.
Creating stack PM.
Stack pagesize is 16k.
Stack placement policy is PlacementDefault.
Setting stack PM.
there are now 2 threads
there are now 3 threads
there are now 4 threads
there are now 5 threads

---

Caution: The preceding script example is for illustrative purposes only and has not been evaluated for use in a production environment.

Why do I see “stack traceback” information when my MPI job aborts?

More information can be found in the **MPI(1)** man page in descriptions of the **MPI_COREDUMP** and **MPI_COREDUMP_DEBUGGER** environment variables.
Chapter 10

Array Services

This chapter includes the following topics:

• "About Array Services" on page 87
• "Retrieving the Array Services Release Notes" on page 88
• "Managing Local Processes" on page 89
• "Using Array Services Commands" on page 90
• "Array Services Commands" on page 91
• "Obtaining Information About the Array" on page 94
• "Additional Array Configuration Information" on page 97
• "Configuring Array Commands" on page 103

About Array Services

The SGI Array Services software enables parallel applications to run on multiple hosts in a cluster, or array. Array Services provides cluster job launch capabilities for SGI Message Passing Toolkit jobs.

The array can consist of the following:

• Multiple single system images (SSIs) on an SGI UV system
• Multiple compute nodes plus a service node on an SGI ICE or SGI ICE X system
• Multiple physical machines

An array system is bound together with a high-speed network and the Array Services software. Array users can access the system with familiar commands for job control, login and password management, and remote execution. Array Services facilitates global session management, array configuration management, batch processing, message passing, system administration, and performance visualization.

The Array Services software package includes the following:
An array daemon that runs on each node. The daemon groups logically related processes together across multiple nodes. The process groups create a global process namespace across the array, facilitate accounting, and facilitate administration.

The daemon maintains information about node configuration, process IDs, and process groups. Array daemons on the nodes cooperate with each other.

An array configuration database. The database describes the array configuration and provides reference information for array daemons and user programs. Each node hosts a copy of the array configuration database.

Commands, libraries, and utilities such as ainfo(1), arshell(1), and others.

The Message Passing Interface (MPI) of SGI MPI uses Array Services to launch parallel applications.

SGI includes MUNGE software in the SGI MPI software distribution. This optional, open-source product provides secure Array Services functionality. MUNGE allows a process to authenticate the UID and GID of another local or remote process within a group of hosts that have common users and groups. MUNGE authentication, which also includes the Array Services data exchanged in the array, is encrypted. For more information about MUNGE, see the MUNGE website, which is at the following location:

http://dun.github.io/munge/

The Array Services package requires that the process sets service be installed and running. This package is provided in the sgi-procset RPM. You can type the following commands to verify that the process sets service is installed and running:

```
# rpm -q sgi-procset
# /etc/init.d/procset status
```

Retrieving the Array Services Release Notes

The following procedure explains how to find the array services release note information.
Procedure 10-1 To retrieve Array Services release note information

1. Type the following command to retrieve the location of the Array Services release notes:

   ```
   # rpm -qi sgi-arraysvcs
   /usr/share/doc/sgi-arraysvcs-3.7/README.relnotes
   ```

2. Use a text editor or other command to display the file that the `rpm(8)` command returns.

Managing Local Processes

Each UNIX process has a process identifier (PID), a number that identifies that process within the node where it runs. It is important to realize that a PID is local to the node; so it is possible to have processes in different nodes using the same PID numbers.

Within a node, processes can be logically grouped in process groups. A process group is composed of a parent process together with all the processes that it creates. Each process group has a process group identifier (PGID). Like a PID, a PGID is defined locally to that node, and there is no guarantee of uniqueness across the array.

Monitoring Local Processes and System Usage

You query the status of processes using the system command `ps`. To generate a full list of all processes on a local system, use a command such as the following:

   ```
   ps -elfj
   ```

You can monitor the activity of processes using the command `top` for an ASCII display in a terminal window.

Scheduling and Killing Local Processes

You can schedule commands to run at specific times using the `at` command. You can kill or stop processes using the `kill` command. To destroy the process with PID 13032, use a command such as the following:

   ```
   kill -KILL 13032
   ```
Summary of Local Process Management Commands

Table 10-1 on page 90 summarizes information about local process management.

Table 10-1 Information Sources: Local Process Management

<table>
<thead>
<tr>
<th>Topic</th>
<th>Man Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Process ID and process group</td>
<td>intro(2)</td>
</tr>
<tr>
<td>Listing and monitoring processes</td>
<td>ps(1), top(1)</td>
</tr>
<tr>
<td>Running programs at low priority</td>
<td>nice(1), batch(1)</td>
</tr>
<tr>
<td>Running programs at a scheduled time</td>
<td>at(1)</td>
</tr>
<tr>
<td>Terminating a process</td>
<td>kill(1)</td>
</tr>
</tbody>
</table>

Using Array Services Commands

When an application starts processes on more than one node, the PID and PGID are no longer adequate to manage the application. The Array Services commands enable you to view the entire array and to control the processes on multinode programs.

You can type Array Services commands from any workstation connected to an array system. You do not have to be logged in to an array node. Table 10-2 on page 90 shows the commands that are common to Array Services operations.

Table 10-2 Common Array Services Commands

<table>
<thead>
<tr>
<th>Topic</th>
<th>Man Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Array Services Overview</td>
<td>array_services(5)</td>
</tr>
<tr>
<td>ainfo command</td>
<td>ainfo(1)</td>
</tr>
<tr>
<td>array command</td>
<td>array(1) or arrayd.conf(4)</td>
</tr>
<tr>
<td>arshell command</td>
<td>arshell(1)</td>
</tr>
<tr>
<td>newsess command</td>
<td>newsess(1)</td>
</tr>
</tbody>
</table>
About Array Sessions
Array Services is composed of a daemon—a background process that is started at boot time in every node—and a set of commands such as ainfo(1). The commands call on the daemon process in each node to get the information they need.

One concept that is basic to Array Services is the array session, which is a term for all the processes of one application, wherever they may execute. Normally, your login shell, with the programs you start from it, constitutes an array session. A batch job is an array session, and you can create a new shell with a new array session identity.

About Names of Arrays and Nodes
Each node is a server, and as such each node has a hostname. An array system as a whole has a name, too. In most installations there is only a single array, and you never need to specify which array you mean. However, it is possible to have multiple arrays available on a network, and you can direct Array Services commands to a specific array.

About Authentication Keys
It is possible for the array administrator to establish an authentication code, which is a 64-bit number, for all or some of the nodes in an array. There can be a single authentication code number for each node. Your system administrator can tell you if this is necessary.

When authentication keys are implemented, you need to specify the authentication key as the argument to the -s option on the command line of each Array Services command. The code applies to any command entered at that node or addressed to that node.

Array Services Commands
The Array Services package includes an array daemon, an array configuration database, and several commands. Some utilities enable you to retrieve information about the array. Other utilities let the administrator query and manipulate distributed array applications. The Array Services commands are as follows:
The `ainfo(1)`, `array(1)`, and `arshell(1)` commands accept a common set of options plus some command-specific options. Table 10-3 on page 92 summarizes the common options. The default values of some options are set by environment variables.

**Table 10-3** Array Services Command Option Summary

<table>
<thead>
<tr>
<th>Option</th>
<th>Used In</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-a array</td>
<td>ainfo, array</td>
<td>Specify a particular array when more than one is accessible.</td>
</tr>
<tr>
<td>-D</td>
<td>ainfo, array</td>
<td>Send commands to other nodes directly, rather than through <code>arrayd</code> daemon.</td>
</tr>
<tr>
<td>-F</td>
<td>ainfo, array</td>
<td>Forward commands to other nodes through the <code>arrayd</code> daemon.</td>
</tr>
<tr>
<td>-Kl number</td>
<td>ainfo, array</td>
<td>Authentication key for the local node. This is a 64-bit number.</td>
</tr>
<tr>
<td>-Kr number</td>
<td>ainfo, array</td>
<td>Authentication key for the remote node. This is a 64-bit number.</td>
</tr>
</tbody>
</table>
| -l       | ainfo, array  | Execute in context of the destination node, not necessarily the current node. This option letter is a lowercase letter “L”, for “local”.
### Specifying a Single Node

The `-l` and `-s` options work together. The `-l` option restricts the scope of a command to the node where the command is executed. This option is a lowercase letter “L”, for “local”. By default, that is the node where the command is entered. When `-l` is not used, the scope of a query command is all nodes of the array. The `-s` option directs the command to be executed on a specified node of the array. These options work together in query commands as follows:

- To query all nodes as seen by the local node, use neither option.
- To query only the local node, use only `-l`.
- To query all nodes as seen by a specified node, use only `-s`.
- To query only a particular node, use both `-s` and `-l`.

### Common Environment Variables

The Array Services commands depend on environment variables to define default values for the less-common command options. These variables are summarized in Table 10-4.
Table 10-4 Array Services Environment Variables

<table>
<thead>
<tr>
<th>Variable Name</th>
<th>Use</th>
<th>Default When Undefined</th>
</tr>
</thead>
<tbody>
<tr>
<td>ARRAYD_FORWARD</td>
<td>When defined with a string starting with the letter y, all commands default to forwarding through the array daemon (option -F).</td>
<td>Commands default to direct communication (option -D).</td>
</tr>
<tr>
<td>ARRAYD_PORT</td>
<td>The port (socket) number monitored by the array daemon on the destination node.</td>
<td>The standard number of 5434, or the number given with option -p.</td>
</tr>
<tr>
<td>ARRAYDLOCALKEY</td>
<td>Authentication key for the local node (option -Kl).</td>
<td>No authentication unless the -Kl option is used.</td>
</tr>
<tr>
<td>ARRAYDREMOTEKEY</td>
<td>Authentication key for the destination node (option -Kr).</td>
<td>No authentication unless the -Kr option is used.</td>
</tr>
<tr>
<td>ARRAYD</td>
<td>The destination node, when not specified by the -s option.</td>
<td>The local node, or the node given with -s.</td>
</tr>
</tbody>
</table>

Obtaining Information About the Array

Any user of an array system can use Array Services commands to check the hardware components and the software workload of the array. The commands needed are ainfo and array.

Learning Array Names

If your network includes more than one array system, you can use ainfo arrays at one array node to list all the array names that are configured, as in the following example.

homegrown% ainfo arrays
Arrays known to array services daemon
ARRAY DevArray
   IDENT 0x3381
ARRAY BigDevArray
   IDENT 0x7456
ARRAY test
Array names are configured into the array database by the administrator. Different arrays might know different sets of other array names.

**Learning Node Names**

You can use `ainfo machines` to learn the names and some features of all nodes in the current array, as in the following example.

```
homegrown 175% ainfo -b machines
machine homegrown homegrown 5434 192.48.165.36 0
machine disarray disarray 5434 192.48.165.62 0
machine datarray datarray 5434 192.48.165.64 0
machine tokyo tokyo 5434 150.166.39.39 0
```

In this example, the `-b` option of `ainfo` is used to get a concise display.

**Learning Node Features**

You can use `ainfo nodeinfo` to request detailed information about one or all nodes in the array. To get information about the local node, use `ainfo -l nodeinfo`. However, to get information about only a particular other node, for example node `tokyo`, use `-l` and `-s`, as in the following example:

```
homegrown 181% ainfo -s tokyo -l nodeinfo
Node information for server on machine "tokyo"
MACHINE tokyo
VERSION 1.2
8 PROCESSOR BOARDS
  BOARD: TYPE 15  SPEED 190
    CPU:  TYPE 9  REVISION 2.4
    FPU:  TYPE 9  REVISION 0.0
...
16 IP INTERFACES  HOSTNAME tokyo  HOSTID 0xc01a5035
  DEVICE et0  NETWORK 150.166.39.0  ADDRESS 150.166.39.39 UP
  DEVICE atm0  NETWORK 255.255.255.255  ADDRESS 0.0.0.0 UP
  DEVICE atm1  NETWORK 255.255.255.255  ADDRESS 0.0.0.0 UP
...
0 GRAPHICS INTERFACES
MEMORY
```
Learning User Names and Workload

The system commands who(1), top(1), and uptime(1) are commonly used to get information about users and workload on one server. The array(1) command offers array-wide equivalents to these commands.

Learning User Names

To get the names of all users logged in to the whole array, use array who. To learn the names of users logged in to a particular node, for example tokyo, use -l and -s, as in the following example:

homegrown 180\%  \texttt{array -s tokyo -l who}

\begin{verbatim}
joecd  tokyo  frummage.eng.sgi -tcsh
joecd  tokyo  frummage.eng.sgi -tcsh
benf  tokyo  einstein.ued.sgi. /bin/tcsh
yohn  tokyo  rayleigh.eng.sgi vi +153 fs/procfs/prd
...
\end{verbatim}

The preceding example has been edited for brevity and security.

Learning Workload

Two variants of the array command return workload information. The array-wide equivalent of uptime is array uptime, as follows:

\begin{verbatim}
homegrown 181\%  array uptime
   homegrown:  up 1 day,  7:40,  26 users,  load average: 7.21, 6.35, 4.72
disarray:  up 2:53,  0 user,  load average: 0.00, 0.00, 0.00
dataarray: up 5:34,  1 user,  load average: 0.00, 0.00, 0.00
      tokyo:  up 7 days, 9:11,  17 users,  load average: 0.15, 0.31, 0.29
homegrown 182\%  array -l -s tokyo uptime
      tokyo:  up 7 days, 9:11,  17 users,  load average: 0.12, 0.30, 0.28
\end{verbatim}
The command `array top` lists the processes that are currently using the most CPU time. The output identifies each process by its internal array session handle (ASH) value. The following is example output:

```
homegrown 183% array top
ASH Host PID User %CPU Command
----------------------------------------------------------------
0x1111ffff00000000 homegrown 5 root 1.20 vfs_sync
0x1111ffff000001e9 homegrown 1327 arraysvcs 1.19 atop
0x1111ffff000001e9 tokyo 19816 arraysvcs 0.73 atop
0x1111ffff000001e9 disarray 1106 arraysvcs 0.47 atop
0x1111ffff000001e9 datarray 1423 arraysvcs 0.42 atop
0x1111ffff00000000 homegrown 20 root 0.41 ShareII
0x1111ffff000000c0 homegrown 29683 kchang 0.37 ld
0x1111ffff0000001e homegrown 1324 root 0.17 arrayd
0x1111ffff00000000 homegrown 229 root 0.14 routed
0x1111ffff00000000 homegrown 19 root 0.09 pdflush
0x1111ffff000001e9 disarray 1105 arraysvcs 0.02 atopm
```

The `-l` and `-s` options can be used to select data about a single node, as usual.

**Additional Array Configuration Information**

The system administrator has to initialize the array configuration database, a file that is used by the Array Services daemon in executing almost every `ainfo` and `array` command.

**Security Considerations for Standard Array Services**

The array services daemon, `arrayd(1M)`, runs as root. As with other system services, if it is configured carelessly it is possible for arbitrary and possibly unauthorized user to disrupt or even damage a running system.

By default, most array commands are executed using the user, group, and project ID of either the user that issued the original command, or `arraysvcs`. When adding new array commands to `arrayd.conf`, or modifying existing ones, always use the most restrictive IDs possible in order to minimize trouble if a hostile or careless user were to run that command. Avoid adding commands that run with more powerful IDs, such as user `root` or group `sys`, than the user. If such commands are necessary,
analyze them carefully to ensure that an arbitrary user would not be granted any more privileges than expected, much the same as one would analyze a setuid program.

In the default array services configuration, the arrayd daemon allows all the local requests to access arrayd but not the remote requests. In order to let the remote requests access the arrayd, the AUTHENTICATION parameter needs to be set to NONE in the /usr/lib/array/arrayd.auth file. By default it is set to NOREMOTE. When the AUTHENTICATION parameter is set to NONE, the arrayd daemon assumes that a remote user will accurately identify itself when making a request. In other words, if a request claims to be coming from user abc, the arrayd daemon assumes that it is in fact from user abc and not somebody spoofing abc. This should be adequate for systems that are behind a network firewall or otherwise protected from hostile attack, and in which all the users inside the firewall are presumed to be non-hostile. On systems for which this is not the case, because they are attached to a public network or because individual machines cannot be trusted, the Array Services AUTHENTICATION parameter should be set to NOREMOTE. When AUTHENTICATION is set to NONE, all requests from remote systems are authenticated using a mechanism that involves private keys that are known only to the super-users on the local and remote systems. Requests originating on systems that do not have these private keys are rejected. For more details, see the section on authentication information in the arrayd.conf(4) man page.

The arrayd daemon does not support mapping user, group or project names between two different namespaces; all members of an array are assumed to share the same namespace for users, groups, and projects. Thus, if systems A and B are members of the same array, username abc on system A is assumed to be the same user as username abc on system B. This is most significant in the case of username root. Authentication should be used if necessary to prevent access to an array by machines using a different namespace.

About the Uses of the Configuration Files

The configuration files are read by the Array Services daemon when it starts. Typically, the daemon starts in each node during the system startup. You can also run the daemon from a command line in order to check the syntax of the configuration files.

The configuration files contain the following data, all of which is needed by ainfo and array:
• The names of array systems, including the current array but also any other arrays on which a user could run an Array Services command. ainfo reports this information.

• The names and types of the nodes in each named array, especially the hostnames that would be used in an Array Services command. ainfo reports this information.

• The authentication keys, if any, that must be used with Array Services commands. The -Kl and -Kr command options use this information. For more information, see "Array Services Commands" on page 91.

• The commands that are valid with the array command.

About Configuration File Format and Contents

A configuration file is a readable text file. The file contains entries of the following four types, which are detailed in later topics.

Array definition       Describes this array and other known arrays, including array names and the node names and types.

Command definition     Specifies the usage and operation of a command that can be invoked through the array command.

Authentication          Specifies authentication numbers that must be used to access the array.

Local option            Options that modify the operation of the other entries or arrayd.

Blank lines, white space, and comment lines beginning with a pound character (#) can be used freely for readability. Entries can be in any order in any of the files read by arrayd.

Besides punctuation, entries are formed with a keyword-based syntax. Keyword recognition is not case-sensitive; however keywords are shown in uppercase in this text and in the man page. The entries are primarily formed from keywords, numbers, and quoted strings, as detailed in the man page arrayd.conf(4).
Loading Configuration Data

The Array Services daemon, `arrayd`, can take one or more filenames as arguments. It reads them all, and treats them like logical continuations. In effect, it concatenates them. If no filenames are specified, it reads `/usr/lib/array/arrayd.conf` and `/usr/lib/array/arrayd.auth`. A different set of files, and any other `arrayd` command-line options, can be written into the file `/etc/config/arrayd.options`, which is read by the startup script that launches `arrayd` at boot time.

Since configuration data can be stored in two or more files, you can combine different strategies, for example:

- One file can have different access permissions than another. Typically, `/usr/lib/array/arrayd.conf` is world-readable and contains the available `array` commands, while `/usr/lib/array/arrayd.auth` is readable only by root and contains authentication codes.

- One node can have different configuration data than another. For example, certain commands might be defined only in certain nodes; or only the nodes used for interactive logins might know the names of all other nodes.

- You can use NFS-mounted configuration files. You could put a small configuration file on each machine to define the array and authentication keys, but you could have a larger file defining `array` commands that is NFS-mounted from one node.

After you modify the configuration files, you can make `arrayd` reload them by killing the daemon and restarting it in each machine. The script `/etc/init.d/array` supports this operation:

To kill daemon, execute this command:

```
/etc/init.d/array stop
```

To kill and restart the daemon in one operation; perform the following command:

```
/etc/init.d/array restart
```

The Array Services daemon in any node knows only the information in the configuration files available in that node. This can be an advantage, in that you can limit the use of particular nodes; but it does require that you take pains to keep common information synchronized. “Designing New Array Commands” on page 108 summarizes an automated way to do this.
About Substitution Syntax

The `arrayd.conf(4)` man page explains the syntax rules for forming entries in the configuration files. An important feature of this syntax is the use of several kinds of text substitution, by which variable text is substituted into entries when they are executed.

Most of the supported substitutions are used in command entries. These substitutions are performed dynamically, each time the `array` command invokes a subcommand. At that time, substitutions insert values that are unique to the invocation of that subcommand. For example, the value `%USER` inserts the user ID of the user who is invoking the `array` command. Such a substitution has no meaning except during execution of a command.

Substitutions in other configuration entries are performed only once, at the time the configuration file is read by `arrayd`. Only environment variable substitution makes sense in these entries. The environment variable values that are substituted are the values inherited by `arrayd` from the script that invokes it, which is `/etc/init.d/array`.

Testing Configuration Changes

The configuration files contain many sections and options. The Array Services command `ascheck` performs a basic sanity check of all configuration files in the array.

After making a change, you can test an individual configuration file for correct syntax by executing `arrayd` as a command with the `-c` and `-f` options. For example, suppose you have just added a new command definition to `/usr/lib/array/arrayd.local`. You can check its syntax with the following command:

```
arrayd -c -f /usr/lib/array/arrayd.local
```

When testing new commands for correct operation, you need to see the warning and error messages produced by `arrayd` and processes that it may spawn. The `stderr` messages from a daemon are not normally visible. You can make them visible by the following procedure:

1. On one node, kill the daemon, as follows:

```
# /etc/init.d/array stop
```

```
2. In one shell window on that node, start `arrayd` with the options `-n` `-v`, as follows:

```
# /usr/sbin/arrayd -n -v
```

Instead of moving into the background, it remains attached to the shell terminal.

**Note:** Although `arrayd` becomes functional in this mode, it does not refer to `/etc/config/arrayd.options`, so you need to specify explicitly all command-line options, such as the names of nonstandard configuration files.

3. From another shell window on the same or other nodes, issue `ainfo` and `array` commands to test the new configuration data. Diagnostic output appears in the `arrayd` shell window.

4. Terminate `arrayd` and use the following command to restart it as a daemon:

```
# /usr/sbin/arrayd -v
```

During steps 1, 2, and 4, the test node might not respond to `ainfo` and `array` commands, so warn users that the array is in test mode.

**Specifying Arrayname and Machine Names**

The following lines are a simple example of an array definition within an `arrayd.conf` file:

```
array simple
  machine congo
  machine niger
  machine nile
```

The array name `simple` is the value the user must specify in the `-a` option. For more information, see "Array Services Commands" on page 91.

One array name should be specified in a `DESTINATION ARRAY` local option as the default array and reported by `ainfo dflt`. Local options are listed under "Configuring Local Options" on page 107.

**Specifying IP Addresses and Ports**

The simple `machine` subentries shown in the example are based on the assumption that the hostname is the same as the machine’s name to domain name services (DNS).
If a machine’s IP address cannot be obtained from the given hostname, provide a hostname subentry to specify either a fully qualified domain name (FQDN) or an IP address, as follows:

```
array simple
  machine congo
    hostname congo.engr.hitech.com
    port 8820
  machine niger
    hostname niger.engr.hitech.com
  machine nile
    hostname "198.206.32.85"
```

The preceding example shows how to use the port subentry to specify that arrayd in a particular machine uses a different socket number than the default of 5434.

### Specifying Additional Attributes

If you want the ainfo command to display certain strings, you can insert these values as subentries to the array entry. The following are some examples of attributes:

```
array simple
  array_attribute config_date="04/03/96"
  machine a_node
    machine_attribute aka="congo"
    hostname congo.engr.hitech.com
```

**Tip:** You can write code that fetches any array name, machine name, or attribute string from any node in the array.

### Configuring Array Commands

The user can invoke arbitrary system commands on single nodes using the arshell command. The user can also launch MPI programs that automatically distribute over multiple nodes. However, the only way to launch coordinated system programs on all nodes at once is to use the array command. This command does not accept any system command; it only permits execution of commands that the administrator has configured into the Array Services database.
You can define any set of commands that your users need. You have complete control over how any single array node executes a command. For example, the definition can be different in different nodes. A command can simply invoke a standard system command, or, since you can define a command as invoking a script, you can make a command arbitrarily complex.

**Operation of Array Commands**

When a user invokes the `array` command, the subcommand and its arguments are processed by the destination node specified by `-s`. Unless the `-l` option was given, that daemon also distributes the subcommand and its arguments to all other array nodes that it knows about. Remember that the destination node might be configured with only a subset of nodes. At each node, `arrayd` searches the configuration database for a `COMMAND` entry with the same name as the `array` subcommand.

In the following example, the subcommand `uptime` is processed by `arrayd` in node `tokyo`:

```
array -s tokyo uptime
```

When `arrayd` finds the subcommand valid, it distributes it to every node that is configured in the default array at node `tokyo`.

The `COMMAND` entry for `uptime` is distributed in this form. You can read it in the file `/usr/lib/array/arrayd.conf`.

```
command uptime # Display uptime/load of all nodes in array
   invoke /usr/lib/array/auptime %LOCAL
```

The `INVOKE` subentry tells `arrayd` how to execute this command. In this case, it executes a shell script `/usr/lib/array/auptime`, passing it one argument, the name of the local node. This command is executed at every node, with `%LOCAL` replaced by that node’s name.

**Summary of Command Definition Syntax**

Look at the basic set of commands distributed with Array Services. This command set resides in `/usr/lib/array/arrayd.conf`. Each `COMMAND` entry is defined using the subentries shown in Table 10-5, which the `arrayd.conf(4)` man page also describes.
<table>
<thead>
<tr>
<th>Keyword</th>
<th>Meaning of Following Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>COMMAND</td>
<td>The name of the command as the user gives it to array.</td>
</tr>
<tr>
<td>INVOKE</td>
<td>A system command to be executed on every node. The argument values can be literals, or arguments given by the user, or other substitution values.</td>
</tr>
<tr>
<td>MERGE</td>
<td>A system command to be executed only on the distributing node, to gather the streams of output from all nodes and combine them into a single stream.</td>
</tr>
<tr>
<td>USER</td>
<td>The user ID under which the INVOKE and MERGE commands run. Usually given as USER %USER, so as to run as the user who invoked array.</td>
</tr>
<tr>
<td>GROUP</td>
<td>The group name under which the INVOKE and MERGE commands run. Usually given as GROUP %GROUP, so as to run in the group of the user who invoked array.</td>
</tr>
<tr>
<td>PROJECT</td>
<td>The project under which the INVOKE and MERGE commands run. Usually given as PROJECT %PROJECT, so as to run in the project of the user who invoked array.</td>
</tr>
<tr>
<td>OPTIONS</td>
<td>A variety of options to modify this command. For more information, see Table 10-7.</td>
</tr>
</tbody>
</table>

The system commands called by INVOKE and MERGE must be specified as full pathnames because arrayd has no defined execution path. As with a shell script, these system commands are often composed from a few literal values and many substitution strings. The substitutions that are supported, all of which are documented in detail in the arrayd.conf(4) man page, are summarized in Table 10-6.
Table 10-6  Substitutions Used in a COMMAND Definition

<table>
<thead>
<tr>
<th>Substitution</th>
<th>Replacement Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>%1..%9; %ARG(n);</td>
<td>Argument tokens from the user’s subcommand. %OPTARG does not produce an error message if the specified argument is omitted.</td>
</tr>
<tr>
<td>%ALLARGS; %OPTARG(n)</td>
<td></td>
</tr>
<tr>
<td>%USER, %GROUP, %PROJECT</td>
<td>The effective user ID, effective group ID, and project of the user who invoked array.</td>
</tr>
<tr>
<td>%REALUSER, %REALGROUP</td>
<td>The real user ID and real group ID of the user who invoked array.</td>
</tr>
<tr>
<td>%ASH</td>
<td>The internal array session handle (ASH) number under which the INVOKE or MERGE command is to run.</td>
</tr>
<tr>
<td>%PID(ash)</td>
<td>List of PID values for a specified ASH. %PID(%ASH) is a common use.</td>
</tr>
<tr>
<td>%ARRAY</td>
<td>The array name, either default or as given in the -a option.</td>
</tr>
<tr>
<td>%LOCAL</td>
<td>The hostname of the executing node.</td>
</tr>
<tr>
<td>%ORIGIN</td>
<td>The full domain name of the node where the array command ran and the output is to be viewed.</td>
</tr>
<tr>
<td>%OUTFILE</td>
<td>List of names of temporary files, each containing the output from one node’s INVOKE command. Valid only in the MERGE subentry.</td>
</tr>
</tbody>
</table>

The OPTIONS subentry permits a number of important modifications of the command execution. Table 10-7 summarizes these.

Table 10-7  Options of the COMMAND Definition

<table>
<thead>
<tr>
<th>Keyword</th>
<th>Effect on Command</th>
</tr>
</thead>
<tbody>
<tr>
<td>LOCAL</td>
<td>Do not distribute to other nodes. Effectively forces the -l option.</td>
</tr>
<tr>
<td>NEWSESSION</td>
<td>Execute the INVOKE command under a newly created ASH. %ASH in the INVOKE line is the new ASH. The MERGE command runs under the original ASH, and %ASH substitutes as the old ASH in that line.</td>
</tr>
</tbody>
</table>
Keyword Effect on Command

SETRUID Set both the real and effective user ID from the USER subentry. Typically, USER only sets the effective UID.

SETRGID Set both the real and effective group ID from the GROUP subentry. Typically, GROUP sets only the effective GID.

QUIET Discard the output of INVOKE, unless a MERGE subentry is given. If a MERGE subentry is given, pass INVOKE output to MERGE as usual, and discard the MERGE output.

NOWAIT Discard the output and return as soon as the processes are invoked. Do not wait for completion. A MERGE subentry is ineffective.

Configuring Local Options

The LOCAL entry specifies options to arrayd itself. The most important options are summarized in Table 10-8.

Table 10-8 Subentries of the LOCAL Entry

<table>
<thead>
<tr>
<th>Subentry</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIR</td>
<td>Pathname for the arrayd working directory, which is the initial, current working directory of INVOKE and MERGE commands. The default is /usr/lib/array.</td>
</tr>
<tr>
<td>DESTINATION ARRAY</td>
<td>Name of the default array, used when the user omits the -a option. When only one ARRAY entry is given, it is the default destination.</td>
</tr>
<tr>
<td>USER, GROUP, PROJECT</td>
<td>Default values for COMMAND execution when USER, GROUP, or PROJECT are omitted from the COMMAND definition.</td>
</tr>
<tr>
<td>HOSTNAME</td>
<td>Value returned in this node by %LOCAL. Default is the hostname.</td>
</tr>
<tr>
<td>PORT</td>
<td>Socket to be used by arrayd.</td>
</tr>
</tbody>
</table>
If you do not supply LOCAL USER, GROUP, and PROJECT values, the default values for USER and GROUP are arraysvcs.

The HOSTNAME entry is needed whenever the hostname command does not return a node name as specified in the ARRAY MACHINE entry. In order to supply a LOCAL HOSTNAME entry unique to each node, each node needs an individualized copy of at least one configuration file.

**Designing New Array Commands**

A basic set of commands is distributed in the file/usr/lib/array/arrayd.conf.template. You should examine this file carefully before defining commands of your own. You can define new commands which then become available to the users of the array system.

Typically, a new command will be defined with an INVOKE subentry that names a script written in sh, csh, or Perl syntax. You use the substitution values to set up arguments to the script. You use the USER, GROUP, PROJECT, and OPTIONS subentries to establish the execution conditions of the script.

Within the invoked script, you can write any amount of logic to verify and validate the arguments and to execute any sequence of commands. For an example of a script in Perl, see /usr/lib/array/aps, which is invoked by the array ps command.

**Note:** Perl is a particularly interesting choice for array commands, since Perl has native support for socket I/O. In principle at least, you could build a distributed application in Perl in which multiple instances are launched by array and coordinate and exchange data using sockets. Performance would not rival the highly tuned MPI libraries, but development would be simpler.

The administrator has need for distributed applications as well, since the configuration files are distributed over the array. Here is an example of a distributed command to reinitialize the Array Services database on all nodes at once. The script to be executed at each node, called /usr/lib/array/arrayd-reinit would read as follows:

```bash
#!/bin/sh

# Script to reinitialize arrayd with a new configuration file
```
# Usage: arrayd-reinit <hostname:new-config-file>
sleep 10        # Let old arrayd finish distributing
rcp $1 /usr/lib/array/
/etc/init.d/array restart
exit 0

The script uses rcp to copy a specified file, presumably a configuration file such as arrayd.conf, into /usr/lib/array. This fails if %USER is not privileged. Then the script restarts arrayd to reread configuration files.

The command definition is as follows:

```
command reinit
#################################################################
# NOTE: The example shown is for illustrative purposes only and #
# has not been evaluated for use in a production environment.  #
#################################################################
   invoke /usr/lib/array/arrayd-reinit %ORIGIN:%1
user  %USER
   group %GROUP
   options nowait   # Exit before restart occurs!
```

The INVOKE subentry calls the restart script shown above. The NOWAIT option prevents the daemon’s waiting for the script to finish because the script kills the daemon.

Caution: The preceding examples are for illustrative purposes only and have not been evaluated for use in a production environment.
Chapter 11

Using the SGI MPT Plugin for Nagios

This chapter includes the following topics:

• "About the SGI MPT Plugin for Nagios" on page 111
• "Installing the SGI MPT Nagios Plugin on the Admin Node" on page 112
• "(Optional) Installing the SGI MPT Nagios Plugin on a Rack Leader Controller (RLC) Node" on page 115
• "Viewing SGI MPT Messages From Within Nagios and Clearing the Messages" on page 116
• "(Optional) Modifying the Notification Email" on page 119

About the SGI MPT Plugin for Nagios

Nagios is a web-based system monitoring tool that SGI automatically installs on SGI ICE cluster computer systems. Nagios enables you to monitor the cluster infrastructure. When you install the optional SGI MPT plugin for Nagios, the SGI MPT system log messages that typically appear in /var/log/messages also appear in the Nagios graphical user interface (GUI). The plugin scans the system log for messages that SGI MPT has logged, and in the Nagios GUI, the plugin displays the number of error messages and warning messages that the plugin encountered in the scan.

The following topics provide more information about the SGI MPT plugin for Nagios:

• "Installing the SGI MPT Nagios Plugin on the Admin Node" on page 112
• "(Optional) Installing the SGI MPT Nagios Plugin on a Rack Leader Controller (RLC) Node" on page 115
• "Viewing SGI MPT Messages From Within Nagios and Clearing the Messages" on page 116
• "(Optional) Modifying the Notification Email" on page 119
Installing the SGI MPT Nagios Plugin on the Admin Node

The following procedure explains how to install the SGI MPT Nagios plugin on the admin node.

Procedure 11-1  To install the SGI MPT Nagios plugin on the admin node

1. Locate the SGI Performance Suite installation DVD, and insert the DVD into the DVD reader on the admin node.

2. Log into the admin node as the root user.

3. Change to the RPM repository directory.

4. Type one of the following commands to install the plugin:

   a. On RHEL 7 systems or RHEL 6 systems, type the following command:
      
      ```
      # yum install checkmpt-plugin
      ```

   b. On SLES 12 systems or SLES 11 systems, type the following commands:
      
      ```
      # zypper in checkmpt-plugin
      ```

      The preceding commands install the following files:
      
      `/opt/sgi/mpt/checkmpt-plugin/README`
      `/opt/sgi/nagios/libexec/check_mpt`

5. Use a text editor to open file `/opt/sgi/mpt/checkmpt-plugin/README`, and leave the file open in a window on your desktop.

   This file contains a shorthand version of these installation instructions. Some steps in this installation procedure require you to insert specific lines into specific files, and it is easiest to copy the lines out of the README file and modify them as this procedure explains.

6. Type the following command to edit file `sudoers`:

   ```
   # visudo
   ```

7. Copy the following lines from the README file to the end of the sudoers file, and replace `<nagiosuser>` and `<PLUGINSDIR>` with values that are valid at your site:

   ```
   # check_mpt plugin for Nagios (needs access to syslogs)
   <nagiosuser> ALL=NOPASSWD: <PLUGINSDIR>/check_mpt
   ```
# end check_mpt

Replace the variables in the preceding lines as follows:

- Replace `<nagiosuser>` with the Nagios username assigned when Nagios was installed. By default, this username is `nagios`.
- Replace `<PLUGINSDIR>` with the directory in which the Nagios plugin resides. By default, this is `/opt/sgi/nagios/libexec`.

8. Save and close the `sudoers` file.

9. Use a text editor to open file `commands.cfg`.

   By default, this file resides in the following directory:
   
   `/opt/sgi/nagios/etc/objects`

9. Copy the following lines from the `README` file to the end of the `commands.cfg` file:

   ```
   # check_mpt command definition
   define command {
       command_name check_mpt
       command_line sudo $USER1$/check_mpt -W $ARG1$ -E $ARG2$
   }
   # end check_mpt
   
   You do not need to assign values to `$ARG1$` or `$ARG2$`. A later step in this procedure populates these arguments with values.
   
11. Save and close the `commands.cfg` file.

12. Use a text editor to open file `localhost.cfg`.

   By default, this file resides in the following directory:
   
   `/opt/sgi/nagios/etc/objects`

13. Copy the following lines from the `README` file to the end of the `localhost.cfg` file:

   ```
   # check_mpt service definition
   define service {
       use
       local-service
       host_name
       localhost
   }
   ```
Using the SGI MPT Plugin for Nagios

```plaintext
service_description  check_mpt
check_command         check_mpt!10!5
max_check_attempts    2
normal_check_interval 2
retry_check_interval  1
}
# end of check_mpt
```

The key lines in the preceding module have the following effects:

<table>
<thead>
<tr>
<th>Line</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>use local-service</td>
<td>Use the generic Nagios template.</td>
</tr>
<tr>
<td>host_name localhost</td>
<td>Run on localhost or similar.</td>
</tr>
<tr>
<td>service_description check_mpt</td>
<td>Declare the service name.</td>
</tr>
<tr>
<td>check_command check_mpt!10!5</td>
<td>Is CRITICAL if &gt;10 warnings / &gt;5 errors.</td>
</tr>
<tr>
<td>max_check_attempts 2</td>
<td>If !OK, try check again.</td>
</tr>
<tr>
<td>normal_check_interval 2</td>
<td>Run check every 2 minutes.</td>
</tr>
<tr>
<td>retry_check_interval 1</td>
<td>Retry every 1 minute.</td>
</tr>
</tbody>
</table>

14. Save and close file `localhost.cfg`.

15. Type the following command to verify the changes you made and to make sure that there are no conflicts:

```
nagios_dir/bin/nagios -v nagiosdir/etc/nagios.cfg
```

For `nagios_dir`, specify the Nagios home directory. By default, this directory is `/opt/sgi/nagios`.

16. Restart Nagios on the node.

This command differs, depending on your platform, as follows:

- To restart Nagios on RHEL 7 and SLES 12 platforms, type the following command:

  ```
  # systemctl restart Nagios
  ```
• To restart Nagios on RHEL 6 and SLES 11 platforms, type the following command:

    # service nagios restart

You need to restart Nagios after you change any of the Nagios .cfg files.

17. On the admin node, use a shell command to set the following environment variable:

    MPI_SYSLOG_COPY=1

    For example:

    # set MPI_SYSLOG_COPY=1

    Make sure to set this value in your shell before you run any SGI MPI or SGI SHMEM applications.

18. (Optional) Leave the DVD in the admin node’s DVD reader, and proceed to the following:

    "(Optional) Installing the SGI MPT Nagios Plugin on a Rack Leader Controller (RLC) Node" on page 115

(Opposional) Installing the SGI MPT Nagios Plugin on a Rack Leader Controller (RLC) Node

In addition to the admin node, you can also install the plugin on one or more RLCs. The installation procedure is very similar to the procedure that explains how to install the plugin on the admin node. After you install the plugin on an RLC, you can start Nagios on that RLC to monitor (1) the messages on that RLC and (2) the messages related to that RLC’s compute nodes.

The following procedure explains how to install the plugin on an RLC.

Procedure 11-2 To install the SGI MPT plugin on an RLC

1. From the admin node, use the ssh command to log into one of the RLCs as the root user.

2. Use the information in the following steps to install the plugin on the RLC:

   • Procedure 11-1, step 4 on page 112
Viewing SGI MPT Messages From Within Nagios and Clearing the Messages

The following procedure explains how to retrieve SGI MPT messages and clear SGI MPT messages.

**Procedure 11-3** To retrieve and clear SGI MPT messages

1. Log into one of the cluster nodes.
   
   If you log into the admin node and start Nagios from the admin node, Nagios displays information for the whole cluster.
   
   If you log into one of the RLCs and start Nagios from one of the RLCs, Nagios displays information for that RLC and its subordinate nodes.

2. Start Nagios.
   
   Type one of the following URLs into your browser:
   
   - To start Nagios on the admin node, type the following:
     
     http://admin_name/nagios
     
     For **admin_name**, type the hostname or IP address of the admin node.
   
   - To start Nagios on one of the RLCs, type the following:
     
     http://admin_name/rlc_name/nagios
     
     For **admin_name**, type the hostname or IP address of the admin node.
     
     For **rlc_name**, type the hostname or IP address of the RLC.

3. Type in the Nagios user’s username and password.
   
   By default, the username is **nagiosadmin**. By default, the password is **sgisgi**.

4. Look for SGI MPT information in the Nagios interface.
   
   By default, the plugin scans the messages in the /var/log/messages and reports messages to Nagios, as follows:
• If you installed the plugin on the admin node, the plugin sends messages to Nagios for the admin node.

• If you installed the plugin on one or more RLCs, the plugin sends messages to Nagios for the RLC and the RLC compute nodes. You need to start Nagios on the RLC to observe the messages related to that RLC.

Figure 11-1 on page 117 shows how an SGI MPT message appears in the Nagios interface.

![Figure 11-1 A Critical SGI MPT Message in Nagios](image)

If you click an SGI MPT message from within the Nagios interface, you retrieve more information about the message. For example, Figure 11-1 on page 117 provides more information about this example.
5. Use administrator commands to remedy the error conditions, if needed.

6. On the admin node, run the `check_mpt` command to clear the messages that Nagios reported.

   If you installed the plugin on the RLCs, run the `check_mpt` on RLCs, too.

   The MPT plugin works by scanning `/var/log/messages`, from beginning to end. To stop the plugin from repeatedly scanning the log file, a file offset is preserved. After you run the `check_mpt` command, the changes appear in Nagios after the next scan.

   The following examples show how to use options to the `check_mpt` command to direct the plugin to scan the system log according to your site preferences.

   Example 1. To direct the plugin to scan for only newly logged messages, use the `-C` option. The `-C` option clears all current message counts and requests that Nagios continue its scan for new messages. Also, the `-C` parameter changes the Nagios CRITICAL and WARNING status back to OK after you correct the reported error condition. To use this option, type the following command:

   ```bash
   # check_mpt -C
   ```
Example 2. The \(-X\) parameter directs the plugin to start a new scan of /var/log/messages, clears the MPT message counts, and resets the offsets to 0. You can run `check_mpt` with the \(-X\) parameter after each log rotation. This command is as follows:

```bash
# check_mpt -X
```

The `check_mpt` command accepts additional parameters. For more information on these parameters, type the following command to retrieve a usage statement:

```bash
# check_mpt -h
```

(Optional) Modifying the Notification Email

In addition to the notifications that Nagios reports in the Nagios GUI, Nagios also sends email notifications of alert conditions. If you modify the Nagios email configuration file, the Nagios email can include hostname information, which can let you identify the node upon which the error condition occurred more easily.

The `commands.cfg` file contains the following:

```bash
# 'notify-service-by-email-long' command definition
define command {
    command_name notify-service-by-email-long
    command_line /usr/bin/printf "%b" "***** Nagios *****
    Notification Type: $NOTIFICATIONTYPE$
    Service: $SERVICEDESC$
    Host: $HOSTALIAS$
    Address: $HOSTADDRESS$
    State: $SERVICESTATE$
    Date/Time: $LONGDATETIME$
    Additional Info: $SERVICEOUTPUT$ | /usr/bin/mail -s "**$NOTIFICATIONTYPE$ Service Alert: $HOSTALIAS$/$SERVICEDESC$ is $SERVICESTATE$ /*" $CONTACTEMAIL$ }
```

If you change `$HOSTALIAS$` to hostname, the Nagios emails include the hostname of the node upon which the error condition occurred. For example, the following file shows this enhancement:

```bash
# 'notify-service-by-email-long' command definition
define command {
    command_name notify-service-by-email-long
    command_line /usr/bin/printf "%b" "***** Nagios *****
    Notification Type: $NOTIFICATIONTYPE$
    Service: $SERVICEDESC$
    Host: 'hostname'
    Address: $HOSTADDRESS$
    State: $SERVICESTATE$
    Date/Time: $LONGDATETIME$
    Additional Info: $SERVICEOUTPUT$ | /usr/bin/mail -s "**$NOTIFICATIONTYPE$ Service Alert: $HOSTALIAS$/$SERVICEDESC$ is $SERVICESTATE$ /*" $CONTACTEMAIL$ }
```
Using the SGI MPT Plugin for Nagios

Info:\n\n$SERVICEOUTPUT$\n\n$LONGSERVICEOUTPUT$" | /usr/bin/mail -s "**$NOTIFICATIONTYPE$ Service Alert: $HOSTALIAS$/$SERVICEDESC$ is $SERVICESTATE$ **" $CONTACTEMAIL$
}

For more information about Nagios and the Nagios email reporting feature, see your Nagios documentation.
Appendix A

Guidelines for Using SGI MPT on a Virtual Machine Within an SGI UV Computer System

This appendix section includes the following topics:

- "About SGI MPT on a Virtual Machine" on page 121
- "Installing Software Within the Virtual Machine (VM)" on page 121
- "Adjusting SGI UV Virtual Machine System Settings" on page 122
- "Running SGI MPI Programs From Within a Virtual Machine (VM)" on page 124

About SGI MPT on a Virtual Machine

You can configure a virtual machine (VM) on an SGI UV system. The VM creates a general-purpose computer, and MPT can run on that computer. When you use SGI MPT from within a VM, however, you can expect differences in the computing environment and differences with regard to your application’s behavior.

For information about how to configure a VM on an SGI system, see the documentation for Red Hat Enterprise Linux (RHEL) or for SLES.

If you are an administrator, use the information in the following topics to configure the VM environment appropriately:

- "Installing Software Within the Virtual Machine (VM)" on page 121
- "Adjusting SGI UV Virtual Machine System Settings" on page 122

If you are an application developer, use the information in the following topic to understand how your program might behave differently when running from within a VM:

- "Running SGI MPI Programs From Within a Virtual Machine (VM)" on page 124

Installing Software Within the Virtual Machine (VM)

The following procedure explains the software that you need to install in the VM in order for MPI programs to run on the VM.
Procedure A-1 To install the software for MPI programs

1. Install and configure the operating system (RHEL or SLES) and the SGI Foundation Software on the SGI UV computer.

   For installation information, see the *SGI UV System Software Installation and Configuration Guide*.

2. Install and configure the VM according to your operating system vendor’s instructions.

   Note that RHEL and SLES do not support InfiniBand technology from within a VM. Other OFED providers support InfiniBand technology from within a VM through single-root I/O virtualization (SR-IOV), but SGI does not support SR-IOV or other alternatives to the distribution-supplied OFED.

3. (Optional) Install the SGI Foundation Software into the VM.

   For installation information, see the *SGI UV System Software Installation and Configuration Guide*.

4. Install the SGI Performance Suite software into the VM.

   For installation information, see the SGI Performance Suite release notes.

5. Install SGI MPT into the VM.

   For installation information, see Chapter 2, "Getting Started" on page 21.

Adjusting SGI UV Virtual Machine System Settings

For best performance, SGI recommends to change certain operating system settings after the software installation is complete.

The following procedure explains how to adjust the number of files that can be open at a given time.

Procedure A-2 To adjust system settings

1. Log into the SGI UV system as the root user.

2. Type `cpumap` command to retrieve the number of cores on the SGI UV computer.
For example:

```
# cpumap
This is an SGI UV
model name : Genuine Intel(R) CPU @ 2.60GHz
Architecture : x86_64
cpu MHz : 2600.072
cache size : 20480 KB (Last Level)

Total Number of Sockets : 16
Total Number of Cores : 128 (8 per socket)
Hyperthreading : ON
Total Number of Physical Processors : 128
Total Number of Logical Processors : 256 (2 per Phys Processor)
```

UV Information
```
HUB Version: UVHub 3.0
Number of Hubs: 16
Number of connected Hubs: 16
Number of connected NUMAlink ports: 128
```

The Total Number of Cores line reveals that there are 128 cores, 8 per socket.

3. Display the contents of the `/etc/sysctl.conf` file.

For example, type the following command:

```
# less /etc/sysctl.conf
...  
fs.file-max = 8204481
...
```

4. (Conditional) Use a text editor to open file `sysctl.conf` and increase the value of the `fs.file-max` parameter in the `/etc/sysctl.conf` file.

Perform this step if the number of cores on your computer is greater than 512 and the `fs.file-max` parameter is set to less than 10,000,000.

For optimum performance within a VM, set the `fs.file-max` parameter that is at least 10000000 on SGI UV systems with 512 cores or more.
Running SGI MPI Programs From Within a Virtual Machine (VM)

The following list explains some of the differences between running an MPI or SHMEM program on native SGI hardware versus running an MPI or SHMEM program from within a VM hosted by an SGI UV system:

- Hardware-dependent features might not exist on a VM.

When you run an MPI program on a VM, the environment detects the virtual nature of the platform and ignores any SGI hardware-specific features. The following hardware features are not available to an application that runs in a VM: NUMAlink, Superpages, the SGI UV timer, the HUB ASIC, hardware performance counters, and global reference units (GRUs). In addition, processor-specific performance diagnostics are limited.

If your application uses hardware technologies that are not specific to SGI systems, you can expect that the VM can honor those non-specific technologies.

- Topology characteristics might be different.

An application that relies on the topology of an SGI system needs to be run on a VM that was configured with topology that mimics the SGI computer system. MPI programs do not automatically use special topology characteristics effectively. If the application requires special heuristics for locality and placement, you need to configure that into the VM.

- XPMEM libraries are beneficial in very large VMs.

SGI has tested XPMEM on VMs. XPMEM loads, and your application can call XPMEM routines successfully. However, XPMEM is useful only on systems with very large memory.

- No InfiniBand support.

The RHEL and SLES operating systems do not support InfiniBand technology in VMs. Consult your system administrator to find out if single-root I/O virtualization (SR-IOV) is configured on the VM.
Appendix B

Configuring Array Services Manually

This appendix contains the following topics:

- "About Configuring Array Services Manually" on page 125
- "Configuring Array Services on Multiple Partitions or Hosts" on page 125

About Configuring Array Services Manually

The SGI MPT configuration procedures explain how to configure Array Services in an automated way on SGI UV partitioned systems and on SGI ICE X systems. The information in this appendix section explains how to configure Array Services in a manual way, which allows you to make customizations if necessary.

Configuring Array Services on Multiple Partitions or Hosts

The following procedure explains how to configure Array Services to run on multiple hosts, such as exist on an SGI UV partitioned system or an SGI ICE X system.

**Procedure B-1** To configure Array Services for multiple hosts

1. Log in as root on one of the hosts you want to include in the array.
   
   You must be logged in as an administrator to perform this procedure.
   
   For example, on an SGI ICE X system, log into one of the service nodes. You can include service nodes and compute nodes in the array.

2. (Optional) Install the MUNGE package from the SGI MPI software distribution.
   
   The optional MUNGE software package enables additional security for Array Services operations.
   
   During MUNGE installation, make sure of the following:
   
   - The MUNGE key that is used is the same across all the nodes in the array.
   
   The MUNGE key resides in `/etc/munge/munge.key`. 
• You configure a good time clock source, such as an NTP server. MUNGE depends on time synchronization across all nodes in the array.

To install MUNGE, use one of the following commands:
• On Red Hat Enterprise Linux platforms: `yum install munge`
• On SUSE Linux Enterprise Server platforms: `zypper install munge`

For more information about how to install MUNGE, see the SGI MPI release notes.


4. Edit the `/usr/lib/array/arrayd.conf` file to list the machines in your cluster.

This file enables you to configure many characteristics of an array services environment. The required specifications are as follows:
• The array name.
• The hostnames of the array participants.
• A default destination array.

For more information about the additional characteristics that you can specify in the `arrayd.conf` file, see the `arrayd.conf(4)` man page.

For an example `arrayd.conf` file, see file `/usr/lib/array/arrayd.conf.template`.

Example 1. The following lines specify an array name (`sgicluster`) and two hostnames. Specify each hostname on its own line. `array` and `machine` are keywords in the file.

```
array sgicluster
  machine host1
  machine host2
```

Example 2. The following line sets a default array name.

```
destination array sgicluster
```

5. Save and close file `/usr/lib/array/arrayd.conf`.

6. Use a text editor to open file `/usr/lib/array/arrayd.auth`.
7. Search for the string AUTHENTICATION NOREMOTE, and insert a # character in column 1 to comment out the line.

8. Enable the security level under which you want Array Services to operate.

   This step specifies the authentication mechanism to use when Array Services messages pass between the Array Services daemons. Possible security levels are NONE, SIMPLE, or MUNGE, as follows:

   • If no authentication is required, remove the # character from column 1 of the AUTHENTICATION NONE line.

   • To enable simple authentication, ensure that there is no # in column 1 of the AUTHENTICATION SIMPLE line. This is the default.

   • To enable authentication through MUNGE, remove the # character from column 1 of the AUTHENTICATION MUNGE line.

   Make sure that MUNGE has been installed, as prescribed earlier in this procedure.

   For information about the authentication methods, see the arrayd.auth(4) man page.


10. (Optional) Reset the default user account or the default array port.

   By default, the Array Services installation and configuration process sets the following defaults in the /usr/lib/array/arrayd.conf configuration file:

   • A default user account of arraysvcs.

   Array Services requires that a user account exist on all hosts in the array for the purpose of running certain Array Services commands. If you create a different account, make sure to update the arrayd.conf file and set the user account permissions correctly on all hosts.

   • A default port number of 5434.

   The /etc/services file contains a line that defines the arrayd service and port number as follows:

   sgi-arrayd 5434/tcp  # SGI Array Services daemon
You can set any value for the port number, but all systems mentioned in the `arrayd.conf` file must use the same value.

11. Type the following command to restart Array Services:

```
/etc/init.d/array restart
```

12. Repeat the preceding steps on the other hosts or copy the `/usr/lib/array/arrayd.conf` and `/usr/lib/array/arrayd.auth` files to the other hosts.

The Array Services feature requires that the configuration files on each participant host include the list of host participants and the authentication method. The files can contain additional, host-specific information.
Index

A

Argument checking, 43
Array Services, 91
array configuration database, 91
array daemon, 91
arrayconfig_tempo command, 15
authentication key, 91
commands, 91
ainfo, 91
array, 91
arshell, 91
common environment variables, 93
concepts
array session, 91
configuring, 15
global process namespace, 87
ibarray, 91
local process management commands, 90
at, 90
batch, 90
intro, 90
kill, 90
nice, 90
ps, 90
top, 90
managing local processes, 89
monitoring processes and system usage, 89
names of arrays and nodes, 91
release notes, 88
scheduling and killing local processes, 89
security considerations, 97
specifying a single node, 93
using an array, 87
using array services commands, 90
arrayconfig_tempo command, 15

B

Berkeley Lab Checkpoint/Restart (BLCR), 51
installation, 51
using with SGI MPT, 52

C

Cache coherent non-uniform memory access (ccNUMA) systems, 25, 67
ccNUMA
See also "cache coherent non-uniform memory access", 25, 67
Checkpoint/restart, 51
Code hangs, 83
Combining MPI with tools, 85
Configuring Array Services, 15
Configuring SGI MPT
adjusting file descriptor limits, 5, 11
OFED, 9

D

Debuggers
idb and gdb, 44

F

Frequently asked questions, 81

G

Getting started, 21
Global reference unit (GRU), 65

I

Internal statistics, 78

M

Memory placement and policies, 57
Memory use size problems, 84
MPI jobs, suspending, 68
MPI launching problems, 85
MPI on SGI UV systems, 65
  general considerations, 66
  job performance types, 66
  other ccNUMA performance issues, 67
MPI performance profiling, 72
  environment variables, 76
  results file, 73
MPI_REQUEST_MAX too small, 83
mpirun command
  to launch application, 27
  mpirun failing, 81

O

OFED configuration for SGI MPT, 9

P

PerfBoost, 47
  environment variables, 47
  MPI supported functions, 48
  using, 47
Perfcatch utility
  results file, 73
  See also "MPI performance profiling", 72
  using, 72

Profiling interface, 77
Profiling MPI applications, 71
  MPI internal statistics, 78
  profiling interface, 77
Programs
  compiling and linking, 21
    GNU compilers, 24
    Intel compiler, 24
    Open 64 compiler with hybrid
      MPI/OpenMP applications, 24
debugging methods, 43
  launching with mpirun, 27
  launching with PBS, 25
  launching with Torque, 26
  SHMEM programming model, 29
  with TotalView, 43

R

Running MPI Jobs with a workload manager, 25

S

SGI MPT software installation, 84
SGI SHMEM applications, 29
SGI UV hub, 65
SHMEM information, 84
Single copy optimization
  avoiding message buffering, 56
  using the XPMEM driver, 56
Stack traceback information, 86
stdout and/or stderr not appearing, 84
System configuration
  Configuring Array Services, 15
  configuring SGI MPT
    adjusting file descriptor limits, 5, 11
T

TotalView, 43
Troubleshooting, 81
Tuning
  avoiding message buffering, 56
  buffer resources, 55
  enabling single copy, 56
  for running applications across multiple hosts, 61
  for running applications over the InfiniBand Interconnect, 63
  memory placement and policies, 57
  MPI/OpenMP hybrid codes, 59
  reducing run-time variability, 54
  using dplace, 59
  using MPI_DSM_CPULIST, 57
  using MPI_DSM_DISTRIBUTE, 58
  using MPI_DSM_VERBOSE, 59
  using the XPMEM driver, 56

U

Unpinning memory, 68
Using PBS Professional to launch application, 25
Using Torque to launch application, 26