New Features

This revision includes the following updates:

- Removed references to the Unified Parallel C (UPC) product.
- Revised chapter 5, Data Placement Tools for SGI UV Computers
- Added information about the *madvise* keyword for transparent huge pages.
- Miscellaneous editorial and technical corrections.
# Record of Revision

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Supports the SGI Performance Suite 1.1 release. |
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Supports the SGI Performance Suite 1.7 release and includes a correction to the PerfSocket installation documentation. |
| 007     | May 2014  
Supports the SGI Performance Suite 1.8 release. |
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About This Guide

This publication provides information about how to tune C and Fortran application programs that you compiled with an Intel compiler on an SGI® UV™ series system that hosts either the Red Hat Enterprise Linux (RHEL) or the SUSE Linux Enterprise Server (SLES) operating system. Some parts of this manual are also applicable to other SGI X86-64 based systems, such as the SGI® ICE™ X and SGI® Rackable™ systems.

This guide is written for experienced programmers who are familiar with Linux commands and with either the C or Fortran programming languages. The focus in this document is on achieving the highest possible performance by exploiting the features of your SGI system. The material assumes that you know the basics of software engineering and that you are familiar with standard methods and data structures. If you are new to programming or software design, this guide will not be of use to you.

Related SGI Publications

The release notes for the SGI Foundation Suite and the SGI Performance Suite list SGI publications that pertain to the specific software packages in those products. The release notes reside in a text file in the /docs directory on the product media. For example, SGI-MPI-1.x-readme.txt. After installation, the release notes and other product documentation reside in the /usr/share/doc/packages/product directory.

All SGI publications are available on the Technical Publications Library at http://docs.sgi.com. The following publications provide information about Linux implementations on SGI systems:

- **SGI UV System Software Installation and Configuration Guide**
  
  Explains how to install the operating system on an SGI UV system. This manual also includes information about basic configuration features such as CPU frequency scaling and partitioning.

- **SGI Cpuset Software Guide**
  
  Explains how to use cpusets within your application program. Cpusets restrict processes within a program to specific processors or memory nodes.

- **Message Passing Toolkit (MPT) User Guide**
Describes the industry-standard message passing protocol optimized for SGI computers. This manual describes how to tune the run-time environment to improve the performance of an MPI message passing application on SGI computers. The tuning methods do not involve application code changes.

- **MPIInside Reference Guide**
  Documents the SGI MPIInside MPI profiling tool.

- SGI hardware documentation.
  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.

  The following procedure explains how to retrieve a list of hardware manuals for your system.

**Procedure 0-1** To retrieve hardware documentation

1. Type the following URL into the address bar of your browser:
   
   docs.sgi.com

2. In the search box on the Techpubs Library, narrow your search as follows:
   
   - In the search field, type the model of your SGI system.
     
     For example, type one of the following: "UV 2000", "ICE X", Rackable.
     
     Remember to enclose hardware model names in quotation marks (" ") if the hardware model name includes a space character.

   - Check **Search only titles**.

   - Check **Show only 1 hit/book**.

   - Click search.
Related Publications From Other Sources

Compilers and performance tool information for software that runs on SGI Linux systems is available from a variety of sources. The following additional documents might be useful to you:

- http://sourceware.org/gdb/documentation/

  GDB: The GNU Project Debugger website with documentation, such as, Debugging with GDB, GDB User Manual, and so on.

- http://www.intel.com/cd/software/products/asmo-na/eng/perflib/219780.htm; documentation for Intel compiler products can be downloaded from this website.

  Intel Software Network page with links to Intel documentation, such as, Intel Professional Edition Compilers, Intel Thread Checker, Intel Vtune Performance Analyzer, and various Intel cluster software solutions.


- Intel provides specific tuning information tutorial for Nehalem (Intel Xeon 5500) at http://software.intel.com/sites/webinar/tuning-your-application-for-nehalem/.

- Intel provides information for Westmere (Intel Xeon 5600) at http://www.intel.com/itcenter/products/xeon/5600/index.htm


  Intel Software Network page with information specific to Intel Vtune Performance Analyzer including links to documentation.


- Information about the OpenMP Standard can be found at http://openmp.org/wp/.

  The OpenMP API specification for parallel programming website is found here.
Obtaining Publications

You can obtain SGI documentation in the following ways:

- You can access the SGI Technical Publications Library at the following website:
  http://docs.sgi.com

  Various formats are available. This library contains the most recent and most
  comprehensive set of online books, release notes, man pages, and other
  information.

- You can view man pages by typing `man title` at a command line.

Conventions

The following conventions are used in this documentation:

- Brackets enclose optional portions of a command or directive line.

- This fixed-space font denotes literal items such as commands, files, routines, path names, signals, messages, and programming language structures.

- Ellipses indicate that a preceding element can be repeated.

- This bold, fixed-space font denotes literal items that the user enters in interactive sessions. (Output is shown in nonbold, fixed-space font.)

- Italic typeface denotes variable entries and words or concepts being defined.

- Man page section identifiers appear in parentheses after man page names.

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:
  
  techpubs@sgi.com

- 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

System Overview

Tuning an application involves making your program run its fastest on the available hardware. The first step is to make your program run as efficiently as possible on a single processor system and then consider ways to use parallel processing.

This chapter provides an overview of concepts involved in working in parallel computing environments.

An Overview of SGI System Architecture

For information about system architecture, see the hardware manuals that are available on the Tech Pubs Library at the following website:

http://docs.sgi.com

The Basics of Memory Management

Virtual memory (VM), also known as virtual addressing, is used to divide a system’s relatively small amount of physical memory among the potentially larger amount of logical processes in a program. It does this by dividing physical memory into pages, and then allocating pages to processes as the pages are needed.

A page is the smallest unit of system memory allocation. Pages are added to a process when either a page fault occurs or an allocation request is issued. Process size is measured in pages and two sizes are associated with every process: the total size and the resident set size (RSS). The number of pages being used in a process and the process size can be determined by using either the ps(1) or the top(1) command.

Swap space is used for temporarily saving parts of a program when there is not enough physical memory. The swap space may be on the system drive, on an optional drive, or allocated to a particular file in a filesystem. To avoid swapping, try not to overburden memory. Lack of adequate memory limits the number and the size of applications that can run simultaneously on the system, and it can limit system performance. Access time to disk is orders of magnitude slower than access to random access memory (RAM). A system that runs out of memory and uses swap to disk while running a program will have its performance seriously affected, as
swapping will become a major bottleneck. Be sure your system is configured with enough memory to run your applications.

Linux is a demand paging operating system, using a least-recently-used paging algorithm. Pages are mapped into physical memory when first referenced, and pages are brought back into memory if swapped out. In a system that uses demand paging, the operating system copies a disk page into physical memory only if an attempt is made to access it, that is, a page fault occurs. A page fault handler algorithm does the necessary action. For more information, see the `mmap(2)` man page.
Chapter 2

The SGI Compiling Environment

This chapter provides an overview of the SGI compiling environment on the SGI family of servers and covers the following topics:

- "Compiler Overview" on page 3
- "Environment Modules" on page 4
- "Library Overview" on page 5
- "About Debugging" on page 19

The remainder of this book provides more detailed examples of the use of the SGI compiling environment elements.

Compiler Overview

You can obtain an Intel Fortran compiler or an Intel C/C++ compiler from Intel Corporation or from SGI. For more information, see one of the following links:


In addition, the GNU Fortran and C compilers are available on SGI systems.

For example, the following is the general format for the Fortran compiler command line:

```
% ifort [options] filename.extension
```

An appropriate filename extension is required for each compiler, according to the programming language used (Fortran, C, C++, or FORTRAN 77).

Some common compiler options are:

- `-o filename`: renames the output to `filename`.
- `-g`: produces additional symbol information for debugging.
- `-O[level]`: invokes the compiler at different optimization levels, from 0 to 3.
The SGI Compiling Environment

- `-l directory_name`: looks for include files in `directory_name`.
- `-c`: compiles without invoking the linker; this options produces an `.o` file only.

Many processors do not handle denormalized arithmetic (for gradual underflow) in hardware. The support of gradual underflow is implementation-dependent. Use the `-ftz` option with the Intel compilers to force the flushing of denormalized results to zero.

Note that frequent gradual underflow arithmetic in a program causes the program to run very slowly, consuming large amounts of system time (this can be determined with the `time` command). In this case, it is best to trace the source of the underflows and fix the code; gradual underflow is often a source of reduced accuracy anyway.

`prctl(1)` allows you to query or control certain process behavior. In a program, `prctl` tracks where floating point errors occur.

Environment Modules

A `module` is a user interface that provides for the dynamic modification of a user’s environment. By loading a module, a user does not have to change environment variables in order to access different versions of the compilers, loaders, libraries and utilities that are installed on the system.

Modules can be used in the SGI compiling environment to customize the environment. If the use of modules is not available on your system, its installation and use is highly recommended.

To view which modules are available on your system, use the following command (for any shell environment):

```
% module avail
```

To load modules into your environment (for any shell), use the following commands:

```
% module load intel-compilers-latest mpt/2.04
```

**Note:** The above commands are for example use only; the actual release numbers may vary depending on the version of the software you are using. See the release notes that are distributed with your system for the pertinent release version numbers.
The module help command provides a list of all arguments accepted, as follows:

```
sys:~> module help
```

Modules Release 3.1.6 (Copyright GNU GPL v2 1991):
Available Commands and Usage:
- `add|load` modulefile [modulefile ...]
- `rm|unload` modulefile [modulefile ...]
- `switch|swap` modulefile modulefile2
- `display|show` modulefile [modulefile ...]
- `avail` [modulefile [modulefile ...]]
- `use [-a|--append]` dir [dir ...]
- `unuse` dir [dir ...]
- update
- purge
- list
- clear
- `help` [modulefile [modulefile ...]]
- `whatis` [modulefile [modulefile ...]]
- `apropos|keyword` string
- `initadd` modulefile [modulefile ...]
- `initprepend` modulefile [modulefile ...]
- `initrm` modulefile [modulefile ...]
- `initswitch` modulefile modulefile2
- `initlist`
- `initclear`

----

For details about using modules, see the `module(1)` man page.

**Library Overview**

*Libraries* are files that contain one or more object (.o) files. Libraries are used to simplify local software development by hiding compilation details. Libraries are sometimes also called *archives.*

The SGI compiling environment contains several types of libraries; an overview about each library is provided in this subsection.
Static Libraries

Static libraries are used when calls to the library components are satisfied at link time by copying text from the library into the executable. To create a static library, use the `ar(1)`, or an archiver command.

To use a static library, include the library name on the compiler’s command line. If the library is not in a standard library directory, be sure to use the `-L` option to specify the directory and the `-l` option to specify the library filename.

To build an application to have all static versions of standard libraries in the application binary, use the `-static` option on the compiler command line.

Dynamic Libraries

Dynamic libraries are linked into the program at run time and when loaded into memory can be accessed by multiple programs. Dynamic libraries are formed by creating a Dynamic Shared Object (DSO).

Use the link editor command (`ld(1)`) to create a dynamic library from a series of object files or to create a DSO from an existing static library.

To use a dynamic library, include the library on the compiler’s command line. If the dynamic library is not in one of the standard library directories, use the `-L path` and `-l library_shortname` compiler options during linking. You must also set the `LD_LIBRARY_PATH` environment variable to the directory where the library is stored before running the executable.

C/C++ Libraries

The following C/C++ libraries are provided with the Intel compiler:

- `libguide.a, libguide.so`: for support of OpenMP-based programs.
- `libsvml.a`: short vector math library
- `libirc.a`: Intel’s support for Profile-Guided Optimizations (PGO) and CPU dispatch
- `libimf.a, libimf.so`: Intel’s math library
- `libcpprts.a, libcpprts.so`: Dinkumware C++ library
• libunwind.a, libunwind.so: Unwinder library
• libcxa.a, libcxa.so: Intel’s runtime support for C++ features

SHMEM Message Passing Libraries

The SHMEM application programming interface is implemented by the libsma library and is part of the Message Passing Toolkit (MPT) product on SGI systems. The SHMEM programming model consists of library routines that provide low-latency, high-bandwidth communication for use in highly parallelized, scalable programs. The routines in the SHMEM application programming interface (API) provide a programming model for exchanging data between cooperating parallel processes. The resulting programs are similar in style to Message Passing Interface (MPI) programs. The SHMEM API can be used either alone or in combination with MPI routines in the same parallel program.

A SHMEM program is SPMD (single program, multiple data) in style. The SHMEM processes, called processing elements or PEs, all start at the same time, and they all run the same program. Usually the PEs perform computation on their own subdomains of the larger problem, and periodically communicate with other PEs to exchange information on which the next computation phase depends.

The SHMEM routines minimize the overhead associated with data transfer requests, maximize bandwidth, and minimize data latency. Data latency is the period of time that starts when a PE initiates a transfer of data and ends when a PE can use the data.

SHMEM routines support remote data transfer through put operations, which transfer data to a different PE, get operations, which transfer data from a different PE, and remote pointers, which allow direct references to data objects owned by another PE. Other operations supported are collective broadcast and reduction, barrier synchronization, and atomic memory operations. An atomic memory operation is an atomic read-and-update operation, such as a fetch-and-increment, on a remote or local data object.

For details about using the SHMEM routines, see the intro_shmem(3) man page or the Message Passing Toolkit (MPT) User’s Guide.
Tuning an application involves determining the source of performance problems and then rectifying those problems to make your programs run their fastest on the available hardware. Performance gains usually fall into one of three categories of measured time:

- **User CPU time**: time accumulated by a user process when it is attached to a CPU and is executing.
- **Elapsed (wall-clock) time**: the amount of time that passes between the start and the termination of a process.
- **System time**: the amount of time performing kernel functions like system calls, `sched_yield`, for example, or floating point errors.

Any application tuning process involves the following steps:

1. Analyzing and identifying a problem
2. Locating where in the code the problem is
3. Applying an optimization technique

This chapter describes the process of analyzing your code to determine performance bottlenecks. See Chapter 6, "Performance Tuning" on page 61, for details about tuning your application for a single processor system and then tuning it for parallel processing.

**Determining System Configuration**

One of the first steps in application tuning is to determine the details of the system that you are running. Depending on your system configuration, different options might or might not provide good results.

The `topology(1)` command displays general information about SGI systems, with a focus on node information. This can include node counts for blades, node IDs, NASIDs, memory per node, system serial number, partition number, UV Hub versions, CPU to node mappings, and general CPU information. The `topology` command is installed by the `pcp-sgi` RPM package.
The following is example output from two `topology(1)` commands:

```
uv-sys:~ # topology
System type: UV2000
System name: harp34-sys
Serial number: UV2-00000034
Partition number: 0
  8 Blades
  256 CPUs
  16 Nodes
235.82 GB Memory Total
15.00 GB Max Memory on any Node
  1 BASE I/O Riser
  2 Network Controllers
  2 Storage Controllers
  2 USB Controllers
  1 VGA GPU

uv-sys:~ # topology --summary --nodes --cpus
System type: UV2000
System name: harp34-sys
Serial number: UV2-00000034
Partition number: 0
  8 Blades
  256 CPUs
  16 Nodes
235.82 GB Memory Total
15.00 GB Max Memory on any Node
  1 BASE I/O Riser
  2 Network Controllers
  2 Storage Controllers
  2 USB Controllers
  1 VGA GPU

<table>
<thead>
<tr>
<th>Index</th>
<th>ID</th>
<th>NASID</th>
<th>CPUS</th>
<th>Memory</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>r001i11b00h0</td>
<td>0</td>
<td>16</td>
<td>15316 MB</td>
</tr>
<tr>
<td>1</td>
<td>r001i11b00h1</td>
<td>2</td>
<td>16</td>
<td>15344 MB</td>
</tr>
<tr>
<td>2</td>
<td>r001i11b01h0</td>
<td>4</td>
<td>16</td>
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<tr>
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<td>r001i11b01h1</td>
<td>6</td>
<td>16</td>
<td>15344 MB</td>
</tr>
<tr>
<td>4</td>
<td>r001i11b02h0</td>
<td>8</td>
<td>16</td>
<td>15344 MB</td>
</tr>
</tbody>
</table>
```
The cpumap(1) command displays logical CPUs and shows relationships between them in a human-readable format. Aspects displayed include hyperthread relationships, last level cache sharing, and topological placement. The cpumap command gets its information from /proc/cpuinfo, the /sys/devices/system directory structure, and /proc/sgi_uv/topology. When creating cpusets, the Socket numbers reported in the output section Processor Numbering on Socket(s) corresponds to the mems argument you would use in the definition of a cpuset. The cpuset mems argument is the list of memory nodes that tasks in the cpuset are allowed to use. For more information, see the SGI Cpuset Software Guide available at http://docs.sgi.com. The following is example output:

uv# cpumap
Thu Sep 19 10:17:21 CDT 2013
harp34-sys.americas.sgi.com
This is an SGI UV

model name : Genuine Intel(R) CPU @ 2.60GHz
Architecture : x86_64
cpu MHz : 2599.946
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

Hub-Processor Mapping

<table>
<thead>
<tr>
<th>Hub Location</th>
<th>Processor Numbers -- HyperThreads in ()</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 r001i11b00h0</td>
<td>0 1 2 3 4 5 6 7</td>
</tr>
<tr>
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    ( 208 209 210 211 212 213 214 215 )
11 r001i11b05h1  88 89 90 91 92 93 94 95
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15 r001i11b07h1 120 121 122 123 124 125 126 127
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Processor Numbering on Node(s)

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</tr>
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## 3: Performance Analysis and Debugging

### Sharing of Last Level (3) Caches

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### HyperThreading

#### Shared Processors

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<td>(49, 177)</td>
<td>(50, 178)</td>
<td>(51, 179)</td>
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The `x86info(1)` command displays x86 CPU diagnostics information. Type one of the following commands to load the `x86info(1)` command if the command is not already installed:

- On Red Hat Enterprise Linux (RHEL) systems, type the following:

  ```
  # yum install x86info.x86_64
  ```

- On SUSE Linux Enterprise Server (SLES) systems, type the following:

  ```
  # zypper install x86info
  ```

The following is an example of `x86info(1)` command output:

```
uv44-sys:~ # x86info
x86info v1.25. Dave Jones 2001-2009
Feedback to.

Found 64 CPUs
```

```
-----------------------------------------------
CPU #1
EFamily: 0 EModel: 2 Family: 6 Model: 46 Stepping: 6
CPU Model: Unknown model.
```
3: Performance Analysis and Debugging

Processor name string: Intel(R) Xeon(R) CPU E7520 @ 1.87GHz
Type: 0 (Original OEM)  Brand: 0 (Unsupported)
Number of cores per physical package=16
Number of logical processors per socket=32
Number of logical processors per core=2
APIC ID: 0x0  Package: 0  Core: 0  SMT ID 0
--------------------------------------------------------------------------

CPU #2
EFamily: 0  EModel: 2  Family: 6  Model: 46  Stepping: 6
CPU Model: Unknown model.
Processor name string: Intel(R) Xeon(R) CPU E7520 @ 1.87GHz
Type: 0 (Original OEM)  Brand: 0 (Unsupported)
Number of cores per physical package=16
Number of logical processors per socket=32
Number of logical processors per core=2
APIC ID: 0x6  Package: 0  Core: 0  SMT ID 6
--------------------------------------------------------------------------

CPU #3
EFamily: 0  EModel: 2  Family: 6  Model: 46  Stepping: 6
CPU Model: Unknown model.
Processor name string: Intel(R) Xeon(R) CPU E7520 @ 1.87GHz
Type: 0 (Original OEM)  Brand: 0 (Unsupported)
Number of cores per physical package=16
Number of logical processors per socket=32
Number of logical processors per core=2
APIC ID: 0x10  Package: 0  Core: 0  SMT ID 16
--------------------------------------------------------------------------

...  

You can also use the `uname` command, which returns the kernel version and other machine information. For example:

```
uv44-sys:~ # uname -a
Linux uv44-sys 2.6.32.13-0.4.1.1559.0.PTF-default #1 SMP 2010-06-15 12:47:25 +0200 x86_64 x86_64 x86_64
```

For more system information, change directory to the
`/sys/devices/system/node/node0/cpu0/cache` directory and list the contents.
For example:

```
v44-sys:/sys/devices/system/node/node0/cpu0/cache # ls
index0  index1  index2  index3
```
Change directory to `index0` and list the contents, as follows:

```bash
uv44-sys:/sys/devices/system/node/node0/cpu0/cache/index0 # ls
```

**Sources of Performance Problems**

There are usually three areas of program execution that can have performance slowdowns:

- **CPU-bound processes**: processes that are performing slow operations (such as `sqrt` or floating-point divides) or non-pipelined operations such as switching between add and multiply operations.

- **Memory-bound processes**: code which uses poor memory strides, occurrences of page thrashing or cache misses, or poor data placement in NUMA systems.

- **I/O-bound processes**: processes which are waiting on synchronous I/O, formatted I/O, or when there is library or system level buffering.

Several profiling tools can help pinpoint where performance slowdowns are occurring. The following sections describe some of these tools.

**Profiling with perf**

Linux Performance Events provides a performance analysis framework for systems that use Intel Xeon Phi technology. It includes hardware-level CPU performance monitoring unit (PMU) features, software counters, and tracepoints.

Before you use these profiling tools, make sure the `perf` RPM is installed. The `perf` RPM comes with the your operating system and is not an SGI product.

For more information, see the following man pages: `perf(1)`, `perf-stat(1)`, `perf-top(1)`, `perf-record(1)`, `perf-report(1)`, `perf-list(1)`. The `perf` RPM includes these man pages.

**Profiling with PerfSuite**

PerfSuite is a set of tools, utilities, and libraries that you can use to analyze application software performance Linux-based systems. You can use PerfSuite tools to
perform performance-related activities, ranging from assistance with compiler optimization reports to hardware performance counting, profiling, and MPI usage summarization. PerfSuite is Open Source software. It is approved for licensing under the University of Illinois/NCSA Open Source License (OSI-approved).

For more information, see one of the following websites:

- http://perfsuite.ncsa.uiuc.edu/
- http://perfsuite.sourceforge.net/
- http://www.ncsa.illinois.edu/UserInfo/Resources/Software/Tools/PerfSuite/, which hosts NCSA-specific information about using PerfSuite tools

The `psrun` utility is a PerfSuite command line utility that gathers hardware performance information on an unmodified executable. For more information, see http://perfsuite.ncsa.uiuc.edu/psrun/.

**Other Performance Analysis Tools**

The following tools might be useful to you when you try to optimize your code:

- **The Intel® VTune™ Amplifier XE**, which is a performance and thread profiler. This tool does remote sampling experiments. The VTune data collector runs on the Linux system and an accompanying GUI runs on an IA-32 Windows machine, which is used for analyzing the results. VTune allows you to perform interactive experiments while connected to the host through its GUI. An additional tool, the Performance Tuning Utility (PTU), requires the Intel VTune license.

  For information about Intel VTune Amplifier XE, see the following URL:

- **Intel Inspector XE**, which is a memory and thread debugger. For information about Intel Inspector XE, see the following:


- **Intel Advisor XE**, which is a threading design and prototyping tool. For information about Intel Advisor XE, see the following:

About Debugging

Several debuggers are available on SGI platforms. Information in the following list explains how to access the debuggers:

- Intel® Debugger for Linux, the Intel symbolic debugger. This debugger is based on the Eclipse graphical user interface (GUI). You can run the Intel Debugger for Linux from the `idb` command. This debugger works with the Intel® C and C++ compilers, the Intel® Fortran90 and FORTRAN 77 compilers, and the GNU compilers. This product is available if your system is licensed for the Intel compilers.

  To run the debugger in command line mode, start the debugger with the `idbc` command.

  You can use the Intel Debugger for Linux with both single-threaded applications, multithreaded applications, serial code, and parallel code.

  If you specify the `-gdb` option on the `idb` command, the shell command line provides user commands and debugger output similar to the GNU debugger.

  For more information, see the following:


- GDB, the GNU debugger. The GDB debugger supports C, C++, Fortran, and Modula-2 programs. Use the `gdb` command to start GDB. To use GDB through a GUI, use the `ddd` command.

  When compiling with C and C++, include the `-g` option on the compiler command line. The `-g` option produces the `dwarf2` symbols database that GDB uses.

  When using GDB for Fortran debugging, include the `-g` and `-O0` options. Do not use `gdb` for Fortran debugging when compiling with `-O1` or higher. The standard GDB debugger does not support Fortran 95 programs. To debug Fortran 95 programs, download and install the `gdbf95` patch from the following website:


  To verify that you have the correct version of GDB installed, use the `gdb -v` command. The output should appear similar to the following:

  GNU gdb 5.1.1 FORTRAN95-20020628 (RC1)
  Copyright 2012 Free Software Foundation, Inc.
For a complete list of GDB commands, use the `help` option or see the following user guide:

http://sources.redhat.com/gdb/onlinedocs/gdb_toc.html

Note that the current instances of GDB do not report `ar.ec` registers correctly. If you are debugging rotating, register-based, software-pipelined loops at the assembly code level, try using the Intel Debugger for Linux.

- TotalView, which is a licensed graphical debugger that you can use with MPI programs. For information about TotalView, see the following:
  
  http://www.roguewave.com

In addition to the preceding debuggers, you can start the Intel Debugger and GDB with the `ddd` command. The `ddd` command starts the Data Display Debugger, a GNU product that provides a graphical debugging interface.

The following topics provide more information:

- "Using the Intel Debugger" on page 20
- "Using TotalView" on page 21
- "Using the Data Display Debugger" on page 22

**Using the Intel Debugger**

The Intel Debugger `idb` is part of Intel® Composer XE. You are asked during the installation if you want to install it or not. The `idb` command starts the graphical user interface (GUI). The `idbc` command starts the command line interface.

The following figure shows the GUI:
Figure 3-1 Intel® Debugger GUI

Using TotalView

Figure 3-2 on page 22 shows a TotalView session.
Using the Data Display Debugger

The Data Display Debugger provides a graphical debugging interface to an arbitrary command line debugger. You use the `ddd` command to start this interface. Specify the `--debugger` option to specify the debugger you want to use. For example, specify `--debugger "idb"` to specify the Intel Debugger. The default debugger is `gdb`.

When the debugger loads, the Data Display Debugger screen appears divided into panes that show the following information:

Figure 3-2 TotalView Session
• Array inspection
• Source code
• Disassembled code
• A command line window to the debugger engine

These panes can be switched on and off from the View menu.

Some commonly used commands can be found on the menus. In addition, the following actions can be useful:

• Select an address in the assembly view, click the right mouse button, and select lookup. The \texttt{gdb} command is executed in the command pane and it shows the corresponding source line.

• Select a variable in the source pane and click the right mouse button. The current value is displayed. Arrays are displayed in the array inspection window. You can print these arrays to PostScript by using the Menu>Print Graph option.

• You can view the contents of the register file, including general, floating-point, NaT, predicate, and application registers by selecting Registers from the Status menu. The Status menu also allows you to view stack traces or to switch OpenMP threads.
This chapter describes several tools that you can use to monitor system performance. The tools are divided into two general categories: system monitoring tools and nonuniform memory access (NUMA) tools.

System monitoring tools include the `topology(1)`, `top(1)` commands and the Performance Co-Pilot `pmchart(1)` command and other operating system commands such as the `vmstat(1)`, `iostat(1)` command and the `sar(1)` commands that can help you determine where system resources are being spent.

The `gtopology(1)` command displays a 3D scene of the system interconnect using the output from the `topology(1)` command.

**System Monitoring Tools**

You can use system utilities to better understand the usage and limits of your system. These utilities allow you to observe both overall system performance and single-performance execution characteristics. This section covers the following topics:

- "Hardware Inventory and Usage Commands" on page 25
- "Performance Co-Pilot Monitoring Tools" on page 29
- "System Usage Commands" on page 32
- "Memory Statistics and `nodeinfo` Command" on page 36

**Hardware Inventory and Usage Commands**

This section describes hardware inventory and usage commands and covers the following topics:

- "`topology(1)` Command" on page 25
- "`gtopology(1)` Command" on page 26

**`topology(1)` Command**

The `topology(1)` command provides topology information about your system.
Applications programmers can use the `topology` command to help optimize execution layout for their applications. For more information, see the `topology(1)` man page.

For an example of the `topology` command’s output, see "Determining System Configuration" on page 9.

**gtopology(1) Command**

The `gtopology(1)` command is included as part of the `sgi-pcp` package of the SGI Accelerate, part of SGI Performance Suite software. It displays a 3D scene of the system interconnect using the output from the `topology(1)` command. See the man page for more details.

Figure 4-1 on page 27, shows the ring topology (the eight nodes are shown in pink, the NUMAlink connections in cyan) of an SGI system with 16 CPUs.
Figure 4-1 Ring Topology of an System with 16 CPUs

Figure 4-2 on page 28, shows the fat-tree topology of an SGI system with 32 CPUs. Again, nodes are the pink cubes. Routers are shown as blue spheres (if all ports are used) otherwise, yellow.
Figure 4-2 An SGI System with 32 CPUs Fat-tree Topology

Figure 4-3 on page 29, shows an SGI system with 512 CPUs. The dual planes of the fat-tree topology are clearly visible.
Performance Co-Pilot Monitoring Tools

This section describes Performance Co-Pilot monitoring tools and covers the following topics:

- "hubstats(1) Command" on page 30
- "linkstat-uv(1) Command" on page 30
- "Other Performance Co-Pilot Monitoring Tools" on page 30
hubstats(1) Command

The hubstats(1) command monitors NUMAlink traffic, directory cache operations, and global reference unit (GRU) traffic statistics on SGI UV systems. It does not work on any other platform. For more information, see the hubstats(1) man page.

linkstat-uv(1) Command

The linkstat-uv(1) command monitors NUMAlink traffic and error rates on SGI UV systems. It does not work on any other platform. This command returns information about packets and Mbytes sent/received on each NUMAlink in the system, as well as error rates. It is useful as a performance monitoring tool and as a tool for helping you to diagnose and identify faulty hardware. For more information, see the linkstat-uv(1) man page.

Note that this command is specific to SGI UV systems and does not return the same information as the linkstat(1) command.

Other Performance Co-Pilot Monitoring Tools

In addition to the UV specific tools described above, the pcp and pcp-sgi packages also provide numerous other performance monitoring tools, both graphical and text-based. It is important to remember that all of the performance metrics displayed by any of the tools described in this chapter can also be monitored with other tools such as pmchart(1), pmval(1), pminfo(1) and others. Additionally, the pmlogger(1) command can be used to capture Performance Co-Pilot archives, which can then be "replayed" during a retrospective performance analysis.

A very brief description of other Performance Co-Pilot monitoring tools follows. See the associated man page for each tool for more details.

- **pmchart(1)** — graphical stripchart tool, chiefly used for investigative performance analysis.

- **pmgsys(1)** — graphical tool showing miniature CPU, Disk, Network, LoadAvg and memory/swap in a miniature display, for example, useful for permanent residence on your desktop for the servers you care about.

- **pmgcluster(1)** — pmgsys, but for multiple hosts and thus useful for monitoring a cluster of hosts or servers.

- **clustervis(1)** — 3D display showing per-CPU and per-Network performance for multiple hosts.
- **nfsvis(1)** – 3D display showing NFS client/server traffic, grouped by NFS operation type.
- **nodevis(1)** – 3D display showing per-node CPU and memory usage.
- **webvis(1)** – 3D display showing per-httpd traffic.
- **dkvis(1)** – 3D display showing per-disk traffic, grouped by controller.
- **diskstat(1)** – command line tool for monitoring disk traffic.
- **topdisk(1)** – command line, curses-based tool, for monitoring disk traffic.
- **topsys(1)** – command line, curses-based tool, for monitoring processes making a large numbers of system calls or spending a large percentage of their execution time in system mode using assorted system time measures.
- **pmgxvm(1)** – miniature graphical display showing XVM volume topology and performance statistics.
- **osvis(1)** – 3D display showing assorted kernel and system statistics.
- **pmdumptext(1)** – command line tool for monitoring multiple performance metrics with a highly configurable output format. Therefore, it is a useful tools for scripted monitoring tasks.
- **pmval(1)** – command line tool, similar to **pmdumptext(1)**, but less flexible.
- **pminfo(1)** – command line tool, useful for printing raw performance metric values and associated help text.
- **pmprobe(1)** – command line tool useful for scripted monitoring tasks.
- **pmie(1)** – a performance monitoring inference engine. This is a command line tool with an extraordinarily powerful underlying language. It can also be used as a system service for monitoring and reporting on all sorts of performance issues of interest.
- **pmieconf(1)** – command line tool for creating and customizing "canned" **pmie(1)** configurations.
- **pmlogger(1)** – command line tool for capturing Performance Co-Pilot performance metrics archives for replay with other tools.
- **pmlogger_daily(1)** and **pmlogger_check(1)** – cron driven infrastructure for automated logging with **pmlogger(1)**.
4: Monitoring Tools

- pmcd(1) – the Performance Co-Pilot metrics collector daemon
- PCPIntro(1) - introduction to Performance Co-Pilot monitoring tools, generic command line usage and environment variables
- PMAPI(3) - introduction to the Performance Co-Pilot API libraries for developing new performance monitoring tools
- PMDA(3) - introduction to the Performance Co-Pilot Metrics Domain Agent API, for developing new Performance Co-Pilot agents
- topology(1) – displays general information about SGI UV systems, with a focus on node information. This includes counts of the CPUs, nodes, routers and memory, as well as, various I/O devices. More detailed information is available for node IDs, NASIDs, memory per node, system serial number, partition number, UV Hub versions, CPU to node mappings, I/O device descriptions, and general CPU and I/O information.

System Usage Commands

The following topics show several commands that can be used to determine user load, system usage, and active processes:

- "Using the **w** command" on page 32
- "Using the **ps**(1) Command" on page 33
- "Using the **top**(1) Command" on page 34
- "Using the **vmstat**(8) Command" on page 34
- "Using the **iostat**(1) command" on page 34
- "Using the **sar**(1) command" on page 35

Using the **w** command

To obtain a high-level view of system usage that includes information about who is logged into the system, use the **w**(1) command, as follows:

```
uv44-sys:~ # w
15:47:48 up 2:49, 5 users, load average: 0.04, 0.27, 0.42
USER  TTY      LOGIN@   IDLE   JCPU   PCPU WHAT
root  pts/0    13:10   1:41m   0.07s  0.07s -bash
```
The `w` command’s output shows who is on the system, the duration of user sessions, processor usage by user, and currently executing user commands. The output consists of two parts:

- The first output line shows the current time, the length of time the system has been up, the number of users on the system, and the average number of jobs in the run queue in the last one, five, and 15 minutes.

The rest of the output from the `w` command shows who is logged into the system, the duration of each user session, processor usage by user, and each user’s current process command line.

### Using the `ps(1)` Command

To determine active processes, use the `ps(1)` command, which displays a snapshot of the process table.

The `ps -A r` command example that follows returns all the processes currently running on a system:

```
[user@profit user]# ps -A r
PID TTY STAT TIME COMMAND
211116 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211117 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211118 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211119 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211120 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211121 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211122 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211123 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211124 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211125 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211126 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211127 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211128 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211129 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211130 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
211131 pts/0 R+ 4:08 /usr/diags/bin/olconf RunTIME=5
```
Using the `top(1)` Command

To monitor running processes, use the `top(1)` command. This command displays a sorted list of top CPU utilization processes.

Using the `vmstat(8)` Command

The `vmstat(8)` command reports virtual memory statistics. It reports information about processes, memory, paging, block IO, traps, and CPU activity. For more information, see the `vmstat(8)` man page.

In the following `vmstat(8)` command, the `10` specifies a 10–second delay between updates.

```
uv44-sys:~ # vmstat 10
procs -----------memory---------- ---swap-- -----io---- --system-- -----cpu-----
r  b  swpd free  buff  cache  si  so  bi  bo  in  cs  us  sy  id  wa  st
2  0   0 235984032 418748 8649568 0  0  0  0  0  0  100  0  0
1  0   0 236054400 418748 8645216 0  0  0  4809 256729 3401  0  100  0  0
1  0   0 236188016 418748 8649904 0  0  0  448  256200  631  0  100  0  0
2  0   0 236202976 418748 8645104 0  0  0  341  256201 1117  0  100  0  0
1  0   0 236088720 418748 8649360 0  0  0  847  257104 6152  0  100  0  0
1  0   0 235990944 418748 8648460 0  0  0  240  257085  5960  0  100  0  0
1  0   0 236049568 418748 8645100 0  0  0  4849  256749 3604  0  100  0  0
```

Without the `delay` parameter, which is `10` in this example, the output returns averages since the last reboot. Additional reports give information on a sampling period of length `delay`. The process and memory reports are instantaneous in either case.

Using the `iostat(1)` command

The `iostat(1)` command monitors system input/output device loading by observing the time the devices are active, relative to their average transfer rates. You can use information from the `iostat` command to change system configuration information to better balance the input/output load between physical disks. For more information, see the `iostat(1)` man page.
In the following `iostat(1)` command, the 10 specifies a 10–second interval between updates:

```
# iostat 10
```

```
Linux 2.6.32-430.el6.x86_64 (harp34-sys) 02/21/2014 _x86_64_ (256 CPU)

avg-cpu: %user %nice %system %iowait %steal %idle
        46.24  0.01  0.67  0.01  0.00  53.08

Device: tps Blk_read/s Blk_wrtn/s Blk_read Blk_wrtn
sda  53.66  23711.65  23791.93  21795308343  21869098736
sdb  0.01   0.02     0.00      17795          0

avg-cpu: %user %nice %system %iowait %steal %idle
        99.96  0.00  0.04  0.00  0.00   0.00

Device: tps Blk_read/s Blk_wrtn/s Blk_read Blk_wrtn
sda  321.20  149312.00  150423.20    1493120     1504232
sdb  0.00   0.00     0.00       0           0

avg-cpu: %user %nice %system %iowait %steal %idle
        99.95  0.00  0.05  0.00  0.00   0.00

Device: tps Blk_read/s Blk_wrtn/s Blk_read Blk_wrtn
sda  305.19  146746.05  148453.95    1468928     1486024
sdb  0.00   0.00     0.00       0           0

```

Using the `sar(1)` command

The `sar(1)` command returns the content of selected, cumulative activity counters in the operating system. Based on the values in the `count` and `interval` parameters, the command writes information `count` times spaced at the specified `interval`, which is in seconds. For more information, see the `sar(1)` man page. The following example shows the `sar(1)` command with a request for information about CPU 1, a count of 10, and an interval of 10:

```
uv44-sys:~ # sar -P 1 10 10
Linux 2.6.32-416.el6.x86_64 (harp34-sys) 09/19/2013 _x86_64_ (256 CPU)

11:24:54 AM CPU %user %nice %system %iowait %steal %idle
11:25:04 AM 1  0.20  0.00  0.10  0.00  0.00   99.70
```

007–5646–007 35
4: Monitoring Tools

nodeinfo(1) is a tool for monitoring per-node NUMA memory statistics on SGI UV systems. The nodeinfo tool reads /sys/devices/system/node/*/meminfo and /sys/devices/system/node/*/numastat on the local system to gather NUMA memory statistics.

Sample memory statistic from the nodeinfo(1) command are, as follows:

```text
uv44-sys:~ # nodeinfo
Memory Statistics Tue Oct 26 12:01:58 2010

node          Total      Free      Used    Dirty  Anon   Slab    hit  miss  foreign    interlv  local  remote
0 16757488 16277084  480404   52 34284  36288  20724   0   0   0   20720  4
1 16777216 16433988  343228   68 6772  17708  4477    0   0   0   3381 1096
2 16777216 16438568  338648   76 6908  12620  1804    0   0   0   708 1093
3 16760832 16429844  330988  56 2820  16836  1804    0   0   0   708 1094
4 16777216 16444408  332808  88 10124  13588  1517    0   0   0   417 1100
5 16760832 16430300  330532  72 1956  17304  4546    0   0   0   3453 1093
6 16777216 16430788  346428  36 3236  15292  3961    0   0   0   2864 1097
7 16760832 16435532  325300  44 1220  14800  3917    0   0   0   2877 1094
TOT 13414888 131320512 2828336 492 67320 144436 42802    0   0   0  35129 7673
Press "h" for help
```

From an interactive nodeinfo session, enter h for a help statement:

Display memory statistics by node.
q quit
+ Increase starting node number. Used only if more nodes than will fit in the current window.

11:25:14 AM 1 10.10 0.00 0.30 0.00 0.00 89.60
11:25:24 AM 1 99.70 0.00 0.30 0.00 0.00 0.00
11:25:34 AM 1 99.70 0.00 0.30 0.00 0.00 0.00
11:25:44 AM 1 8.99 0.00 0.60 0.00 0.00 90.41
11:25:54 AM 1 0.10 0.00 0.20 0.00 0.00 99.70
11:26:04 AM 1 38.70 0.00 0.10 0.00 0.00 61.20
11:26:14 AM 1 99.80 0.00 0.10 0.00 0.00 0.10
11:26:24 AM 1 80.42 0.00 0.70 0.00 0.00 18.88
11:26:34 AM 1 0.10 0.00 0.20 0.00 0.00 99.70
Average: 1 43.78 0.00 0.29 0.00 0.00 55.93
- Decrease starting node number. Used only if more nodes than will fit in the current window.
b Start output with node 0.
e Show highest node number.
k show sizes in KB.
m show sizes in MB.
p show sizes in pages.
t Change refresh rate.
A Show/Hide memory policy stats.
H Show/Hide hugepage info.
L Show/Hide LRU Queue stats.

Field definitions:

hit - page was allocated on the preferred node
miss - preferred node was full. Allocation occurred on THIS node by a process running on another node that was full
foreign - Preferred node was full. Had to allocate somewhere else.

interlv - allocation was for interleaved policy

local - page allocated on THIS node by a process running on THIS node
remote - page allocated on THIS node by a process running on ANOTHER node

(press any key to exit from help screen)
Chapter 5

Data Process and Placement Tools

This chapter contains the following topics:

- “About Nonuniform Memory Access (NUMA) Computers” on page 39
- “About the Data and Process Placement Tools” on page 41

About Nonuniform Memory Access (NUMA) Computers

On symmetric multiprocessor (SMP) computers, all data is visible from all processors. Each processor is functionally identical and has equal time access to every memory address. That is, all processors have equally fast (symmetric) access to memory. These types of systems are easy to assemble and have limited scalability due to memory access times.

NUMA computers also have a shared address space. In both cases, there is a single shared memory space and a single operating system instance. However, in an SMP computer, each processor is functionally identical and has equal time access to every memory address.

In contrast, a NUMA system has a shared address space, but the access time to memory varies over physical address ranges and between processing elements. The Intel Xeon 7500 series processor (i7 architecture) is an example of NUMA architecture. Each processor has its own memory and can address the memory attached to another processor through the Quick Path Interconnect (QPI).

The SGI UV 1000 series is a family of multiprocessor, distributed shared memory (DSM) computer systems that initially scale from 32 to 2560 Intel processor cores, as a cache-coherent single system image (SSI). The SGI UV 100 series is a family of multiprocessor distributed shared memory (DSM) computer systems that initially scale from 16 to 768 Intel processor cores as a cache-coherent SSI. The SGI UV 2000 series scales from 32 to 4,096 Intel processor cores as a cache-coherent SSI.

There are two levels of NUMA: intranode, managed by the Intel QPI, and internode, managed through the SGI HUB ASIC and SGI NUMAlink 5 or SGI NUMAlink 6.

The following topics explain other aspects of the SGI NUMA computers:

- “Distributed Shared Memory (DSM)” on page 40
Scalability is the measure of how the work done on a computing system changes as you add CPUs, memory, network bandwidth, I/O capacity, and other resources. Many factors, for example, memory latency across the system, can affect scalability.

In the SGI UV 2000, SGI UV 1000, and SGI UV 100 series systems, memory is physically distributed both within and among the IRU enclosures, which consist of the compute, memory, and I/O blades. However, memory is accessible to and shared by all devices, connected by NUMAlink, within the single-system image (SSI). In other words, all components connected by NUMAlink share a single Linux operating system, and they operate and share the memory fabric of the system. Memory latency is the amount of time required for a processor to retrieve data from memory. Memory latency is lowest when a processor accesses local memory.

The following are the terms used to refer to the types of memory within a system:

- If a processor accesses memory that is on a compute node blade, that memory is referred to as the node’s local memory.
- If processors access memory located on other blade nodes within the IRU or within other NUMAlink IRUs, the memory is referred to as remote memory.
- The total memory within the NUMAlink system is referred to as global memory.

ccNUMA Architecture

As the name implies, the cache-coherent non-uniform memory access (ccNUMA) architecture has two parts, cache coherency and nonuniform memory access, which the following topics describe:

- "Cache Coherency" on page 40
- "Non-uniform Memory Access (NUMA)” on page 41

Cache Coherency

The SGI UV systems use caches to reduce memory latency. Although data exists in local or remote memory, copies of the data can exist in various processor caches throughout the system. Cache coherency keeps the cached copies consistent.
To keep the copies consistent, the ccNUMA architecture uses directory-based coherence protocol. In directory-based coherence protocol, each block of memory (64 bytes) has an entry in a table that is referred to as a directory. Like the blocks of memory that they represent, the directories are distributed among the compute and memory blade nodes. A block of memory is also referred to as a cache line.

Each directory entry indicates the state of the memory block that it represents. For example, when the block is not cached, it is in an unowned state. When only one processor has a copy of the memory block, it is in an exclusive state. And when more than one processor has a copy of the block, it is in a shared state. A bit vector indicates the caches that may contain a copy.

When a processor modifies a block of data, the processors that have the same block of data in their caches must be notified of the modification. The SGI UV systems use an invalidation method to maintain cache coherence. The invalidation method purges all unmodified copies of the block of data, and the processor that wants to modify the block receives exclusive ownership of the block.

Non-uniform Memory Access (NUMA)

In DSM systems, memory is physically located at various distances from the processors. As a result, memory access times (latencies) are different or nonuniform. For example, it takes less time for a processor blade to reference its locally installed memory than to reference remote memory.

About the Data and Process Placement Tools

For cc-NUMA systems, like the SGI UV systems, performance degrades when the application accesses remote memory, versus local memory. Because the Linux operating system has a tendency to migrate processes, SGI recommends that you use the data and process placement tools.

Special optimization applies to SGI UV systems to exploit multiple paths to memory, as follows:

- By default, all pages are allocated with a first touch policy.
- The initialization loop, if executed serially, gets pages from single node.

Perform initialization in parallel, such that each processor initializes data that it is likely to access later for calculation.
In the parallel loop, multiple processors access that one memory.

Figure 5-1 on page 42, shows how to code to get good data placement.

**Figure 5-1 Coding to Get Good Data Placement**

The `dplace(1)` tool, the `taskset(1)` command, and the cpuset tools are built upon the cpusets API. These tools enable your applications to avoid poor data locality caused by process or thread drift from CPU to CPU. The `omplace(1)` tool works like the `dplace(1)` tool and is designed for use with OpenMP applications. The differences among these tools are as follows:

- The `taskset(1)` command restricts execution to the listed set of CPUs when you specify the `-c` or `--cpu-list` option. The process is free to move among the CPUs that you specify.

- The `dplace(1)` tool differs from `taskset(1)` in that `dplace(1)` binds processes to specified CPUs in round-robin fashion. After a process is pinned, it does not
migrate, so you can use this for high performance and reproducibility of parallel codes.

- Cpusets are named subsets of system cpus/memories and are used extensively in batch environments. For more information about cpusets, see the *SGI Cpuset Software Guide*.

The following topics provide more information about the data and process placement utilities:

- "cpusets and cgroups" on page 43
- "dplace Command" on page 44
- "omplace Command" on page 50
- "taskset Command" on page 51
- "numactl Command" on page 53
- "dlook Command" on page 53

**cpusets and cgroups**

SGI systems support both cgroups and cpusets. cpusets are a subsystem of cgroups, and these two facilities are as follows:

- The cpuset facility is a workload manager tool that permits a system administrator to restrict the number of processor and memory resources that a process or set of processes can use. A cpuset defines a list of CPUs and memory nodes. A process contained in a cpuset can execute only on the CPUs in that cpuset and can only allocate memory on the memory nodes in that cpuset. Essentially, a cpuset provides you with a CPU and a memory container or a soft partition within which you can run sets of related tasks. Using cpusets on an SGI UV system improves cache locality and memory access times and can substantially improve an application's performance and runtime repeatability.

  For information about cpusets, see the *SGI Cpuset Software Guide*.

- cgroups allow you to exert finer control over memory than cpusets. If you use cgroups, be aware that their use can result in a 1–5% memory overhead penalty. If you use a batch scheduler, verify that it supports cgroups before you configure cgroups.
For general information about cpusets and cgroups, see one of the following websites:


**dplace Command**

You can use the `dplace(1)` command to improve the performance of processes running on your SGI nonuniform memory access (NUMA) machine.

By default, memory is allocated to a process on the node on which the process is executing. If a process moves from node to node while it is running, a higher percentage of memory references are made to remote nodes. Because remote accesses typically have higher access times, performance can degrade. CPU instruction pipelines also have to be reloaded.

The `dplace(1)` command specifies scheduling and memory placement policies for the process. You can use the `dplace` command to bind a related set of processes to specific CPUs or nodes to prevent process migrations. In some cases, this improves performance because a higher percentage of memory accesses are made to local nodes.

Processes always execute within a cpuset. The cpuset specifies the CPUs on which a process can run. By default, processes usually execute in a cpuset that contains all the CPUs in the system. For information about cpusets, see the *SGI Cpuset Software Guide*.

The `dplace` command creates a placement container that includes all the CPUs, or a subset of CPUs, of a cpuset. The `dplace` process is placed in this container and, by default, is bound to the first CPU of the cpuset associated with the container. Then `dplace` invokes `exec` to execute the command.

The command executes within this placement container and remains bound to the first CPU of the container. As the command forks child processes, the child processes inherit the container and are bound to the next available CPU of the container.

If you do not specify a placement file, `dplace` binds processes sequentially in a round-robin fashion to CPUs of the placement container. For example, if the current cpuset consists of physical CPUs 2, 3, 8, and 9, the first process launched by `dplace` is bound to CPU 2. The first child process forked by this process is bound to CPU 3. The next process, regardless of whether it is forked by a parent or a child, is bound to CPU 8, and so on. If more processes are forked than there are CPUs in the cpuset, binding starts over with the first CPU in the cpuset.
For more information about `dplace(1)`, see the `dplace(1)` man page. The `dplace(1)` man page also includes examples of how to use the command.

**Example 5-1 Using the `dplace` command with MPI Programs**

The following command improves the placement of MPI programs on NUMA systems and verifies placement of certain data structures of a long-running MPI program:

```
% mpirun -np 64 /usr/bin/dplace -s1 -c 0-63 ./a.out
```

The `-s1` parameter causes `dplace(1)` to start placing processes with the second process, `p1`. The first process, `p0`, is not placed because it is associated with the job launch, not with the job itself. The `-c 0-63` parameter causes `dplace(1)` to use processors 0–63.

You can then use the `dlook(1)` command to verify placement of the data structures in another window on one of the slave thread PIDs. For more information about the `dlook` command, see "`dlook Command` on page 53 and the `dlook(1)` man page.

**Example 5-2 Using `dplace` command with OpenMP Programs**

The following command runs an OpenMP program on logical CPUs 4 through 7 within the current cpuset:

```
% efc -o prog -openmp -O3 program.f
% setenv OMP_NUM_THREADS 4
% dplace -c4-7 ./prog
```

**Example 5-3 Using `dplace` command with OpenMP Programs**

The `dplace(1)` command has a static load balancing feature, so you do not have to supply a CPU list. To place `prog1` on logical CPUs 0 through 3 and `prog2` on logical CPUs 4 through 7, type the following:

```
% setenv OMP_NUM_THREADS 4
% dplace ./prog1 &
% dplace ./prog2 &
```

You can use the `dplace -q` command to display the static load information.

**Example 5-4 Using the `dplace` command with Linux commands**

The following examples assume that you run the `dplace` commands from a shell that runs in a cpuset consisting of physical CPUs 8 through 15.
**Command**  
**Run Location**

<table>
<thead>
<tr>
<th>Command</th>
<th>Run Location</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>dplace -c2 date</code></td>
<td>Runs the date command on physical CPU 10.</td>
</tr>
<tr>
<td><code>dplace make linux</code></td>
<td>Runs gcc and related processes on physical CPUs 8 through 15.</td>
</tr>
<tr>
<td><code>dplace -c0-4,6 make linux</code></td>
<td>Runs gcc and related processes on physical CPUs 8 through 12 or 14.</td>
</tr>
<tr>
<td><code>taskset 4,5,6,7 dplace app</code></td>
<td>The taskset command restricts execution to physical CPUs 12 through 15. The dplace command sequentially binds processes to CPUs 12 through 15.</td>
</tr>
</tbody>
</table>

**Example 5-5 Using dplace and a debugger for verification**

To use the dplace command accurately, you should know how your placed tasks are being created in terms of the fork, exec, and pthread_create calls. Determine whether each of these worker calls are an MPI rank task or are groups of pthreads created by rank tasks. Here is an example of two MPI ranks, each creating three threads:

```bash
cat <<EOF > placefile
firsttask cpu=0
  exec name=mpiapp cpu=1
  fork name=mpiapp cpu=4-8:4 exact
  thread name=mpiapp oncpu=4 cpu=5-7 exact thread name=mpiapp oncpu=8 cpu=9-11 exact EOF

# mpirun is placed on cpu 0 in this example
# the root mpiapp is placed on cpu 1 in this example

# or, if your version of dplace supports the "cpurel=" option:
# firsttask cpu=0
# fork name=mpiapp cpu=4-8:4 exact
# thread name=mpiapp oncpu=4 cpurel=1-3 exact

# create 2 rank tasks, each will pthread_create 3 more
```
# ranks will be on 4 and 8
# thread children on 5,6,7 9,10,11

dplace -p placefile mpirun -np 2 ~cpw/bin/mpiapp -P 3 -l

exit

You can use the debugger to determine if it is working. It should show two MPI rank applications, each with three pthreads, as follows:

```
>> pthreads | grep mpiapp
px *(task_struct *)e00002343c528000 17769 17769 17763 0  mpiapp
   member task: e000013817540000 17795 17769 17763 0  5 mpiapp
   member task: e000013473aa8000 17796 17769 17763 0  6 mpiapp
   member task: e000013817c68000 17798 17769 17763 0  mpiapp
px *(task_struct *)e0000234704f0000 17770 17770 17763 0  mpiapp
   member task: e000023466ed8000 17794 17770 17763 0  9 mpiapp
   member task: e00002384cce0000 17797 17770 17763 0  mpiapp
   member task: e00002342c448000 17799 17770 17763 0  mpiapp
```

You can also use the debugger to see a root application, the parent of the two MPI rank applications, as follows:

```
>> ps | grep mpiapp
0xe00000340b30000 1139 17763 17729 1 0xc800000 - mpiapp
0xe00002343c528000 1139 17769 17763 0 0xc800040 - mpiapp
0xe0000234704f0000 1139 17770 17763 0 0xc800040 8 mpiapp
```

These are placed as specified:

```
>> oncpus e00002343c528000 e000013817540000 e000013473aa8000
>> e000013817c68000 e0
000234704f0000 e000023466ed8000 e00002384cce0000 e00002342c448000
  task: 0xe00002343c528000 mpiapp cpus_allowed: 4
  task: 0xe000013817540000 mpiapp cpus_allowed: 5
  task: 0xe000013473aa8000 mpiapp cpus_allowed: 6
  task: 0xe000013817c68000 mpiapp cpus_allowed: 7
  task: 0xe0000234704f0000 mpiapp cpus_allowed: 8
  task: 0xe000023466ed8000 mpiapp cpus_allowed: 9
  task: 0xe00002384cce0000 mpiapp cpus_allowed: 10
  task: 0xe00002342c448000 mpiapp cpus_allowed: 11
```
Example 5-6 Using dplace for compute thread placement troubleshooting

Sometimes compute threads do not end up on unique processors when using commands such as `dplace(1)` or `profile.pl`. For information about Perfsuite, see the following:

"Profiling with Perfsuite" on page 17

In this example, a user used the `dplace -s1 -c0-15` command to bind 16 processes to run on 0-15 CPUs. However, output from the `top(1)` command shows only 13 CPUs running with CPUs 13, 14, and 15 still idle, and CPUs 0, 1 and 2 are shared with 6 processes.

263 processes: 225 sleeping, 18 running, 3 zombie, 17 stopped
CPU states: cpu user nice system irq softirq iowait idle

<table>
<thead>
<tr>
<th></th>
<th>user</th>
<th>nice</th>
<th>system</th>
<th>irq</th>
<th>softirq</th>
<th>iowait</th>
<th>idle</th>
</tr>
</thead>
<tbody>
<tr>
<td>total</td>
<td>1265.6%</td>
<td>0.0%</td>
<td>28.8%</td>
<td>0.0%</td>
<td>11.2%</td>
<td>0.0%</td>
<td>291.2%</td>
</tr>
<tr>
<td>cpu00</td>
<td>100.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu01</td>
<td>90.1%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>9.7%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu02</td>
<td>99.9%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu03</td>
<td>99.9%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu04</td>
<td>100.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu05</td>
<td>100.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu06</td>
<td>100.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu07</td>
<td>88.4%</td>
<td>0.0%</td>
<td>10.6%</td>
<td>0.0%</td>
<td>0.8%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu08</td>
<td>100.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu09</td>
<td>99.9%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu10</td>
<td>99.9%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu11</td>
<td>88.1%</td>
<td>0.0%</td>
<td>11.2%</td>
<td>0.0%</td>
<td>0.6%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
<tr>
<td>cpu12</td>
<td>99.7%</td>
<td>0.0%</td>
<td>0.2%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
<td>0.0%</td>
</tr>
</tbody>
</table>
cpu13 0.0% 0.0% 2.5% 0.0% 0.0% 0.0% 97.4%
cpu14 0.8% 0.0% 1.6% 0.0% 0.0% 0.0% 97.5%
cpu15 0.0% 0.0% 2.4% 0.0% 0.0% 0.0% 97.5%
Mem: 60134432k av, 15746912k used, 44387520k free, 0k shrd, 672k buff
351024k active, 13594288k inactive
Swap: 2559968k av, 0k used, 2559968k free
2652128k cached

PID USER PRI NI SIZE RSS SHARE STAT %CPU %MEM TIME CPU COMMAND
7653 ccao 25 0 115G 586M 114G R 99.9 0.9 0:08 3 mocassin
7656 ccao 25 0 115G 586M 114G R 99.9 0.9 0:08 6 mocassin
7654 ccao 25 0 115G 586M 114G R 99.8 0.9 0:08 4 mocassin
7655 ccao 25 0 115G 586M 114G R 99.8 0.9 0:08 5 mocassin
7658 ccao 25 0 115G 586M 114G R 99.8 0.9 0:08 8 mocassin
7659 ccao 25 0 115G 586M 114G R 99.8 0.9 0:08 9 mocassin
7660 ccao 25 0 115G 586M 114G R 99.8 0.9 0:08 10 mocassin
7662 ccao 25 0 115G 586M 114G R 99.7 0.9 0:08 12 mocassin
7657 ccao 25 0 115G 586M 114G R 88.5 0.9 0:07 7 mocassin
7661 ccao 25 0 115G 586M 114G R 88.3 0.9 0:07 11 mocassin
7649 ccao 25 0 115G 586M 114G R 55.2 0.9 0:04 2 mocassin
7651 ccao 25 0 115G 586M 114G R 54.1 0.9 0:03 1 mocassin
7650 ccao 25 0 115G 586M 114G R 50.0 0.9 0:04 0 mocassin
An application can start some threads executing for a very short time, yet the threads still have taken a token in the CPU list. Then, when the compute threads are finally started, the list is exhausted and restarts from the beginning. Consequently, some threads end up sharing the same CPU. To bypass this, try to eliminate the ghost thread creation, as follows:

- Check for a call to the system function. This is often responsible for the placement failure due to unexpected thread creation. If all the compute processes have the same name, you can do this by issuing a command such as the following:

  ```
  % dplace -c0-15 -n compute-process-name ...
  ```

- You can also run `dplace -e -c0-32` on 16 CPUs to understand the pattern of the thread creation. If this pattern is the same from one run to the other (unfortunately race between thread creation often occurs), you can find the right flag to `dplace`. For example, if you want to run on CPU 0-3, with `dplace -e -c0-16` and you see that threads are always placed on CPU 0, 1, 5, and 6, then `dplace -e -c0,1,x,x,x,2,3` or `dplace -x24 -c0-3` (24 =11000, place the 2 first and skip 3 before placing) should place your threads correctly.

**omplace Command**

The `omplace` command controls the placement of MPI processes and OpenMP threads. This command is a wrapper script for `dplace`. Use `omplace`, rather than `dplace`, if your application uses MPI, OpenMP, pthreads, or hybrid MPI/OpenMP and MPI/pthreads codes. The `omplace` command generates the proper `dplace` placement file syntax automatically. It also supports some unique options, such as block-strided CPU lists.

The `omplace` command causes the successive threads in a hybrid MPI/OpenMP job to be placed on unique CPUs. The CPUs are assigned in order from the effective CPU list within the containing cpuset. The CPU placement is performed by dynamically generating a placement file and invoking `dplace` with the MPI job launch.
For example, to run two MPI processes with four threads per process, and to display
the generated placement file, type a command similar to the following:

```
# mpirun -np 2 omplace -nt 4 -vv ./a.out
```

The preceding command places the threads as follows:

```
rank 0 thread 0 on CPU 0
rank 0 thread 1 on CPU 1
rank 0 thread 2 on CPU 2
rank 0 thread 3 on CPU 3
rank 1 thread 0 on CPU 4
rank 1 thread 1 on CPU 5
rank 1 thread 2 on CPU 6
rank 1 thread 3 on CPU 7
```

For more information, see the `omplace(1)` man page and the *Message Passing Toolkit (MPT) User’s Guide*.

**taskset Command**

You can use the `taskset(1)` command to perform the following tasks:

- Restricting execution to a list of CPUs. Use the `-c` parameter and the
  `--cpu-list` parameter.

- Retrieving or setting the CPU affinity of a process. Use the following parameters:

  ```
  taskset [options] mask command [arg]...
  taskset [options] -p [mask] pid
  ```

-Launching a new command with a specified CPU affinity.

*CPU affinity* is a scheduler property that bonds a process to a given set of CPUs on
the system. The Linux scheduler honors the given CPU affinity and runs the process
only on the specified CPUs. The process does not run on any other CPUs. Note that
the scheduler also supports natural CPU affinity in which the scheduler attempts to
keep processes on the same CPU as long as practical for performance reasons.
Forcing a specific CPU affinity is useful only in certain applications.

The CPU affinity is represented as a bitmask, with the lowest order bit corresponding
to the first logical CPU and the highest order bit corresponding to the last logical
CPU. The `mask` parameter can specify more CPUs than are present. In other words, it
might be true that not all CPUs specified in the mask exist on a given system. A retrieved mask reflects only the bits that correspond to CPUs physically on the system. If the mask does not correspond to any valid CPUs on the system, the mask is invalid, and the system returns an error. The masks are typically specified in hexadecimal notation. For example:

<table>
<thead>
<tr>
<th>mask specification</th>
<th>CPUs specified</th>
</tr>
</thead>
<tbody>
<tr>
<td>0x00000001</td>
<td>Processor #0</td>
</tr>
<tr>
<td>0x00000003</td>
<td>Processors #0 and #1</td>
</tr>
<tr>
<td>0xFFFFFFFF</td>
<td>All processors (#0 through #31)</td>
</tr>
</tbody>
</table>

When taskset(1) returns, it is guaranteed that the given program has been scheduled to a valid CPU.

The taskset(1) command does not pin a task to a specific CPU. It only restricts a task so that it does not run on any CPU that is not in the CPU list. For example, if you use taskset(1) to launch an application that forks multiple tasks, it is possible that the scheduler initially assigns multiple tasks to the same CPU, even though there are idle CPUs that are in the CPU list. Scheduler load balancing software eventually distributes the tasks so that CPU-bound tasks run on different CPUs. However, the exact placement is not predictable and can vary from run to run. After the tasks are evenly distributed, a task can jump to a different CPU. This outcome can affect memory latency as pages that were node-local before the jump can be remote after the jump.

If you are running an MPI application, SGI recommends that you do not use the taskset(1) command because the taskset(1) command can pin the MPI shepherd process, which wastes a CPU, and then put the remaining working MPI rank on one of the CPUs that already had some other rank running on it. Instead of taskset(1), SGI recommends that you use the dplace(1) command or the environment variable MPI_DSM_CPULIST. For more information, see "dplace Command" on page 44.

If you are using a batch scheduler that creates and destroys cpusets dynamically, SGI recommends that you use the MPI_DSM_DISTRIBUTE environment variable instead of either MPI_DSM_CPULIST environment variable or the dplace command.

Example 1. The following example shows how to run an MPI program on eight CPUs:

```
# mpirun -np 8 dplace -s1 -c10,11,16-21 myMPlapplication ...
```

Example 2. The following example sets the MPI_DSM_CPULIST variable:

```
# setenv MPI_DSM_CPULIST 10,11,16-21 mpirun -np 8 myMPlapplication ...
```
Example 3. The following example runs an executable on CPU 1. The mask for CPU 1 is 0x2, so type the following:

```
# taskset 0x2 executable_name
```

Example 4. The following example moves PID 14057 to CPU 0. The mask for CPU 0 is 0x1, so type the following:

```
# taskset -p 0x1 14057
```

Example 5. The following example runs an MPI Abaqus/Standard job on an SGI UV system with eight CPUs. Standard input is redirected to /dev/null to avoid a SIGTTIN signal for MPT applications. Type the following:

```
# taskset -c 8-15 ./runme < /dev/null &
```

Example 6. The following example uses the taskset(1) command to lock a given process to a particular CPU, CPU5, and then uses the profile(1) command to profile it. The second command moves the process to another CPU, CPU3. Type the following:

```
# taskset -p -c 5 16269
pid 16269’s current affinity list: 0-15
pid 16269’s new affinity list: 5
# taskset -p 16269 -c 3
pid 16269’s current affinity list: 5
pid 16269’s new affinity list: 3
```

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

**numactl Command**

The numactl(8) command runs processes with a specific NUMA scheduling or memory placement policy. The policy is set for an executable command and inherited by all of its children. In addition numactl(8) can set persistent policy for shared memory segments or files. For more information, see the numactl(8) man page.

**dlook Command**

You can use the dlook(1) command to find out where, in memory, the operating system is placing your application’s pages and how much system and user CPU time
it is consuming. The command allows you to display the memory map and CPU usage for a specified process. For each page in the virtual address space of the process, dlook(1) generates the following information:

- The object that owns the page, such as a file, SYSV shared memory, a device driver, and so on.
- The type of page, such as random access memory (RAM), FETCHOP, IOSPACE, and so on.

If the page type is RAM memory, the following information is displayed:
- Memory attributes, such as, SHARED, DIRTY, and so on
- The node on which the page is located
- The physical address of the page (optional)

**Example 5-7 Using dlook(1) with a PID**

To specify a PID as a parameter to the dlook(1) command, you must be the owner of the process or you must be logged in as the root user. The following dlook(1) command example shows output for the sleep process, with a PID of 191155:

```
$ dlook 191155
```

---

5: Data Process and Placement Tools

---

```bash
Peek:  sleep
Pid:  191155 Fri Sep 27 17:14:01 2013

Process memory map:
00400000-00406000 r-xp 00000000 08:08 262250 /bin/sleep
[0000000000400000-0000000000401000] 1 page on node
4 MEMORY|SHARED
[0000000000401000-0000000000402000] 1 page on node
5 MEMORY|SHARED
[0000000000403000-0000000000404000] 1 page on node
7 MEMORY|SHARED
[0000000000404000-0000000000405000] 1 page on node
8 MEMORY|SHARED

00605000-00606000 rw-p 00005000 08:08 262250 /bin/sleep
[0000000000605000-0000000000606000] 1 page on node
2 MEMORY|RW|DIRTY
```
The `dlook(1)` command generates the name of the process (Peek: `sleep`), the process ID, and time and date it was invoked. It provides total user and system CPU time in seconds for the process.
Under the Process memory map heading, the `dlook(1)` command generates information about a process from the `/proc/pid/cpu` and `/proc/pid/maps` files. On the left, it shows the memory segment with the offsets below in decimal. In the middle of the output, it shows the type of access, time of execution, the PID, and the object that owns the memory, which in this example is `/lib/ld-2.2.4.so`. The characters `s` or `p` indicate whether the page is mapped as sharable (`s`) with other processes or is private (`p`). The right side of the output page shows the number of pages of memory consumed and shows the nodes on which the pages reside. A page is 16,384 bytes.

The node numbers reported by the `dlook(1)` command correspond to Socket numbers reported by the `cpumap(1)` command under the section Processor Numbering on Socket(s). For more information, see the `cpumap(1)` command description in "Determining System Configuration" on page 9.

Dirty memory means that the memory has been modified by a user.

**Example 5-8 Using dlook(1) with a command**

When you pass a command as an argument to `dlook(1)`, you specify the command and optional command arguments. The `dlook(1)` command issues an `exec` call on the command and passes the command arguments. When the process terminates, `dlook(1)` prints information about the process, as shown in the following example:

```bash
$ dlook date

Thu Aug 22 10:39:20 CDT 2002
_______________________________________________________________________________
Exit: date

Process memory map:
2000000000030000-200000000003c000 rw-p 0000000000000000 00:00 0
[2000000000030000-200000000003c000] 3 pages on node 3 MEMORY DIRTY

20000000002dc000-20000000002e4000 rw-p 0000000000000000 00:00 0
[20000000002dc000-20000000002e4000] 2 pages on node 3 MEMORY DIRTY

2000000000324000-2000000000334000 rw-p 0000000000000000 00:00 0
[2000000000324000-2000000000328000] 1 page on node 3 MEMORY DIRTY

4000000000000000-400000000000c000 r-xp 0000000000000000 04:03 9657220 /bin/date
[4000000000000000-400000000000c000] 3 pages on node 1 MEMORY SHARED
```
Example 5-9 Using the dlook(1) command with the -s secs option

If you use the dlook(1) command with the -s secs option, the information is sampled at regular internals. The example command and output are as follows:

$ dlook -s 5 sleep 50
Exit: sleep
Pid: 5617 Thu Aug 22 11:16:05 2002

Process memory map:
2000000000030000-200000000003c000 rw-p 0000000000000000 00:00 0
[2000000000030000-200000000003c000] 3 pages on node 3 MEMORY|DIRTY

2000000000034000-2000000000038000 rw-p 0000000000000000 00:00 0
[2000000000034000-2000000000038000] 1 page on node 3 MEMORY|DIRTY

4000000000000000-4000000000000000 r-xp 0000000000000000 04:03 9657225 /bin/sleep
[4000000000000000-4000000000000000] 2 pages on node 3 MEMORY|SHARED

6000000000004000-6000000000008000 rw-p 0000000000000000 04:03 9657225 /bin/sleep
[6000000000004000-6000000000008000] 1 page on node 3 MEMORY|DIRTY

6000000000008000-6000000000010000 rw-p 0000000000000000 04:03 9657220 /bin/date
[6000000000008000-6000000000010000] 1 page on node 3 MEMORY|DIRTY
Example 5-10 Using the dlook command with the mpirun command

You can run a Message Passing Interface (MPI) job using the mpirun command and generate the memory map for each thread, or you can redirect the output to a file.

In the following example, the output has been abbreviated and bold headings added for easier reading:

```
$ mpirun -np 8 dlook -o dlook.out ft.C.8
```

Contents of dlook.out:

```
Exit: ft.C.8
Pid: 2306 Fri Aug 30 14:33:37 2002

Process memory map:
2000000000030000-200000000003c000 rw-p 0000000000000000 00:00 0
  [2000000000030000-2000000000034000] 1 page on node 21 MEMORY|DIRTY
  [2000000000034000-200000000003c000] 2 pages on node 12 MEMORY|DIRTY|SHARED

2000000000044000-2000000000060000 rw-xp 0000000000000000 00:00 0
  [2000000000044000-2000000000050000] 3 pages on node 12 MEMORY|DIRTY|SHARED
  ...

Exit: ft.C.8
Pid: 2310 Fri Aug 30 14:33:37 2002

Process memory map:
2000000000030000-200000000003c000 rw-p 0000000000000000 00:00 0
  [2000000000030000-2000000000034000] 1 page on node 25 MEMORY|DIRTY
  [2000000000034000-200000000003c000] 2 pages on node 12 MEMORY|DIRTY|SHARED
```
For more information about the dlook(1) command, see the dlook(1) man page.
Performance Tuning

This chapter includes the following topics:

- "About Performance Tuning" on page 61
- "Single Processor Code Tuning" on page 62
- "Multiprocessor Code Tuning" on page 71
- "Understanding Parallel Speedup and Amdahl’s Law" on page 77
- "Gustafson’s Law" on page 82
- "Floating-point Program Performance" on page 83
- "About MPI Application Tuning" on page 83
- "Using Transparent Huge Pages (THPs) in MPI and SHMEM Applications" on page 87
- "Enabling Huge Pages in MPI and SHMEM Applications on Systems Without THP" on page 88

About Performance Tuning

After analyzing your code to determine where performance bottlenecks are occurring, you can turn your attention to making your programs run their fastest. One way to do this is to use multiple CPUs in parallel processing mode. However, this should be the last step. The first step is to make your program run as efficiently as possible on a single processor system and then consider ways to use parallel processing.

Intel provides tuning information, including information about the Intel processors, at the following website:


This chapter describes the process of tuning your application for a single processor system and then tuning it for parallel processing. It also addresses how to improve the performance of floating-point programs and MPI applications.
Single Processor Code Tuning

Several basic steps are used to tune performance of single-processor code:

- Get the expected answers and then tune performance. For details, see "Getting the Correct Results" on page 62.
- Use existing tuned code, such as that found in math libraries and scientific library packages. For details, see "Using Tuned Code" on page 63.
- Determine what needs tuning. For details, see "Determining Tuning Needs" on page 64.
- Use the compiler to do the work. For details, see "Using Compiler Options Where Possible" on page 64.
- Consider tuning cache performance. For details, see "Tuning the Cache Performance" on page 67.
- Set environment variables to enable higher-performance memory management mode. For details, see "Managing Memory" on page 69.

Getting the Correct Results

One of the first steps in performance tuning is to verify that the correct answers are being obtained. Once the correct answers are obtained, tuning can be done. You can verify answers by initially disabling specific optimizations and limiting default optimizations. This can be accomplished by using specific compiler options and by using debugging tools.

The following compiler options emphasize tracing and porting over performance:

- \texttt{-O}: the \texttt{-O0} option disables all optimization. The default is \texttt{-O2}.
- \texttt{-g}: the \texttt{-g} option preserves symbols for debugging. In the past, using \texttt{-g} automatically put down the optimization level. In Intel compiler today, you can use \texttt{-O3} with \texttt{-g}.
- \texttt{-fp-model}: the \texttt{-fp-model} option lets you specify the compiler rules for:
  - Value safety
  - Floating-point (FP) expression evaluation
  - FPU environment access
- Precise FP exceptions
- FP contractions

Default is `-fp-model fast=1`. Note that `-mp` option is an old flag replaced by `-fp-model`.

- `-r`, `-i`: the `-r8` and `-i8` options set default real, integer, and logical sizes to 8 bytes, which are useful for porting codes from Cray, Inc. systems. This explicitly declares intrinsic and external library functions.

Some debugging tools can also be used to verify that correct answers are being obtained. See "Using TotalView" on page 21 for more details.

Managing Heap Corruption Problems

You can use environment variables to check for heap corruption problems in programs that use glibc `malloc/free` dynamic memory management routines.

Set the `MALLOC_CHECK_` environment variable to 1 to print diagnostic messages or to 2 to abort immediately when heap corruption is detected.

Overruns and underruns are circumstances where an access to an array is outside the declared boundary of the array. Underruns and overruns cannot be simultaneously detected. The default behavior is to place inaccessible pages immediately after allocated memory.

Using Tuned Code

Where possible, use code that has already been tuned for optimum hardware performance.

The following mathematical functions should be used where possible to help obtain best results:

- MKL: Intel’s Math Kernel Library. This library includes BLAS, LAPACK, and FFT routines.
- VML: the Vector Math Library, available as part of the MKL package (libmkl_vml_itp.so).
- Standard Math library
Standard math library functions are provided with the Intel compiler’s libimf.a file. If the -lm option is specified, glibc libm routines are linked in first.

Documentation is available for MKL and VML at the following website:

Determining Tuning Needs

Use the following tools to determine what points in your code might benefit from tuning:

- **time**: Use this command to obtain an overview of user, system, and elapsed time.
- **gprof**: Use this tool to obtain an execution profile of your program (a pcsamp profile). Use the -p compiler option to enable gprof use.
- **VTune**: This is an Intel performance monitoring tool. You can run it directly on your SGI UV system. The Linux server/Windows client is useful when you are working on a remote system.
- **psrun** is a PerfSuite command-line utility that allows you to take performance measurements of unmodified executables. psrun takes as input a configuration XML document that describes the desired measurement. For more information, see the following website:
  http://perfsuite.ncsa.uiuc.edu/

For information about other performance analysis tools, see Chapter 3, "Performance Analysis and Debugging" on page 9.

Using Compiler Options Where Possible

Several compiler options can be used to optimize performance. For a short summary of ifort or icc options, use the -help option on the compiler command line. Use the -dryrun option to show the driver tool commands that ifort or icc generate. This option does not actually compile.

Use the following options to help tune performance:

- **-ftz**: Flushes underflow to zero to avoid kernel traps. Enabled by default at -O3 optimization.
- `fno-alias`: Assumes no pointer aliasing. Pointer aliasing can create uncertainty about the possibility that two unrelated names might refer to the identical memory; because of this uncertainty, the compiler assumes that any two pointers can point to the same location in memory. This can remove optimization opportunities, particularly for loops.

Other aliasing options include `-ansi_alias` and `-fno_fnalias`. Note that incorrect alias assertions may generate incorrect code.

- `-ip`: Generates single file, interprocedural optimization; `-ipo` generates multifile, interprocedural optimization.

Most compiler optimizations work within a single procedure (like a function or a subroutine) at a time. This intra-procedural focus restricts optimization possibilities because a compiler is forced to make worst-case assumptions about the possible effects of a procedure. By using inter-procedural analysis, more than a single procedure is analyzed at once and code is optimized. It performs two passes through the code and requires more compile time.

- `-O3`: Enables `-O2` optimizations plus more aggressive optimizations, including loop transformation and prefetching. Loop transformations are found in a transformation file created by the compiler; you can examine this file to see what suggested changes have been made to loops. Prefetch instructions allow data to be moved into the cache before their use. A prefetch instruction is similar to a load instruction.

Note that Level 3 optimization may not improve performance for all programs.


- `-override_limits`: This is an undocumented option that sometimes allows the compiler to continue optimizing when it has hit an internal limit.

- `-prof_gen` and `-prof_use`: Generates and uses profiling information. These options require a three-step compilation process:
  1. Compile with proper instrumentation using `-prof_gen`.
  2. Run the program on one or more training datasets.
  3. Compile with `-prof_use`, which uses the profile information from the training run.
-S: Compiles and generates an assembly listing in the .s files and does not link. The assembly listing can be used in conjunction with the output generated by the -opt_report option to try to determine how well the compiler is optimizing loops.

-vec-report: For information specific to the vectorizer. Intel Xeon 7500 series processors can perform short vector operations which provides a powerful performance boost.

-fast: equivalent to writing: -ipo -O3 -no-prec-div -static -xHos

-­xHost: Can generate instructions for the highest instruction set and processor available on the compilation host.

Specific processor architecture to compile for: -xSSE4.2 for Nehalem EP/EX, for example. Useful if compiling in a different system than an SGI UV.

-xSSE4.2: Can generate Intel® SSE4 Efficient Accelerated String and Text Processing instructions supported by Intel® Core i7 processors. Can generate Intel® SSE4 Vectorizing Compiler and Media Accelerator, Intel® SSSE3, SSE3, SSE2, and SSE instructions and it can optimize for the Intel® Core™ processor family.

Another important feature of new Intel compilers is the Source Checker, which is enabled using the flag -diag-enable + options. The source checker is a compiler feature that provides advanced diagnostics based on detailed analysis of source code. It performs static global analysis to find errors in software that go undetected by the compiler itself. It is a general source code analysis tool that provides an additional diagnostic capability to help you debug your programs. You can use source code analysis options to detect potential errors in your compiled code.

Specific processor architecture to compile for: -xSSE4.2 for Nehalem EP/EX Useful if compiling in a different system than an SGI UV.

-xSSE4.2: Can generate Intel® SSE4 Efficient Accelerated String and Text Processing instructions supported by Intel® Core i7 processors. Can generate Intel® SSE4 Vectorizing Compiler and Media Accelerator, Intel® SSSE3, SSE3, SSE2, and SSE instructions and it can optimize for the Intel® Core™ processor family. Another important feature of new Intel compilers is the Source Checker, which is enabled using the flag -diag-enable + options. The source checker is a compiler feature that provides advanced diagnostics based on detailed analysis of source code. It performs static global analysis to find errors in software that go undetected by the compiler itself. general source code analysis tool that provides an additional diagnostic.
capability to help you debug your programs. You can use source code analysis options to detect potential errors in your compiled code including the following:

- Incorrect usage of OpenMP directives
- Inconsistent object declarations in different program units
- Boundary violations
- Uninitialized memory
- Memory corruptions
- Memory Leaks
- Incorrect usage of pointers and allocatable arrays
- Dead code and redundant executions
- Typographical errors or uninitialized variables
- Dangerous usage of unchecked input

Source checker analysis performs a general overview check of a program for all possible values simultaneously. This is in contrast to run-time checking tools that execute a program with a fixed set of values for input variables; such checking tools cannot easily check all edge effects. By not using a fixed set of input values, the source checker analysis can check for all possible corner cases. In fact, you do not need to run the program for Source Checker, the analysis is performed at compilation time. Only requirement is a successful compilation. Important caveat: Limitations of Source Checker Analysis: Since the source checker does not perform full interpretation of analyzed programs, it can generate so called false-positive messages. This is a fundamental difference between the compiler and source checker generated errors; in the case of the source checker, you decide whether the generated error is legitimate and needs to be fixed.

**Tuning the Cache Performance**

The processor cache stores recently-used information in a place where it can be accessed quickly. This discussion uses the following terms:

- A *cache line* is the minimum unit of transfer from next-higher cache into this one.
- A *cache hit* is reference to a cache line which is present in the cache.
A cache miss is reference to a cache line which is not present in this cache level and must be retrieved from a higher cache (or memory or swap space).

The hit time is the time to access the upper level of the memory hierarchy, which includes the time needed to determine whether the access is a hit or a miss.

A miss penalty is the time to replace a block in the upper level with the corresponding block from the lower level, plus the time to deliver this block to the processor. The time to access the next level in the hierarchy is the major component of the miss penalty.

There are several actions you can take to help tune cache performance:

- Avoid large power-of-2 (and multiples thereof) strides and dimensions that cause cache thrashing. Cache thrashing occurs when multiple memory accesses require use of the same cache line. This can lead to an unnecessary number of cache misses.

To prevent cache thrashing, redimension your vectors so that the size is not a power of two. Space the vectors out in memory so that concurrently accessed elements map to different locations in the cache. When working with two-dimensional arrays, make the leading dimension an odd number; for multidimensional arrays, change two or more dimensions to an odd number.

Consider the following example: a cache in the hierarchy has a size of 256 KB (or 65536 4-byte words). A Fortran program contains the following loop:

```fortran
real data(655360,24)
...
do i=1,23
   do j=1,655360
      diff=diff+data(j,i)-data(j,i+1)
   enddo
enddo
```

The two accesses to `data` are separated in memory by 655360*4 bytes, which is a simple multiple of the cache size; they consequently load to the same location in the cache. Because both data items cannot simultaneously coexist in that cache location, a pattern of replace on reload occurs that considerably reduces performance.

- Use a memory stride of 1 wherever possible. A loop over an array should access array elements from adjacent memory addresses. When the loop iterates through
memory by consecutive word addresses, it uses every word of every cache line in sequence and does not return to a cache line after finishing it.

If memory strides other than 1 are used, cache lines could be loaded multiple times if an array is too large to be held in memory at one time.

- Cache bank conflicts can occur if there are two accesses to the same 16-byte-wide bank at the same time.

  A maximum of four performance monitoring events can be counted simultaneously.

- Group together data that is used at the same time and do not use vectors in your code, if possible. If elements that are used in one loop iteration are contiguous in memory, it can reduce traffic to the cache and fewer cache lines will be fetched for each iteration of the loop.

- Try to avoid the use of temporary arrays and minimize data copies.

**Managing Memory**

Nonuniform memory access (NUMA) uses hardware with memory and peripherals distributed among many CPUs. This allows scalability for a shared memory system but a side effect is the time it takes for a CPU to access a memory location. Because memory access times are nonuniform, program optimization is not always straightforward.

Codes that frequently allocate and deallocate memory through glibc malloc/free calls may accrue significant system time due to memory management overhead. By default, glibc strives for system-wide memory efficiency at the expense of performance.

In compilers up to and including version 7.1.x, to enable the higher-performance memory management mode, set the following environment variables:

```
% setenv MALLOC_TRIM_THRESHOLD_ -1
% setenv MALLOC_MMAP_MAX_ 0
```

Because allocations in ifort using the malloc intrinsic use the glibc malloc internally, these environment variables are also applicable in Fortran codes using, for example, Cray pointers with malloc/free. But they do not work for Fortran 90 allocatable arrays, which are managed directly through Fortran library calls and
placed in the stack instead of the heap. The example, above, applies only to the csh shell and the tcsh shell.

Memory Use Strategies

This section describes some general memory use strategies, as follows:

- **Register reuse**: do a lot of work on the same data before working on new data
- **Cache reuse**: the program is much more efficient if all of the data and instructions fit in cache; if not, try to use what is in cache a lot before using anything that is not in cache.
- **Data locality**: try to access data that is near each other in memory before data that is far.
- **I/O efficiency**: do a bunch of I/O all at once rather than a little bit at a time; do not mix calculations and I/O.

Memory Hierarchy Latencies

Programmers tend to think of memory as a flat, random access storage device. It is critical to understand that memory is a hierarchy to get good performance. Memory latency differs within the hierarchy. Performance is affected by where the data resides. Registers: 0 cycles latency (cycle = 1/freq) L1 cache: 1 cycle L2 cache: 5-6 cycles L3 cache: 12-17 cycles Main memory: 130-1000+ cycles. CPUs which are waiting for memory are not doing useful work. Software should be "hierarchy-aware" to achieve best performance:

- Perform as many operations as possible on data in registers
- Perform as many operations as possible on data in the cache(s)
- Keep data uses spatially and temporally local
- Consider temporal locality and spatial locality

Memory hierarchies take advantage of temporal locality by keeping more recently accessed data items closer to the processor. Memory hierarchies take advantage of spatial locality by moving contiguous words in memory to upper levels of the hierarchy.
Multiprocessor Code Tuning

Before beginning any multiprocessor tuning, first perform single processor tuning. This can often obtain good results in multiprocessor codes also. For details, see "Single Processor Code Tuning" on page 62.

Multiprocessor tuning consists of the following major steps:

- Determine what parts of your code can be parallelized. For background information, see "Data Decomposition" on page 71.
- Choose the parallelization methodology for your code. For details, see "Parallelizing Your Code" on page 72.
- Analyze your code to make sure it is parallelizing properly. For details, see Chapter 3, "Performance Analysis and Debugging" on page 9.
- Check to determine if false sharing exists. False sharing refers to OpenMP, not MPI. For details, see "Fixing False Sharing" on page 75.
- Tune for data placement. For details, see Chapter 5, "Data Process and Placement Tools" on page 39.
- Use environment variables to assist with tuning. For details, see "Environment Variables for Performance Tuning" on page 76.

Data Decomposition

In order to efficiently use multiple processors on a system, tasks have to be found that can be performed at the same time. There are two basic methods of defining these tasks:

- Functional parallelism

  Functional parallelism is achieved when different processors perform different functions. This is a known approach for programmers trained in modular programming. Disadvantages to this approach include the difficulties of defining functions as the number of processors grow and finding functions that use an equivalent amount of CPU power. This approach may also require large amounts of synchronization and data movement.

- Data parallelism
Data parallelism is achieved when different processors perform the same function on different parts of the data. This approach takes advantage of the large cumulative memory. One requirement of this approach, though, is that the problem domain be decomposed. There are two steps in data parallelism:

1. Data decomposition

   *Data decomposition* is breaking up the data and mapping data to processors. Data can be broken up explicitly by the programmer by using message passing (with MPI) and data passing (using the SHMEM library routines) or can be done implicitly using compiler-based MP directives to find parallelism in implicitly decomposed data.

   There are advantages and disadvantages to implicit and explicit data decomposition:

   - **Implicit decomposition advantages**: No data resizing is needed; all synchronization is handled by the compiler; the source code is easier to develop and is portable to other systems with OpenMP or High Performance Fortran (HPF) support.

   - **Implicit decomposition disadvantages**: The data communication is hidden by the user

   - **Explicit decomposition advantages**: The programmer has full control over insertion of communication and synchronization calls; the source code is portable to other systems; code performance can be better than implicitly parallelized codes.

   - **Explicit decomposition disadvantages**: Harder to program; the source code is harder to read and the code is longer (typically 40% more).

2. The final step is to divide the work among processors.

Parallelizing Your Code

The first step in multiprocessor performance tuning is to choose the parallelization methodology that you want to use for tuning. This section discusses those options in more detail.

You should first determine the amount of code that is parallelized. Use the following formula to calculate the amount of code that is parallelized:

\[ p = \frac{N(T(1) - T(N))}{T(1)(N-1)} \]
In this equation, $T(1)$ is the time the code runs on a single CPU and $T(N)$ is the time it runs on $N$ CPUs. Speedup is defined as $T(1)/T(N)$.

If $\frac{\text{speedup}}{N}$ is less than 50% (that is, $N>(2-p)/(1-p)$), stop using more CPUs and tune for better scalability.

CPU activity can be displayed with the `top` or `vmstat` commands or accessed by using the Performance Co-Pilot tools (for example, `pmval kernel_percpu_cpu_user`) or by using the Performance Co-Pilot visualization tools `pmchart`.

Next you should focus on a parallelization methodology, as discussed in the following subsections.

**Use MPT**

You can use the Message Passing Interface (MPI) from the SGI Message Passing Toolkit (MPT). MPI is optimized and more scalable for SGI UV series systems than generic MPI libraries. It takes advantage of the SGI UV architecture and SGI Linux NUMA features. MPT is included with the SGI MPI, part of the SGI Performance Suite software.

Use the `-lmpi` compiler option to use MPI. For a list of environment variables that are supported, see the `mpi` man page.

`MPIO_DIRECT_READ` and `MPIO_DIRECT_WRITE` are supported under Linux for local XFS filesystems in SGI MPT version 1.6.1 and beyond.

MPI provides the MPI-2 standard MPI I/O functions that provide file read and write capabilities. A number of environment variables are available to tune MPI I/O performance. See the `mpi_io(3)` man page for a description of these environment variables.

Performance tuning for MPI applications is described in more detail in the Message Passing Toolkit (MPT) User’s Guide.

**Use OpenMP**

OpenMP is a shared memory multiprocessing API, which standardizes existing practice. It is scalable for fine or coarse grain parallelism with an emphasis on performance. It exploits the strengths of shared memory and is directive-based. The OpenMP implementation also contains library calls and environment variables.
OpenMP is included with the C, C++, and Fortran compilers. To use OpenMP directives, specify the following compiler options:

- `ifort -openmp` or `icc -openmp`: These options use the OpenMP front-end that is built into the Intel compilers. The latest Intel compiler OpenMP runtime name is `libiomp5.so`. The latest Intel compiler also supports the GNU OpenMP OpenMP library as an either/or option (not to be mixed-and-matched with the Intel version).

For details about OpenMP usage see the OpenMP standard, available at http://www.openmp.org/specs.

**OpenMP Nested Parallelism**

This section describes OpenMP nested parallelism. For additional information, see the `dplace(1)` man page.

The following OpenMP nested parallelism output shows 2 primary threads and 4 secondary threads, called master/nested:

```
% cat place_nested
firsttask cpu=0
thread name=a.out oncpu=0 cpu=4 noplace=1 exact onetime thread name=a.out oncpu=0
 cpu=1-3 exact thread name=a.out oncpu=4 cpu=5-7 exact

% dplace -p place_nested a.out
Master thread 0 running on cpu 0
Master thread 1 running on cpu 4
Nested thread 0 of master 0 gets task 0 on cpu 0 Nested thread 1 of master 0 gets task 1 on cpu 1
Nested thread 2 of master 0 gets task 2 on cpu 2 Nested thread 3 of master 0 gets task 3 on cpu 3
Nested thread 0 of master 1 gets task 0 on cpu 4 Nested thread 1 of master 1 gets task 1 on cpu 5
Nested thread 2 of master 1 gets task 2 on cpu 6 Nested thread 3 of master 1 gets task 3 on cpu 7
```

**Use Compiler Options**

Use the compiler to invoke automatic parallelization. Use the `-parallel` and `-par_report` option to the `ifort` or `icc` compiler. These options show which loops were parallelized and the reasons why some loops were not parallelized. If a source file contains many loops, it might be necessary to add the `-override_limits` flag to enable automatic parallelization. The code generated by `-parallel` is based on the OpenMP API, the standard OpenMP environment variables and Intel extensions apply.
There are some limitations to automatic parallelization:

- For Fortran codes, only DO loops are analyzed.
- For C/C++ codes, only for loops using explicit array notation or those using pointer increment notation are analyzed. In addition, for loops using pointer arithmetic notation are not analyzed nor are while or do/while loops. The compiler also does not check for blocks of code that can be run in parallel.

Identifying Parallel Opportunities in Existing Code

Another parallelization optimization technique is to identify loops that have a potential for parallelism, such as the following:

- Loops without data dependencies; a data dependency conflict occurs when a loop has results from one loop pass that are needed in future passes of the same loop.
- Loops with data dependencies because of temporary variables, reductions, nested loops, or function calls or subroutines.

Loops that do not have a potential for parallelism are those with premature exits, too few iterations, or those where the programming effort to avoid data dependencies is too great.

Fixing False Sharing

If the parallel version of your program is slower than the serial version, false sharing might be occurring. False sharing occurs when two or more data items that appear not to be accessed by different threads in a shared memory application correspond to the same cache line in the processor data caches. If two threads executing on different CPUs modify the same cache line, the cache line cannot remain resident and correct in both CPUs, and the hardware must move the cache line through the memory subsystem to retain coherency. This causes performance degradation and reduction in the scalability of the application. If the data items are only read, not written, the cache line remains in a shared state on all of the CPUs concerned. False sharing can occur when different threads modify adjacent elements in a shared array. When two CPUs share the same cache line of an array and the cache is decomposed, the boundaries of the chunks split at the cache line.

You can use the following methods to verify that false sharing is happening:

- Use the performance monitor to look at output from pfmon and the BUS_MEM_READ_BRIL_SELF and BUS_RD_INVAL_ALL_HITM events.
- Use `pfmon` to check DEAR events to track common cache lines.
- Use the Performance Co-Pilot `pmshub` utility to monitor cache traffic and CPU utilization.

If false sharing is a problem, try the following solutions:
- Use the hardware counter to run a profile that monitors storage to shared cache lines. This will show the location of the problem.
- Revise data structures or algorithms.
- Check shared data, static variables, common blocks, and private and public variables in shared objects.
- Use critical regions to identify the part of the code that has the problem.

**Environment Variables for Performance Tuning**

You can use several different environment variables to assist in performance tuning. For details about environment variables used to control the behavior of MPI, see the `mpi(1)` man page.

Several OpenMP environment variables can affect the actions of the OpenMP library. For example, some environment variables control the behavior of threads in the application when they have no work to perform or are waiting for other threads to arrive at a synchronization semantic; other variables can specify how the OpenMP library schedules iterations of a loop across threads. The following environment variables are part of the OpenMP standard:

- `OMP_NUM_THREADS` (The default is the number of CPUs in the system.)
- `OMP_SCHEDULE` (The default is `static`.)
- `OMP_DYNAMIC` (The default is `false`.)
- `OMP_NESTED` (The default is `false`.)

In addition to the preceding environment variables, Intel provides several OpenMP extensions, two of which are provided through the use of the `KMP_LIBRARY` variable. The `KMP_LIBRARY` variable sets the run-time execution mode, as follows:

- If set to `serial`, single-processor execution is used.
• If set to throughput, CPUs yield to other processes when waiting for work. This is the default and is intended to provide good overall system performance in a multiuser environment.

• If set to turnaround, worker threads do not yield while waiting for work. Setting KMP_LIBRARY to turnaround may improve the performance of benchmarks run on dedicated systems, where multiple users are not contending for CPU resources.

If your program gets a segmentation fault immediately upon execution, you may need to increase KMP_STACKSIZE. This is the private stack size for threads. The default is 4 MB. You may also need to increase your shell stacksize limit.

## Understanding Parallel Speedup and Amdahl’s Law

There are two ways to obtain the use of multiple CPUs. You can take a conventional program in C, C++, or Fortran, and have the compiler find the parallelism that is implicit in the code.

You can write your source code to use explicit parallelism, stating in the source code which parts of the program are to execute asynchronously, and how the parts are to coordinate with each other.

When your program runs on more than one CPU, its total run time should be less. But how much less? What are the limits on the speedup? That is, if you apply 16 CPUs to the program, should it finish in 1/16th the elapsed time?

This section covers the following topics:

• "Adding CPUs to Shorten Execution Time" on page 78
• "Understanding Parallel Speedup" on page 78
• "Understanding Amdahl’s Law" on page 79
• "Calculating the Parallel Fraction of a Program" on page 80
• "Predicting Execution Time with n CPUs" on page 81
Adding CPUs to Shorten Execution Time

You can distribute the work your program does over multiple CPUs. However, there is always some part of the program’s logic that has to be executed serially, by a single CPU. This sets the lower limit on program run time.

Suppose there is one loop in which the program spends 50% of the execution time. If you can divide the iterations of this loop so that half of them are done in one CPU while the other half are done at the same time in a different CPU, the whole loop can be finished in half the time. The result: a 25% reduction in program execution time.

The mathematical treatment of these ideas is called Amdahl’s law, for computer pioneer Gene Amdahl, who formalized it. There are two basic limits to the speedup you can achieve by parallel execution:

- The fraction of the program that can be run in parallel, \( p \), is never 100%.
- Because of hardware constraints, after a certain point, there is less and less benefit from each added CPU.

Tuning for parallel execution comes down to doing the best that you are able to do within these two limits. You strive to increase the parallel fraction, \( p \), because in some cases even a small change in \( p \) (from 0.8 to 0.85, for example) makes a dramatic change in the effectiveness of added CPUs.

Then you work to ensure that each added CPU does a full CPU’s work and does not interfere with the work of other CPUs. In the SGI UV architectures this means:

- Spreading the workload equally among the CPUs
- Eliminating false sharing and other types of memory contention between CPUs
- Making sure that the data used by each CPU are located in a memory near that CPU’s node

Understanding Parallel Speedup

If half the iterations of a loop are performed on one CPU, and the other half run at the same time on a second CPU, the whole loop should complete in half the time. For example, consider the typical C loop in Example 6-1.
Example 6-1 Typical C Loop

```c
for (j=0; j<MAX; ++j) {
    z[j] = a[j]*b[j];
}
```

The compiler can automatically distribute such a loop over \( n \) CPUs (with \( n \) decided at run time based on the available hardware), so that each CPU performs \( \frac{MAX}{n} \) iterations.

The speedup gained from applying \( n \) CPUs, \( \text{Speedup}(n) \), is the ratio of the one-CPU execution time to the \( n \)-CPU execution time: \( \text{Speedup}(n) = \frac{T(1)}{T(n)} \). If you measure the one-CPU execution time of a program at 100 seconds, and the program runs in 60 seconds with two CPUs, \( \text{Speedup}(2) = \frac{100}{60} = 1.67 \).

This number captures the improvement from adding hardware. \( T(n) \) ought to be less than \( T(1) \); if it is not, adding CPUs has made the program slower, and something is wrong. So \( \text{Speedup}(n) \) should be a number greater than 1.0, and the greater it is, the better. Intuitively you might hope that the speedup would be equal to the number of CPUs (twice as many CPUs, half the time) but this ideal can seldom be achieved.

Understanding Superlinear Speedup

You expect \( \text{Speedup}(n) \) to be less than \( n \), reflecting the fact that not all parts of a program benefit from parallel execution. However, it is possible, in rare situations, for \( \text{Speedup}(n) \) to be larger than \( n \). When the program has been sped up by more than the increase of CPUs it is known as superlinear speedup.

A superlinear speedup does not really result from parallel execution. It comes about because each CPU is now working on a smaller set of memory. The problem data handled by any one CPU fits better in cache, so each CPU executes faster than the single CPU could do. A superlinear speedup is welcome, but it indicates that the sequential program was being held back by cache effects.

Understanding Amdahl’s Law

There are always parts of a program that you cannot make parallel, where code must run serially. For example, consider the loop. Some amount of code is devoted to setting up the loop, allocating the work between CPUs. This housekeeping must be done serially. Then comes parallel execution of the loop body, with all CPUs running concurrently. At the end of the loop comes more housekeeping that must be done
serially; for example, if \( n \) does not divide \( \text{MAX} \) evenly, one CPU must execute the few iterations that are left over.

The serial parts of the program cannot be speeded up by concurrency. Let \( p \) be the fraction of the program’s code that can be made parallel (\( p \) is always a fraction less than 1.0.) The remaining fraction (1–\( p \)) of the code must run serially. In practical cases, \( p \) ranges from 0.2 to 0.99.

The potential speedup for a program is proportional to \( p \) divided by the CPUs you can apply, plus the remaining serial part, 1–\( p \). As an equation, this appears as Example 6-2.

**Example 6-2** Amdahl’s law: \( Speedup(n) \) Given \( p \)

\[
Speedup(n) = \frac{1}{(p/n) + (1-p)}
\]

Suppose \( p = 0.8 \); then \( Speedup(2) = 1 / (0.4 + 0.2) = 1.67 \), and \( Speedup(4) = 1 / (0.2 + 0.2) = 2.5 \). The maximum possible speedup (if you could apply an infinite number of CPUs) would be 1 / (1–\( p \)). The fraction \( p \) has a strong effect on the possible speedup.

The reward for parallelization is small unless \( p \) is substantial (at least 0.8); or to put the point another way, the reward for increasing \( p \) is great no matter how many CPUs you have. The more CPUs you have, the more benefit you get from increasing \( p \). Using only four CPUs, you need only \( p = 0.75 \) to get half the ideal speedup. With eight CPUs, you need \( p = 0.85 \) to get half the ideal speedup.

There is a slightly more sophisticated version of Amdahl’s law which includes communication overhead, showing also that if the program has no serial part that as we increase the number of cores the amount of computation per core diminishes and the communication overhead (unless there is no communication and we have trivial parallelization) increases, also diminishing the efficiency of the code and the speedup. The equation is:

\[
Speedup(n) = \frac{n}{(1 + a^n(n-1) + n^a(tc/ts))}
\]

Where: \( n \): number of processes \( a \): the fraction of the given task not dividable into concurrent subtasks \( ts \): time to execute the task in a single processor \( tc \): communication overhead If \( a = 0 \) and \( tc = 0 \) (no serial part and no communications) like in a trivial parallelization program, you will get linear speedup.

**Calculating the Parallel Fraction of a Program**

You do not have to guess at the value of \( p \) for a given program. Measure the execution times \( T(1) \) and \( T(2) \) to calculate a measured \( Speedup(2) = T(1) / T(2) \). The
Amdahl’s law equation can be rearranged to yield \( p \) when \( \text{Speedup}(2) \) is known, as in Example 6-3.

**Example 6-3 Amdahl’s law: \( p \) Given \( \text{Speedup}(2) \)**

\[
\begin{align*}
2 & \quad \text{SpeedUp}(2) - 1 \\
1 & \quad \text{SpeedUp}(2)
\end{align*}
\]

\[
p = \frac{2 \quad \text{SpeedUp}(2) - 1}{1 \quad \text{SpeedUp}(2)}
\]

Suppose you measure \( T(1) = 188 \) seconds and \( T(2) = 104 \) seconds.

\[
\text{SpeedUp}(2) = \frac{188}{104} = 1.81
\]

\[
p = 2 \times \frac{(1.81-1)/1.81}{1} = 2 \times (0.81/1.81) = 0.895
\]

In some cases, the \( \text{Speedup}(2) = T(1)/T(2) \) is a value greater than 2; in other words, a superlinear speedup (‘Understanding Superlinear Speedup’ on page 79). When this occurs, the formula in Example 6-3 returns a value of \( p \) greater than 1.0, which is clearly not useful. In this case you need to calculate \( p \) from two other more realistic timings, for example \( T(2) \) and \( T(3) \). The general formula for \( p \) is shown in Example 6-4, where \( n \) and \( m \) are the two CPU counts whose speedups are known, \( n > m \).

**Example 6-4 Amdahl’s Law: \( p \) Given \( \text{Speedup}(n) \) and \( \text{Speedup}(m) \)**

\[
\begin{align*}
\text{Speedup}(n) - \text{Speedup}(m) & \quad \text{Speedup}(n) - \text{Speedup}(m)
\end{align*}
\]

\[
p = \frac{\text{Speedup}(n) - \text{Speedup}(m)}{(1 - 1/n) \times \text{Speedup}(n) - (1 - 1/m) \times \text{Speedup}(m)}
\]

**Predicting Execution Time with \( n \) CPUs**

You can use the calculated value of \( p \) to extrapolate the potential speedup with higher numbers of CPUs. The following example shows the expected time with four CPUs, if \( p = 0.895 \) and \( T(1) = 188 \) seconds:

\[
\begin{align*}
\text{Speedup}(4) &= \frac{1}{((0.895/4)+(1-0.895))} = 3.04 \\
T(4) &= \frac{T(1)}{\text{Speedup}(4)} = \frac{188}{3.04} = 61.8
\end{align*}
\]

The calculation can be made routine using the computer by creating a script that automates the calculations and extrapolates run times.

These calculations are independent of most programming issues such as language, library, or programming model. They are not independent of hardware issues, because Amdahl’s law assumes that all CPUs are equal. At some level of parallelism, adding a CPU no longer affects run time in a linear way. For example, on some
architectures, cache-friendly codes scale closely with Amdahl’s law up to the maximum number of CPUs, but scaling of memory intensive applications slows as the system bus approaches saturation. When the bus bandwidth limit is reached, the actual speedup is less than predicted.

**Gustafson’s Law**

Gustafson’s law proposes that programmers set the size of problems to use the available equipment to solve problems within a practical fixed time. Therefore, if faster (more parallel) equipment is available, larger problems can be solved in the same time. Amdahl’s law is based on fixed workload or fixed problem size. It implies that the sequential part of a program does not change with respect to machine size (for example, the number of processors). However, the parallel part is evenly distributed by \( n \) processors. The effect of Gustafson’s law was to shift research goals to select or reformulate problems so that solving a larger problem in the same amount of time would be possible. In particular, the law redefines efficiency as a need to minimize the sequential part of a program, even if it increases the total amount of computation. The bottom line is that by running larger problems, it is hoped that the bulk of the calculation will increase faster than the serial part of the program, allowing for better scaling. There is a slightly more sophisticated version of Amdahl’s law which includes communication overhead, showing also that if the program has no serial part that as we increase the number of cores the amount of computation per core diminishes and the communication overhead (unless there is not communication and you have trivial parallelization) increases, also diminishing the efficiency of the code and the speedup. The equation is:

\[
\text{Speedup}(n) = \frac{n}{1 + a \ast (n-1) + n \ast (t_c/t_s)}
\]

The preceding equation uses the following variables:

- \( n \): number of processes
- \( a \): the fraction of the given task not dividable into concurrent subtasks
- \( ts \): time to execute the task in a single processor
- \( tc \): communication overhead

If \( a=0 \) and \( t_c=0 \), which indicated no serial part and no communications, like in a trivial parallelization program, the result is linear speedup.
Floating-point Program Performance

Certain floating-point programs experience slowdowns due to excessive floating point traps called Floating-Point Software Assist (FPSWA).

This happens when the hardware cannot complete a floating point operation and requests help (emulation) from software. This happens, for instance, with denormals numbers.

The symptoms are a slower than normal execution, FPSWA message in the system log (run `dmesg`). The average cost of a FPSWA fault is quite high around 1000 cycles/fault.

By default, the kernel prints a message similar to the following in the system log:

```
foo(7716): floating-point assist fault at ip 40000000000200e1
    isr 0000020000000008
```

The kernel throttles the message in order to avoid flooding the console.

It is possible to control the behavior of the kernel on FPSWA faults using the `prctl(1)` command. In particular, it is possible to get a signal delivered at the first FPSWA. It is also possible to silence the console message.

About MPI Application Tuning

When you design your MPI application, make sure to include the following in your design:

- The pinning of MPI processes to CPUs
- The isolating of multiple MPI jobs onto different sets of sockets and Hubs

You can achieve this design by configuring a batch scheduler 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 `MPI_DSM_CPULIST`. For more information about these programming practices, see the following:

- The `omplace(1)` and `dplace(1)` man pages.
• The SGI Cpuset Software Guide.

**MPI Application Communication on SGI Hardware**

On an SGI UV system, the following two transfer methods facilitate MPI communication between processes:

• Shared memory

• The global reference unit (GRU), which is part of the SGI UV Hub ASIC

The SGI UV series systems use a scalable nonuniform memory access (NUMA) architecture to allow the use of thousands of processors and terabytes of RAM in a single Linux operating system instance. As in other large shared memory systems, memory is distributed to processor sockets and accesses to memory are cache coherent. Unlike other systems, SGI UV systems use a network of Hub ASICs connected over NUMALink to scale to more sockets than any other x86-64 system, with excellent performance out of the box for most applications.

When running on SGI UV systems with SGI’s Message Passing Toolkit (MPT), applications can attain higher bandwidth and lower latency for MPI calls than when running on more conventional distributed memory clusters. However, knowing your SGI UV system’s NUMA topology and the performance constraints that it imposes can still help you extract peak performance. For more information about the SGI UV hub, SGI UV compute blades, Intel QPI and SGI NUMALink, see your SGI UV hardware system user guide.

The MPI library chooses the transfer 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 allocates buffer space and control blocks between the logical processors being used by the MPI job.

**MPI Job Problems and Application Design**

The MPI library chooses buffer sizes and communication algorithms in an attempt to deliver the best performance automatically to a wide variety of MPI applications, but user tuning might be needed to improve performance. The following are some application performance problems and some ways that you might be able to improve MPI performance:
Primary HyperThreads are idle.

Most high performance computing MPI programs run best when they use only one HyperThread per core. When an SGI UV system has multiple HyperThreads per core, logical CPUs are numbered such that primary HyperThreads are the high half of the logical CPU numbers. Therefore, the task of scheduling only on the additional HyperThreads may be accomplished by scheduling MPI jobs as if only half the full number exists, 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 command’s output includes the following:

- The number of physical and logical processors
- Whether Hyperthreading is enabled
- How shared processors are paired

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 the `gru_resource(3)` man page.

Message bandwidth is inadequate.

Use either huge pages or transparent huge pages (THP) to ensure that your application obtains optimal message bandwidth.

To specify the use of hugepages, use the `MPI_HUGEPAGE_HEAP_SPACE` environment variable. The `MPI_HUGEPAGE_HEAP_SPACE` environment variable defines the minimum amount of heap space that each MPI process can allocate using huge pages. For information about this environment variable, see the `MPI(1)` man page.

To use THPs, see "Using Transparent Huge Pages (THPs) in MPI and SHMEM Applications" on page 87.

Some programs transfer large messages via the `MPI_Send` function. To enable unbuffered, single-copy transport in these cases, you can set `MPI_BUFFER_MAX` to 0. For information about the `MPI_BUFFER_MAX` environment variable, see the `MPI(1)` man page.

MPI small or near messages are very frequent.
For small fabric hop counts, shared memory message delivery is faster than GRU messages. To deliver all messages within an SGI UV host via shared memory, set MPI_SHARED_NEIGHBORHOOD to host. For more information, see the MPI(1) man page.

- Memory allocations are nonlocal.

MPI application processes normally perform best if their local memory is allocated near the socket assigned to use it. This cannot happen if memory on that socket is exhausted by the application or by other system consumption, for example, file buffer cache. Use the nodeinfo(1) command to view memory consumption on the nodes assigned to your job, and use bcfree(1) to clear out excessive file buffer cache. PBS Professional batch scheduler installations can be configured to issue bcfree commands in the job prologue.

For information about PBS Professional, including the availability of scripts, see the PBS Professional documentation and the bcfree(1) man page.

### MPI Performance Tools

SGI supports several MPI performance tools. You can use the following tools to enhance or troubleshoot MPI program performance:

- **MPInside.** MPInside is an MPI profiling tool that can help you optimize your MPI application. The tool provides information about data transferred between ranks, both in terms of speed and quantity. For information about MPInside, see one of the following:
  - The MPInside(1) man page.
  - The MPInside Reference Guide.

- **SGI 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. The PerfBoost software allows you to run SGI MPT which is a version of MPI optimized for SGI large, shared-memory systems and can take advantage of the UV Hub.


- **SGI PerfCatcher.** SGI PerfCatcher uses a wrapper library to return MPI and SHMEM function profiling information. The information returned includes
percent CPU time, total time spent per function, message sizes, and load
imbalances. For more information, see the following:

- The `perfcatch(1)` man page
- The *Message Passing Toolkit (MPT) User Guide*

## Using Transparent Huge Pages (THPs) in MPI and SHMEM Applications

On SGI UV systems, THP is important because it contributes to attaining the best
GRU-based data transfer bandwidth in Message Passing Interface (MPI) and SHMEM
programs. On newer kernels, the THP feature is enabled by default. If THP is
disabled on your SGI UV system, see "Enabling Huge Pages in MPI and SHMEM
Applications on Systems Without THP" on page 88.

On SGI ICE X systems, if you use a workload manager, such as PBS Professional,
your site configuration might let you enable or disable THP on a per-job basis.

The THP feature can affect the performance of some OpenMP threaded applications
in a negative way. For certain OpenMP applications, some threads in some shared
data structures might be forced to make more nonlocal references because the
application assumes a smaller, 4-KB page size.

The THP feature affects users in the following ways:

- Administrators:

  To activate the THP feature on a system-wide basis, write the keyword `always`
to the following file:

  `/sys/kernel/mm/transparent_hugepage/enabled`

  To create an environment in which individual applications can use THP if memory
  is allocated accordingly within the application itself, type the following:

  `# echo madvise > /sys/kernel/mm/transparent_hugepage/enabled`

  To disable THP, type the following command:

  `# echo never > /sys/kernel/mm/transparent_hugepage/enabled`
If the `khugepaged` daemon is taking a lot of time when a job is running, then
defragmentation of THP might be causing performance problems. You can type
the following command to disable defragmentation:

```bash
# echo never > /sys/kernel/mm/transparent_hugepage/defrag
```

If you suspect that defragmentation of THP is causing performance problems, but
you do not want to disable defragmentation, you can tune the `khugepaged`
daemon by editing the values in
`/sys/kernel/mm/transparent_hugepage/khugepaged`.

- MPI application programmers:

  To determine whether THP is enabled on your system, type the following
  command and note the output:

  ```bash
  % cat /sys/kernel/mm/transparent_hugepage/enabled
  ```

  The output is as follows on a system for which THP is enabled:

  `[always] madvise never`

  In the output, the bracket characters (`[]`) appear around the keyword that is in
effect.

**Enabling Huge Pages in MPI and SHMEM Applications on Systems Without
THP**

If the THP capability is disabled on your SGI UV system, you can use the
`MPI_HUGEPAGE_HEAP_SPACE` environment variable and the
`MPT_HUGEPAGE_CONFIG` command to create huge pages.

The `MPT_HUGEPAGE_CONFIG` command configures the system to allow huge pages to
be available to MPT’s memory allocation interceptors. The
`MPI_HUGEPAGE_HEAP_SPACE` environment variable enables an application to use the
huge pages reserved by the `MPT_HUGEPAGE_CONFIG` command.

For more information, see the `MPI_HUGEPAGE_HEAP_SPACE` environment variable on
the `MPI(1)` man page, or see the `mpt_hugepage_config(1)` man page.
Flexible File I/O

Flexible File I/O (FFIO) provides a mechanism for improving the file I/O performance of existing applications without having to resort to source code changes, that is, the current executable remains unchanged. Knowledge of source code is not required, but some knowledge of how the source and the application software work can help you better interpret and optimize FFIO results. To take advantage of FFIO, all you need to do is to set some environment variables before running your application. This chapter covers the following topics:

- "FFIO Operation" on page 89
- "Environment Variables" on page 90
- "Simple Examples" on page 91
- "Multithreading Considerations" on page 94
- "Application Examples" on page 95
- "Event Tracing" on page 96
- "System Information and Issues" on page 96

**FFIO Operation**

The FFIO subsystem allows you to define one or more additional I/O buffer caches for specific files to augment the Linux kernel I/O buffer cache. The FFIO subsystem then manages this buffer cache for you. In order to accomplish this, FFIO intercepts standard I/O calls like open, read, and write, and replaces them with FFIO equivalent routines. These routines route I/O requests through the FFIO subsystem which utilizes the user defined FFIO buffer cache. FFIO can bypass the Linux kernel I/O buffer cache by communicating with the disk subsystem via direct I/O. This gives you precise control over cache I/O characteristics and allows for more efficient I/O requests. For example, doing direct I/O in large chunks (say 16 megabytes) allows the FFIO cache to amortize disk access. All file buffering occurs in user space when FFIO is used with direct I/O enabled. This differs from the Linux buffer cache mechanism which requires a context switch in order to buffer data in kernel memory. Avoiding this kind of overhead, helps FFIO to scale efficiently. Another important distinction is that FFIO allows you to create an I/O buffer cache dedicated to a specific application.
The Linux kernel, on the other hand, has to manage all the jobs on the entire system with a single I/O buffer cache. As a result, FFIO typically outperforms the Linux kernel buffer cache when it comes to I/O intensive throughput.

Environment Variables

There are only two environment variables that you need to set in order to use FFIO. They are `LD_PRELOAD` and `FF_IO_OPTS`.

In order to enable FFIO to trap standard I/O calls, you must set the `LD_PRELOAD` environment variable.

For SGI systems, perform the following:

```bash
# export LD_PRELOAD="/usr/lib64/libFFIO.so"
```

The `LD_PRELOAD` software is a Linux feature that instructs the linker to preload the indicated shared libraries. In this case, `libFFIO.so` is preloaded and provides the routines which replace the standard I/O calls. An application that is not dynamically linked with the `glibc` library will not work with FFIO, since the standard I/O calls will not be intercepted. To disable FFIO, perform the following:

```bash
unset LD_PRELOAD
```

The FFIO buffer cache is managed by the `FF_IO_OPTS` environment variable. The syntax for setting this variable can be quite complex. A simple method for defining this variable is as follows:

```bash
# export FF_IO_OPTS 'string(eie.direct.mbytes:size:num:lead:share:stride:0)'
```

You can use the following parameters with the `FF_IO_OPTS` environment variable:

- `string` Matches the names of files that can use the buffer cache.
- `size` Number of 4k blocks in each page of the I/O buffer cache.
- `num` Number of pages in the I/O buffer cache.
- `lead` The maximum number of read-ahead pages.
- `share` A value of 1 means a shared cache, 0 means private.
**stride**

Note that the number after the *stride* parameter is always 0.

The following example shows a command that creates a shared buffer cache of 128 pages where each page is 16 megabytes (that is, 4096*4k). The cache has a lead of six pages and uses a stride of one, as follows:

```shell
```

Each time the application opens a file, the FFIO code checks the file name to see if it matches the string supplied by `FF_IO_OPTS`. The file’s path name is not considered when checking for a match against the string. So in the example supplied above, file names like `/tmp/test16` and `/var/tmp/testit` would both be a match.

More complicated usages of `FF_IO_OPTS` are built upon this simpler version. For example, multiple types of file names can share the same cache, as follows:

```shell
```

Multiple caches may also be specified with `FF_IO_OPTS`. In the example that follows, files of the form `output*` and `test*` share a 128 page cache of 16 megabyte pages. The file `special42` has a 256 page private cache of 32 megabyte pages, as follows:

```shell
```

Additional parameters can be added to `FF_IO_OPTS` to create feedback that is sent to standard output. Examples of doing this diagnostic output will be presented in the following section.

### Simple Examples

This section walks you through some simple examples using FFIO.

Assume that `LD_PRELOAD` is set for the correct library and `FF_IO_OPTS` is defined, as follows:

```shell
```

This example uses a small C program called `fio` that reads four megabyte chunks from a file for 100 iterations. When the program runs it produces output, as follows:

```shell
./fio -n 100 /build/testit
Reading 4194304 bytes 100 times to /build/testit
Total time = 7.383761
```
Flexible File I/O

Throughput = 56.804439 MB/sec

It can be difficult to tell what FFIO may or may not be doing even with a simple program such as shown above. A summary of the FFIO operations that occurred can be directed to standard output by making a simple addition to FF_IO_OPTS, as follows:

```
```

This new setting for FF_IO_OPTS generates the following summary on standard output when the program is run:

```
./fio -n 100 /build/testit
Reading 4194304 bytes 100 times to /build/testit
Total time = 7.383761
Throughput = 56.804439 MB/sec

event_close(testit) eie <>syscall (496 mbytes)/( 8.72 s)= 56.85 mbytes/s
oflags=0x0000000000004042=RDWR+CREAT+DIRECT
sector size =4096(bytes)
cb1ks =0 cbits =0x0000000000000000
current file size =512 mbytes high water file size =512 mbytes

function  times  wall time  all mbytes  mbytes delivered  min request  max request  avg request

open      1      0.00      
read      2      0.61      32   32      16      16      16
reada     29      0.01      0   464     464      16      16      16
fcntl
    recall
reada     29      8.11      
other     5      0.00      
flush     1      0.00      
close     1      0.00      
```

Two synchronous reads of 16 megabytes each were issued (for a total of 32 megabytes) and 29 asynchronous reads (reada) were also issued (for a total of 464 megabytes). Additional diagnostic information can be generated by specifying the .diag modifier, as follows:

```
```
The `.diag` modifier may also be used in conjunction with `.event.summary`, the two operate independently from one another, as follows:

```
```

An example of the diagnostic output generated when just the `.diag` modifier is used is, as follows:

```
./fio -n 100 /build/testit
Reading 4194304 bytes 100 times to /build/testit
Total time = 7.383761
Throughput = 56.804439 MB/sec

eie_close EIE final stats for file /build/testit
  eie_close Used shared eie cache 1
  eie_close 128 mem pages of 4096 blocks (4096 sectors), max_lead = 6 pages
  eie_close advance reads used/started : 23/29  79.31%  (1.78 seconds wasted)
  eie_close read hits/total : 0/0  0.00%
  eie_close mbytes transferred parent --> eie --> child sync async
   eie_close 0  0  0 0
  eie_close 400  496  2 29 (0,0)
  eie_close parent <-- eie <-- child

eie_close EIE stats for Shared cache 1
  eie_close 128 mem pages of 4096 blocks
  eie_close advance reads used/started : 23/29  79.31%  (0.00 seconds wasted)
  eie_close write hits/total : 0/0  0.00%
  eie_close read hits/total : 98/100  98.00%
  eie_close mbytes transferred parent --> eie --> child sync async
   eie_close 0  0  0 0
  eie_close 400  496  2 29 (0,0)
```

Information is listed for both the file and the cache. An `mbytes transferred` example is shown below:

```
eie_close mbytes transferred parent --> eie --> child sync async
  eie_close 0  0  0 0
  eie_close 400  496  2 29 (0,0)
```

The last two lines are for write and read operations, respectively. Only for very simple I/O patterns, the difference between (parent --> eie) and (eie --> child) read statistics
Flexible File I/O can be explained by the number of read aheads. For random reads of a large file over a long period of time, this is not the case. All write operations count as async.

**Multithreading Considerations**

FFIO will work with applications that use MPI for parallel processing. An MPI job assigns each thread a number or rank. The master thread has rank 0, while the remaining threads (called slave threads) have ranks from 1 to N-1 where N is the total number of threads in the MPI job. It is important to consider that the threads comprising an MPI job do not (necessarily) have access to each others address space. As a result, there is no way for the different MPI threads to share the same FFIO cache. By default, each thread defines a separate FFIO cache based on the parameters defined by FF_IO_OPTS.

Having each MPI thread define a separate FFIO cache based on a single environment variable (FF_IO_OPTS) can waste a lot of memory. Fortunately, FFIO provides a mechanism that allows the user to specify a different FFIO cache for each MPI thread via the following environment variables:

```
setenv FF_IO_OPTS_RANK0 'result*(eie.direct.mbytes:4096:512:6:1:1:0)'
setenv FF_IO_OPTS_RANK2 'input*(eie.direct.mbytes:2048:64:6:1:1:0)'
...
setenv FF_IO_OPTS_RANKN-1 ...
```

Each rank environment variable is set using the exact same syntax as FF_IO_OPTS and each defines a distinct cache for the corresponding MPI rank. If the cache is designated shared, all files within the same ranking thread will use the same cache. FFIO works with SGI MPI, HP MPI, and LAM MPI. In order to work with MPI applications, FFIO needs to determine the rank of callers by invoking the mpi_comm_rank() MPI library routine. Therefore, FFIO needs to determine the location of the MPI library used by the application. This is accomplished by having the user set one, and only one, of the following environment variables:

```
setenv SGI_MPI /usr/lib
or
setenv LAM_MPI *see below
or
setenv HP_MPI *see below
```
*LAM and HP MPIs are usually distributed via a third party application. The precise paths to the LAM and the HP MPI libraries are application dependent. Please refer to the application installation guide to find the correct path.

In order to use the rank functionality, both the MPI and FF_IO_OPTS_RANK0 environment variables must be set. If either variable is not set, then the MPI threads all use FF_IO_OPTS. If both the MPI and the FF_IO_OPTS_RANK0 variables are defined but, for example, FF_IO_OPTS_RANK2 is undefined, all rank 2 files would generate a no match with FFIO. This means that none of the rank 2 files would be cached by FFIO. In this case, the software does not default to FF_IO_OPTS.

Fortran and C/C++ applications that use the pthreads interface will create threads that share the same address space. These threads can all make use of the single FFIO cache defined by FF_IO_OPTS.

**Application Examples**

FFIO has been deployed successfully with several HPC applications such as Nastran and Abaqus. In a recent customer benchmark, an eight-way Abaqus throughput job ran approximately twice as fast when FFIO was used. The FFIO cache used 16 megabyte pages (that is, page_size = 4096) and the cache size was 8.0 gigabytes. As a rule of thumb, it was determined that setting the FFIO cache size to roughly 10-15% of the disk space required by Abaqus yielded reasonable I/O performance. For this benchmark, the FF_IO_OPTS environment variable was defined by:

```
```

For the MPI version of Abaqus, different caches were specified for each MPI rank, as follows:

```
```

```
```


**Event Tracing**

By specifying the `.trace` option as part of the event parameter the user can enable the event tracing feature in FFIO, as follows:


This option generates files of the form `ffio.events.pid` for each process that is part of the application. By default, event files are placed in `/tmp` but this destination can be changed by setting the `FFIO_TMPDIR` environment variable. These files contain time stamped events for files using the FFIO cache and can be used to trace I/O activity (for example, I/O sizes and offsets).

**System Information and Issues**

Applications written in C, C++, and Fortran are supported. C and C++ applications can be built with either the Intel or gcc compiler. Only Fortran codes built with the Intel compiler will work.

The following restrictions on FFIO must also be observed:

- The FFIO implementation of `pread/pwrite` is not correct (the file offset advances).
- Do not use FFIO to do I/O on a socket.
- Do not link your application with the `librt` asynchronous I/O library.
- Calls that operate on files in `/proc`, `/etc`, and `/dev` are not intercepted by FFIO.
- Calls that operate on `stdin`, `stdout`, and `stderr` are not intercepted by FFIO.
- FFIO is not intended for generic I/O applications such as `vi`, `cp`, or `mv`, and so on.
Chapter 8

I/O Tuning

This chapter describes tuning information that you can use to improve I/O throughput and latency.

Application Placement and I/O Resources

It is useful to place an application on the same node as its I/O resource. For graphics applications, for example, this can improve performance up to 30 percent.

For example, for an SGI UV system with the following devices:

```bash
# gftopology

Serial number: UV-00000021
Partition number: 0
8 Blades
248 CPUs
283.70 Gb Memory Total
5 I/O Risers

<table>
<thead>
<tr>
<th>Blade Location</th>
<th>NASID</th>
<th>PCI Address</th>
<th>X Server Display</th>
<th>Device</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 r001i01b08</td>
<td>0</td>
<td>0000:05:00.0</td>
<td>-</td>
<td>Matrox Pilot</td>
</tr>
<tr>
<td>4 r001i01b12</td>
<td>8</td>
<td>0001:02:01.0</td>
<td>-</td>
<td>SGI Scalable Graphics Capture</td>
</tr>
<tr>
<td>6 r001i01b14</td>
<td>12</td>
<td>0003:07:00.0</td>
<td>Layout0.0</td>
<td>nVidia Quadro FX 5800</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0003:08:00.0</td>
<td>Layout0.1</td>
<td>nVidia Quadro FX 5800</td>
</tr>
<tr>
<td>7 r001i01b15</td>
<td>14</td>
<td>0004:03:00.0</td>
<td>Layout0.2</td>
<td>nVidia Quadro FX 5800</td>
</tr>
</tbody>
</table>
```

For example, to run an OpenGL graphics program, such as glxgears(1), on the third graphics processing unit using numactl(8), type the following command:

```bash
% numactl -N 14 -m 14 /usr/bin/glxgears -display :0.2
```

This example assumes the X server was started with :0 == Layout0.

The --N parameter specifies to run the command on node 14. The -m parameter specifies to allocate memory only from node 14.
You could also use the `dplace(1)` command to place the application, see "dplace Command" on page 44.

## Layout of Filesystems and XVM for Multiple RAIDs

There can be latency spikes in response from a RAID and such a spikes can in effect slow down all of the RAIDs as one I/O completion waits for all of the striped pieces to complete.

These latency spikes impact on throughput may be to stall all the I/O or to delay a few I/Os while others continue. It depends on how the I/O is striped across the devices. If the volumes are constructed as stripes to span all devices, and the I/Os are sized to be full stripes, the I/Os will stall, since every I/O has to touch every device. If the I/Os can be completed by touching a subset of the devices, then those that do not touch a high latency device can continue at full speed, while the stalled I/Os can complete and catch up later.

In large storage configurations, it is possible to lay out the volumes to maximize the opportunity for the I/Os to proceed in parallel, masking most of the effect of a few instances of high latency.

There are at least three classes of events that cause high latency I/O operations, as follows:

- Transient disk delays - one disk pauses
- Slow disks
- Transient RAID controller delays

The first two events affect a single logical unit number (LUN). The third event affects all the LUNs on a controller. The first and third events appear to happen at random. The second event is repeatable.
Chapter 9

Suggested Shortcuts and Workarounds

This chapter contains suggested workarounds and shortcuts that you can use on your SGI system. It covers the following topics:

- "Determining Process Placement" on page 99
- "Resetting System Limits" on page 106
- "Linux Shared Memory Accounting" on page 112
- "OFED Tuning Requirements for SHMEM" on page 113
- "Setting Java Environment Variables" on page 114

Determining Process Placement

This section describes methods that can be used to determine where different processes are running. This can help you understand your application structure and help you decide if there are obvious placement issues.

There are some setup steps to follow before determining process placement (note that all examples use the C shell):

1. Set up an alias as in this example, changing guest to your username:

   ```
   % alias pu "ps -edaf|grep guest"
   % pu
   
   The pu command shows current processes.
   ```

2. Create the .toprc preferences file in your login directory to set the appropriate top options. If you prefer to use the top defaults, delete the .toprc file.

   ```
   % cat <<EOF>> $HOME/.toprc
   YEAbcDgHIjklMnoTP|qrsuzVF
   2mlt
   EOF
   
   ```

3. Inspect all processes and determine which CPU is in use and create an alias file for this procedure. The CPU number is shown in the first column of the top output:
% top -b -n 1 | sort -n | more
% alias topl "top -b -n 1 | sort -n "

Use the following variation to produce output with column headings:
% alias topl "top -b -n 1 | head -4 | tail -1; top -b -n 1 | sort -n"

4. View your files (replacing guest with your username):
% top -b -n 1 | sort -n | grep guest
Use the following variation to produce output with column headings:
% top -b -n 1 | head -4 | tail -1; top -b -n 1 | sort -n grep guest

**Example Using pthreads**

The following example demonstrates a simple usage with a program name of th. It sets the number of desired OpenMP threads and runs the program. Notice the process hierarchy as shown by the PID and the PPID columns. The command usage is the following, where \( n \) is the number of threads:

% th \( n \)
%
% pu

<table>
<thead>
<tr>
<th>UID</th>
<th>PID</th>
<th>PPID</th>
<th>C</th>
<th>STIME</th>
<th>TTY</th>
<th>TIME</th>
<th>CMD</th>
</tr>
</thead>
<tbody>
<tr>
<td>root</td>
<td>13784</td>
<td>13779</td>
<td>0</td>
<td>12:41</td>
<td>pts/3</td>
<td>0:00</td>
<td>login --</td>
</tr>
<tr>
<td>guest1</td>
<td>13785</td>
<td>13784</td>
<td>0</td>
<td>12:41</td>
<td>pts/3</td>
<td>0:00</td>
<td>-csh</td>
</tr>
<tr>
<td>guest1</td>
<td>15062</td>
<td>13785</td>
<td>0</td>
<td>15:23</td>
<td>pts/3</td>
<td>0:00</td>
<td>th 4</td>
</tr>
<tr>
<td>guest1</td>
<td>15063</td>
<td>15062</td>
<td>0</td>
<td>15:23</td>
<td>pts/3</td>
<td>0:00</td>
<td>th 4</td>
</tr>
<tr>
<td>guest1</td>
<td>15064</td>
<td>15063</td>
<td>99</td>
<td>15:23</td>
<td>pts/3</td>
<td>0:01</td>
<td>th 4</td>
</tr>
<tr>
<td>guest1</td>
<td>15065</td>
<td>15063</td>
<td>99</td>
<td>15:23</td>
<td>pts/3</td>
<td>0:01</td>
<td>th 4</td>
</tr>
<tr>
<td>guest1</td>
<td>15066</td>
<td>15063</td>
<td>99</td>
<td>15:23</td>
<td>pts/3</td>
<td>0:01</td>
<td>th 4</td>
</tr>
<tr>
<td>guest1</td>
<td>15067</td>
<td>15063</td>
<td>99</td>
<td>15:23</td>
<td>pts/3</td>
<td>0:01</td>
<td>th 4</td>
</tr>
<tr>
<td>guest1</td>
<td>15068</td>
<td>13857</td>
<td>0</td>
<td>15:23</td>
<td>pts/5</td>
<td>0:00</td>
<td>ps -aef</td>
</tr>
<tr>
<td>guest1</td>
<td>15069</td>
<td>13857</td>
<td>0</td>
<td>15:23</td>
<td>pts/5</td>
<td>0:00</td>
<td>grep guest1</td>
</tr>
</tbody>
</table>
%

% top -b -n 1 | sort -n | grep guest1

<table>
<thead>
<tr>
<th>LC</th>
<th>%CPU</th>
<th>PID</th>
<th>USER</th>
<th>PRI</th>
<th>NI</th>
<th>SIZE</th>
<th>RSS</th>
<th>SHARE</th>
<th>STAT</th>
<th>%MEM</th>
<th>TIME</th>
<th>COMMAND</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>0.0</td>
<td>15072</td>
<td>guest1</td>
<td>16</td>
<td>0</td>
<td>3488</td>
<td>1536</td>
<td>3328</td>
<td>S</td>
<td>0.0</td>
<td>0:00</td>
<td>grep</td>
</tr>
</tbody>
</table>
Now skip the Main and daemon processes and place the rest:

% /usr/bin/dplace -s 2 -c 4-7 th 4
% pu

UID    PID     PPID  C   STIME TTY TIME CMD
root   13784  13779  0  12:41 pts/3 00:00:00 login --
guest1 13785  13784  0  12:41 pts/3 00:00:00 -csh
guest1 15083  15083  0 15:25 pts/3 00:00:00 th 4
guest1 15084  15083  0 15:25 pts/3 00:00:00 th 4
guest1 15085  15084  99 15:25 pts/3 00:00:19 th 4
guest1 15086  15084  99 15:25 pts/3 00:00:19 th 4
guest1 15087  15084  99 15:25 pts/3 00:00:19 th 4
guest1 15088  15084  99 15:25 pts/3 00:00:19 th 4
guest1 15091  13857  0 15:25 pts/5 00:00:00 ps -aef
guest1 15092  13857  0 15:25 pts/5 00:00:00 grep guest1

% top -b -n 1 | sort -n | grep guest1

LC  %CPU  PID USER  PRI  NI  SIZE  RSS  SHARE  STAT %MEM TIME COMMAND
4  99.9  15085 guest1  25  0 15856 2096  6496  R  0.0  0:24 th
5  99.8  15086 guest1  25  0 15856 2096  6496  R  0.0  0:24 th
6  99.9  15087 guest1  25  0 15856 2096  6496  R  0.0  0:24 th
7  99.9  15088 guest1  25  0 15856 2096  6496  R  0.0  0:24 th
8  0.0  15095 guest1  16  0  3488 1536  3328  S  0.0  0:00 grep
12  0.0  15091 guest1  15  0  5872 3664  4592  S  0.0  0:00 csh
12  0.0  15083 guest1  16  0 15856 2096  6496  S  0.0  0:00 th
12  0.0  15084 guest1  15  0 15856 2096  6496  S  0.0  0:00 th
Example Using OpenMP

The following example demonstrates a simple OpenMP usage with a program name of md. Set the desired number of OpenMP threads and run the program, as shown below:

```bash
% alias pu "ps -edaf | grep guest1
% setenv OMP_NUM_THREADS 4
% md
```

The following output is created:

```bash
% pu
```

<table>
<thead>
<tr>
<th>UID</th>
<th>PID</th>
<th>PPID</th>
<th>C</th>
<th>STIME</th>
<th>TTY</th>
<th>TIME</th>
<th>CMD</th>
</tr>
</thead>
<tbody>
<tr>
<td>root</td>
<td>21550</td>
<td>21535</td>
<td>0</td>
<td>21:48</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>login -- guest1</td>
</tr>
<tr>
<td>guest1</td>
<td>21551</td>
<td>21550</td>
<td>0</td>
<td>21:48</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>-csh</td>
</tr>
<tr>
<td>guest1</td>
<td>22183</td>
<td>21551</td>
<td>77</td>
<td>22:39</td>
<td>pts/0</td>
<td>00:00:03</td>
<td>md &lt;-- parent / main</td>
</tr>
<tr>
<td>guest1</td>
<td>22184</td>
<td>22183</td>
<td>0</td>
<td>22:39</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>md &lt;-- daemon</td>
</tr>
<tr>
<td>guest1</td>
<td>22185</td>
<td>22184</td>
<td>0</td>
<td>22:39</td>
<td>pts/0</td>
<td>00:00:00</td>
<td>md &lt;-- daemon helper</td>
</tr>
<tr>
<td>guest1</td>
<td>22186</td>
<td>22184</td>
<td>99</td>
<td>22:39</td>
<td>pts/0</td>
<td>00:00:03</td>
<td>md &lt;-- thread 1</td>
</tr>
<tr>
<td>guest1</td>
<td>22187</td>
<td>22184</td>
<td>94</td>
<td>22:39</td>
<td>pts/0</td>
<td>00:00:03</td>
<td>md &lt;-- thread 2</td>
</tr>
<tr>
<td>guest1</td>
<td>22188</td>
<td>22184</td>
<td>85</td>
<td>22:39</td>
<td>pts/0</td>
<td>00:00:03</td>
<td>md &lt;-- thread 3</td>
</tr>
<tr>
<td>guest1</td>
<td>22189</td>
<td>21956</td>
<td>0</td>
<td>22:39</td>
<td>pts/1</td>
<td>00:00:00</td>
<td>ps -aef</td>
</tr>
<tr>
<td>guest1</td>
<td>22190</td>
<td>21956</td>
<td>0</td>
<td>22:39</td>
<td>pts/1</td>
<td>00:00:00</td>
<td>grep guest1</td>
</tr>
</tbody>
</table>

```bash
% top -b -n 1 | sort -n | grep guest1
```

<table>
<thead>
<tr>
<th>LC</th>
<th>%CPU</th>
<th>PID USER</th>
<th>PRI</th>
<th>NI</th>
<th>SIZE</th>
<th>RSS</th>
<th>SHARE</th>
<th>STAT</th>
<th>%MEM</th>
<th>TIME</th>
<th>COMMAND</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0.0</td>
<td>22192 guest1</td>
<td>16</td>
<td>0</td>
<td>70048 1600 69840 S</td>
<td>0.0</td>
<td>0:00</td>
<td>sort</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>0.0</td>
<td>22193 guest1</td>
<td>16</td>
<td>0</td>
<td>3488 1536 3328 S</td>
<td>0.0</td>
<td>0:00</td>
<td>grep</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>1.6</td>
<td>22191 guest1</td>
<td>15</td>
<td>0</td>
<td>5056 2832 4288 R</td>
<td>0.0</td>
<td>0:00</td>
<td>top</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>98.0</td>
<td>22186 guest1</td>
<td>26</td>
<td>0</td>
<td>26432 2704 4272 R</td>
<td>0.0</td>
<td>0:11</td>
<td>md</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>0.0</td>
<td>22185 guest1</td>
<td>15</td>
<td>0</td>
<td>26432 2704 4272 S</td>
<td>0.0</td>
<td>0:00</td>
<td>md</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>87.6</td>
<td>22188 guest1</td>
<td>25</td>
<td>0</td>
<td>26432 2704 4272 R</td>
<td>0.0</td>
<td>0:10</td>
<td>md</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>0.0</td>
<td>21551 guest1</td>
<td>15</td>
<td>0</td>
<td>5872 3648 4560 S</td>
<td>0.0</td>
<td>0:00</td>
<td>csh</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>0.0</td>
<td>22184 guest1</td>
<td>15</td>
<td>0</td>
<td>26432 2704 4272 S</td>
<td>0.0</td>
<td>0:00</td>
<td>md</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>99.9</td>
<td>22183 guest1</td>
<td>39</td>
<td>0</td>
<td>26432 2704 4272 R</td>
<td>0.0</td>
<td>0:11</td>
<td>md</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
From the notation on the right of the pu list, you can see the \(-x 6\) pattern.

place 1, skip 2 of them, place 3 more \([ 0 1 1 0 0 0 ]\)
now, reverse the bit order and create the dplace \(-x\) mask
\([ 0 0 1 1 0 ] \rightarrow [ 0x06 ] \rightarrow \text{decimal 6}\)

dplace does not currently process hex notation for this bit mask)

The following example confirms that a simple dplace placement works correctly:

\%
setenv OMP_NUM_THREADS 4
%
/usr/bin/dplace \(-x 6\) \(-c\) 4-7 md
%

\%
pu
UID PID PPID C STIME TTY TIME CMD
root 21550 21535 0 21:48 pts/0 00:00:00 login -- guest1
guest1 21551 21550 0 21:48 pts/0 00:00:00 -csh
guest1 22219 21551 93 22:45 pts/0 00:00:05 md
guest1 22220 22219 0 22:45 pts/0 00:00:00 md
guest1 22221 22220 0 22:45 pts/0 00:00:00 md
guest1 22222 22220 93 22:45 pts/0 00:00:05 md
guest1 22223 22220 93 22:45 pts/0 00:00:05 md
guest1 22224 22220 90 22:45 pts/0 00:00:05 md
guest1 22225 21956 0 22:45 pts/1 00:00:00 ps -aef
guest1 22226 21956 0 22:45 pts/1 00:00:00 grep guest1
%

\%
top \(-b\) \(-n\) 1 | sort \(-n\) | grep guest1
%

LC %CPU PID USER PRI NI SIZE RSS SHARE STAT %MEM TIME COMMAND
2 0.0 22228 guest1 16 0 70048 1600 69840 S 0.0 0:00 sort
2 0.0 22229 guest1 16 0 3488 1536 3328 S 0.0 0:00 grep
2 1.6 22227 guest1 15 0 5056 2832 4288 R 0.0 0:00 top
4 0.0 22220 guest1 15 0 28496 2736 21728 S 0.0 0:00 md
4 99.9 22219 guest1 39 0 28496 2736 21728 R 0.0 0:12 md
5 99.9 22222 guest1 25 0 28496 2736 21728 R 0.0 0:11 md
6 99.9 22223 guest1 39 0 28496 2736 21728 R 0.0 0:11 md
7 99.9 22224 guest1 39 0 28496 2736 21728 R 0.0 0:11 md
9 0.0 21551 guest1 15 0 5872 3648 4560 S 0.0 0:00 csh
15 0.0 22221 guest1 15 0 28496 2736 21728 S 0.0 0:00 md
Combination Example (MPI and OpenMP)

For this example, explicit placement using the `dplace -e -c` command is used to achieve the desired placement. If an `x` is used in one of the CPU positions, `dplace` does not explicitly place that process.

If running without a cpuset, the `x` processes run on any available CPU.

If running with a cpuset, you have to renumber the CPU numbers to refer to “logical” CPUs (0 ... n) within the cpuset, regardless of which physical CPUs are in the cpuset. When running in a cpuset, the unplaced processes are constrained to the set of CPUs within the cpuset.

For information about cpusets, see the *SGI Cpuset Software Guide*.

The following example shows a hybrid MPI and OpenMP job with two MPI processes, each with two OpenMP threads and no cpusets:

```bash
% setenv OMP_NUM_THREADS 2
% efc -O2 -o hybrid hybrid.f -lmpi -openmp
% mpirun -v -np 2 /usr/bin/dplace -e -c x,8,9,x,x,x,x,10,11 hybrid
-------------------------
# if using cpusets ...
-------------------------
# we need to reorder cpus to logical within the 8-15 set [0-7]
% cpuset -q omp -A mpirun -v -np 2 /usr/bin/dplace -e -c x,0,1,x,x,x,x,2,3,4,5,6,7 hybrid
# We need a table of options for these pairs. "x" means don’t care. See the dplace man page for more info about the -e option.
# examples at end

-mpirun -v -np 2 /usr/bin/dplace -e -c <as shown> a.out
--- --------------- ---------------------------------------
 2 2 x,0,1,x,x,x,x,2,3
 2 3 x,0,1,x,x,x,x,2,3,4,5
 2 4 x,0,1,x,x,x,x,2,3,4,5,6,7
 4 2 x,0,1,2,3,x,x,x,x,x,x,x,x,4,5,6,7
 4 3 x,0,1,2,3,x,x,x,x,x,x,x,x,x,4,5,6,7,8,9,10,11
```

9: Suggested Shortcuts and Workarounds
Notes: 0 <- 1 -> <- 2 -> <- 3 -> <------ 4

Notes:
0. mpi daemon process
1. mpi child procs, one per np
2. omp daemon procs, one per np
3. omp daemon helper procs, one per np
4. omp thread procs, (OMP_NUM_THREADS - 1) per np

# Example - -np 2 and OMP_NUM_THREADS 2

% setenv OMP_NUM_THREADS 2
% efc -O2 -o hybrid hybrid.f -lmpi -openmp
% mpirun -v -np 2 /usr/bin/dplace -e -c x,8,9,x,x,x,x,10,11 hybrid
% pu

UID       PID      PPID   C   STIME TTY      TIME CMD
root  21550    21535     0  00:00:00 pts/0 login -- guest1
guest1 21551  21550  0  00:32 pts/0  00:00:00 mpirun -v -np 2

/usr/bin/dplace
guest1 23394  23391     2  00:32 pts/0  00:00:00 hybrid <-- mpi daemon
guest1 23401  23394    99  00:32 pts/0  00:00:03 hybrid <-- mpi child 1
guest1 23402  23394    99  00:32 pts/0  00:00:03 hybrid <-- mpi child 2
guest1 23403  23402    0  00:32 pts/0  00:00:00 hybrid <-- omp daemon 2
guest1 23404  23401    0  00:32 pts/0  00:00:00 hybrid <-- omp daemon 1
guest1 23405  23404    0  00:32 pts/0  00:00:00 hybrid <-- omp daemon hlpr 1
guest1 23406  23403    0  00:32 pts/0  00:00:00 hybrid <-- omp daemon hlpr 2
guest1 23407  23403    99  00:32 pts/0  00:00:03 hybrid <-- omp thread 2-1
guest1 23408  23404    99  00:32 pts/0  00:00:03 hybrid <-- omp thread 1-1
guest1 23409  21956    0  00:32 pts/1  00:00:00 ps -aef
guest1 23410  21956    0  00:32 pts/1  00:00:00 grep guest1
% top -b -n 1 | sort -n | grep guest1

<table>
<thead>
<tr>
<th>LC</th>
<th>%CPU</th>
<th>PID</th>
<th>USER</th>
<th>PRI</th>
<th>NI</th>
<th>SIZE</th>
<th>RSS</th>
<th>SHARE</th>
<th>STAT</th>
<th>%MEM</th>
<th>TIME</th>
<th>COMMAND</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.0</td>
<td>21551</td>
<td>guest1</td>
<td>15</td>
<td>0</td>
<td>5904</td>
<td>3712</td>
<td>4592</td>
<td>S</td>
<td>0.0</td>
<td>0:00</td>
<td>csh</td>
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<td>0</td>
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<td>0</td>
<td>883M</td>
<td>9456</td>
<td>882M</td>
<td>S</td>
<td>0.1</td>
<td>0:00</td>
<td>hybrid</td>
</tr>
<tr>
<td>4</td>
<td>0.0</td>
<td>21956</td>
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<td>5856</td>
<td>3616</td>
<td>5664</td>
<td>S</td>
<td>0.0</td>
<td>0:00</td>
<td>csh</td>
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<td>1600</td>
<td>69840</td>
<td>S</td>
<td>0.0</td>
<td>0:00</td>
<td>sort</td>
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<td>guest1</td>
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<td>0</td>
<td>5056</td>
<td>2832</td>
<td>4288</td>
<td>R</td>
<td>0.0</td>
<td>0:00</td>
<td>top</td>
</tr>
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<td>guest1</td>
<td>16</td>
<td>0</td>
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<td>1536</td>
<td>3328</td>
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<td>0.0</td>
<td>0:00</td>
<td>grep</td>
</tr>
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<td>5840</td>
<td>3584</td>
<td>5648</td>
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<td>894M</td>
<td>10M</td>
<td>889M</td>
<td>S</td>
<td>0.1</td>
<td>0:00</td>
<td>hybrid</td>
</tr>
<tr>
<td>8</td>
<td>99.9</td>
<td>23401</td>
<td>guest1</td>
<td>39</td>
<td>0</td>
<td>894M</td>
<td>10M</td>
<td>889M</td>
<td>R</td>
<td>0.1</td>
<td>0:09</td>
<td>hybrid</td>
</tr>
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<td>0</td>
<td>894M</td>
<td>10M</td>
<td>894M</td>
<td>S</td>
<td>0.1</td>
<td>0:00</td>
<td>hybrid</td>
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<td>99.9</td>
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<td>guest1</td>
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<td>0</td>
<td>894M</td>
<td>10M</td>
<td>894M</td>
<td>R</td>
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<td>0:09</td>
<td>hybrid</td>
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<td>0:09</td>
<td>hybrid</td>
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<td>R</td>
<td>0.1</td>
<td>0:09</td>
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<tr>
<td>12</td>
<td>0.0</td>
<td>23391</td>
<td>guest1</td>
<td>15</td>
<td>0</td>
<td>5072</td>
<td>2928</td>
<td>4400</td>
<td>S</td>
<td>0.0</td>
<td>0:00</td>
<td>mpirun</td>
</tr>
<tr>
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<td>23406</td>
<td>guest1</td>
<td>15</td>
<td>0</td>
<td>894M</td>
<td>10M</td>
<td>894M</td>
<td>S</td>
<td>0.1</td>
<td>0:00</td>
<td>hybrid</td>
</tr>
<tr>
<td>14</td>
<td>0.0</td>
<td>23405</td>
<td>guest1</td>
<td>15</td>
<td>0</td>
<td>894M</td>
<td>10M</td>
<td>894M</td>
<td>S</td>
<td>0.1</td>
<td>0:00</td>
<td>hybrid</td>
</tr>
</tbody>
</table>

**Resetting System Limits**

To regulate these limits on a per-user basis for applications that do not rely on `limit.h`, the `limits.conf` file can be modified. System limits that can be modified include maximum file size, maximum number of open files, maximum stack size, and so on. You can view this file as, follows:

[user@machine user]# cat /etc/security/limits.conf
# /etc/security/limits.conf
#
# Each line describes a limit for a user in the form:
# #
# Where:
# can be:
# - an user name
# - a group name, with @group syntax
# - the wildcard *, for default entry
# can have the two values:
# - "soft" for enforcing the soft limits
# - "hard" for enforcing hard limits
#
# can be one of the following:
# - core - limits the core file size (KB)
# - data - max data size (KB)
# - fsize - maximum filesize (KB)
# - memlock - max locked-in-memory address space (KB)
# - nofile - max number of open files
# - rss - max resident set size (KB)
# - stack - max stack size (KB)
# - cpu - max CPU time (MIN)
# - nproc - max number of processes
# - as - address space limit
# - maxlogins - max number of logins for this user
# - priority - the priority to run user process with
# - locks - max number of file locks the user can hold
#
#
##
*
soft core 0
*
hard rss 10000
@@student hard nproc 20
@@faculty soft nproc 20
@@faculty hard nproc 50
ftp hard nproc 0
@@student - maxlogins 4

# End of file

For instructions on how to change these limits, see "Resetting the File Limit Resource Default" on page 107.

Resetting the File Limit Resource Default

Several large user applications use the value set in the limit.h file as a hard limit on file descriptors and that value is noted at compile time. Therefore, some applications may need to be recompiled in order to take advantage of the SGI system hardware.

To regulate these limits on a per-user basis for applications that do not rely on limit.h, the limits.conf file can be modified. This allows the administrator to
set the allowed number of open files per user and per group. This also requires a one-line change to the `/etc/pam.d/login` file.

Follow this procedure to execute these changes:

1. Add the following line to `/etc/pam.d/login`:

   ```
   session required /lib/security/pam_limits.so
   ```

2. Add the following line to `/etc/security/limits.conf`, where `username` is the user’s login and `limit` is the new value for the file limit resource:

   ```
   [username] hard nofile [limit]
   ```

The following command shows the new limit:

```
ulimit -H -n
```

Because of the large number of file descriptors that some applications require, such as MPI jobs, you might need to increase the system-wide limit on the number of open files on your SGI system. The default value for the file limit resource is 1024. The default 1024 file descriptors allows for approximately 199 MPI processes per host. You can increase the file descriptor value to 8196 to allow for more than 512 MPI processes per host by adding the following lines to the `/etc/security/limits.conf` file:

```
* soft nofile 8196
* hard nofile 8196
```

The `ulimit -a` command displays all limits, as follows:

```
sys:~ # ulimit -a
  core file size (blocks, -c) 1
  data seg size (kbytes, -d) unlimited
  scheduling priority (-e) 0
  file size (blocks, -f) unlimited
  pending signals (-i) 511876
  max locked memory (kbytes, -l) 64
  max memory size (kbytes, -m) 55709764
  open files (-n) 1024
  pipe size (512 bytes, -p) 8
  POSIX message queues (bytes, -q) 819200
  real-time priority (-r) 0
  stack size (kbytes, -s) 8192
```
Resetting the Default Stack Size

Some applications will not run well on an SGI system with a small stack size. To set a higher stack limit, follow the instructions in "Resetting the File Limit Resource Default" on page 107 and add the following lines to the `/etc/security/limits.conf` file:

* soft stack 300000
* hard stack unlimited

This sets a soft stack size limit of 300000 KB and an unlimited hard stack size for all users (and all processes).

Another method that does not require root privilege relies on the fact that many MPI implementation use ssh, rsh, or some sort of login shell to start the MPI rank processes. If you merely need to bump up the soft limit, you can modify your shell’s startup script. For example, if your login shell is `bash` then add something like the following to your `.bashrc` file:

```
% ulimit -s 300000
```

Note that SGI MPT MPI allows you to set your stack size limit larger with the `ulimit` or `limit` shell command before launching an MPI program with `mpirun(1)` or `mpiexec_mpt(1)`. MPT will propagate the stack limit setting to all MPI processes in the job.

For more information on default settings, also see "Resetting the File Limit Resource Default" on page 107.

Avoiding Segmentation Faults

The default stack size in the Linux operating system is 8MB (8192 kbytes). This value needs to be increased to avoid "Segmentation Faults" errors.

You can use the `ulimit -a` command to view the stack size, as follows:

```
$ uv44-sys:~ # ulimit -a
  core file size    (blocks, -c) unlimited
```
data seg size (kbytes, -d) unlimited

file size (blocks, -f) unlimited

pending signals (-i) 204800

max locked memory (kbytes, -l) unlimited

max memory size (kbytes, -m) unlimited

open files (-n) 16384

pipe size (512 bytes, -p) 8

POSIX message queues (bytes, -q) 819200

stack size (kbytes, -s) 8192

cpu time (seconds, -t) unlimited

max user processes (-u) 204800

virtual memory (kbytes, -v) unlimited

file locks (-x) unlimited

To change the value, perform a command similar to the following:

uv44-sys:~ # ulimit -s 300000

There is a similar variable for OpenMP programs. If you get a segmentation fault right away while running a program parallelized with OpenMP, a good idea is to increase the KMP_STACKSIZE to a larger size. The default size in Intel Compilers is 4MB.

For example, to increase it to 64MB in csh shell perform the following:

setenv KMP_STACKSIZE 64M

in bash:

export KMP_STACKSIZE=64M
Resetting Virtual Memory Size

The virtual memory parameter `vmemoryuse` determines the amount of virtual memory available to your application. If you are running with csh, use csh commands, such as, the following:

```
limit
limit vmemoryuse 7128960
limit vmemoryuse unlimited
```

The following MPI program fails with a memory-mapping error because of a virtual memory parameter `vmemoryuse` value set too low:

```
% limit vmemoryuse 7128960
% mpirun -v -np 4 ./program
```

```
MPI: libxmpi.so 'SGI MPI 4.9 MPT 1.14 07/18/06 08:43:15'
MPI: libmpi.so 'SGI MPI 4.9 MPT 1.14 07/18/06 08:41:05'
MPI: MPI_MSGS_MAX = 524288
MPI: MPI_BUFS_PER_PROC= 32
mmap failed (memmap_base) for 504972 pages (8273461248 bytes) Killed n
```

The program now succeeds when virtual memory is unlimited:

```
% limit vmemoryuse unlimited
% mpirun -v -np 4 ./program
```

```
MPI: libxmpi.so 'SGI MPI 4.9 MPT 1.14 07/18/06 08:43:15'
MPI: libmpi.so 'SGI MPI 4.9 MPT 1.14 07/18/06 08:41:05'
MPI: MPI_MSGS_MAX = 524288
MPI: MPI_BUFS_PER_PROC= 32
HELLO WORLD from Processor 0
HELLO WORLD from Processor 2
HELLO WORLD from Processor 1
HELLO WORLD from Processor 3
```
If you are running with bash, use bash commands, such as, the following:

```
ulimit -a
ulimit -v 7128960
ulimit -v unlimited
```

**Linux Shared Memory Accounting**

The Linux operating system does not calculate memory utilization in a manner that is useful for certain applications in situations where regions are shared among multiple processes. This can lead to over-reporting of memory and to processes being killed by schedulers erroneously detecting memory quota violation.

The `get_weighted_memory_size` function weighs shared memory regions by the number of processes using the regions. Thus, if 100 processes are each sharing a total of 10GB of memory, the weighted memory calculation shows 100MB of memory shared per process, rather than 10GB for each process.

Because this function applies mostly to applications with large shared-memory requirements, it is located in the SGI NUMA tools package and made available in the `libmemacct` library available from a new package called `memacct`. The library function makes a call to the `numatools` kernel module, which returns the weighted sum back to the library, and then returns back to the application.

The usage statement for the `memacct` call is, as follows:

```
cc ... -lmemacct
#include <sys/types.h>
extern int get_weighted_memory_size(pid_t pid);
```

The syntax of the `memacct` call is, as follows:

```
int *get_weighted_memory_size(pid_t pid);
```

Returns the weighted memory (RSS) size for a `pid`, in bytes. This weights the size of shared regions by the number of processes accessing it. Return -1 when an error occurs and set `errno`, as follows:

- **ESrch**  
  Process `pid` was not found.
- **ENosys**  
  The function is not implemented. Check if `numatools` kernel package is up-to-date.

Normally, the following errors should not occur:
ENOENT Can not open /proc/numatools device file.
EPERM No read permission on /proc/numatools device file.
ENOTTY Inappropriate ioctl operation on /proc/numatools device file.
EFAULT Invalid arguments. The ioctl() operation performed by the function failed with invalid arguments.

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

OFED Tuning Requirements for SHMEM

You can specify the maximum number of queue pairs (QPs) for SHMEM applications when run on large clusters over an OFED fabric, such as InfiniBand. 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

SHMEM codes use the InfiniBand RC protocol for communication between all pairs of processes in the parallel job, which requires a large number of QPs. The log_num_qp parameter defines the log₂ of the number of QPs. The following procedure explains how to specify the log_num_qp parameter.

Procedure 9-1 To specify the log_num_qp parameter

1. Log into one of the hosts upon which you installed the MPT software as the root user.
2. Use a text editor to open file /etc/modprobe.d/libmlx4.conf.
3. Add a line similar to the following to file /etc/modprobe.d/libmlx4.conf:
   
   options mlx4_core log_num_qp=21

   By default, the maximum number of queue pairs on RHEL platforms is $2^{18}$ (262144).

   By default, the maximum number of queue pairs on SLES platforms is $2^{17}$ (131072).

4. Save and close the file.
5. Repeat the preceding steps on other hosts.

**Setting Java Environment Variables**

When Java software starts, it checks the environment in which it is running and configures itself to fit, assuming that it owns the entire environment. The default for some Java implementations (for example, IBM J9 1.4.2) is to start a garbage collection (GC) thread for every CPU it sees. Other Java implementations use other algorithms to decide the number of GCs to start, but the number will generally be 0.5 to 1 times the number of CPUs. On a 1 or 2 socket system, that works out well enough. This strategy does not scale well to large core-count systems, however.

Java command line options allow you to control the number of GC threads that the Java virtual machine (JVM) will use. In many cases, a single GC thread is sufficient, as set in the examples above. In other cases, a larger number may be appropriate and can be set with the applicable environment variable or command line option. Properly tuning the number of GC threads for an application is an exercise in performance optimization, but a reasonable starting point is to use one GC thread per active worker thread.

*For Sun Java (now Oracle Java):*

-XX:ParallelGCThreads

*For IBM Java:*

-Xgcthreads

An example command line option:

```
java -XX:+UseParallelGC -XX:ParallelGCThreads=1
```

The system administrator may choose to implement the option to limit the number of GC threads to a reasonable value with an environment variable set in the global profile, for example the `/etc/profile.local` file so casual Java users can avoid difficulties.

Environment variable settings, are as follows:

*For Sun Java (now Oracle Java):*

`JAVA_OPTIONS="-XX:ParallelGCThreads=1"

*For IBM Java:*
IBM_JAVA_OPTIONS="-Xgcthreads1"
Using PerfSocket

This chapter includes the following topics:

- "About SGI PerfSocket" on page 117
- "Installing and Using PerfSocket" on page 117
- "About Security When Using PerfSocket" on page 120
- "Troubleshooting" on page 120

About SGI PerfSocket

The SGI PerfSocket feature improves an application’s TCP/IP communication within a host. The PerfSocket software intercepts local TCP/IP communication and routes the communication through shared memory, which eliminates much of the overhead incurred when communication data passes through the operating system kernel.

PerfSocket includes a system library, a daemon, a kernel module, and a wrapper command that enables PerfSocket functionality within specified processes. Only processes specifically run with the PerfSocket wrapper command are run with this feature.

SGI includes the PerfSocket technology in the SGI Accelerate™ product within the SGI Performance Suite. After you install SGI Accelerate, you can use the procedure in the following topic to install PerfSocket as an optional feature:

"Installing PerfSocket (Administrator Procedure)" on page 118.

Installing and Using PerfSocket

The SGI Accelerate installation process does not install PerfSocket. The following procedures explain how to install and use PerfSocket:

- "Installing PerfSocket (Administrator Procedure)" on page 118
- "Running an Application With PerfSocket" on page 119
Installing PerfSocket (Administrator Procedure)

When you install the SGI Performance Suite software, the installer does not install PerfSocket. You need to complete the procedure in this topic to install PerfSocket.

The PerfSocket RPM contains the PerfSocket libraries, applications, kernel modules, and man(1) pages. The installer writes the majority of PerfSocket’s files to the /opt/sgi/perfsocket directory.

The following procedure explains how to install PerfSocket and how to start the PerfSocket daemon on an SGI UV computer system.

Procedure 10-1 To install Perfsocket on an SGI UV system

1. Log in as root.

2. Install the PerfSocket software.

   This command sequence differs, depending on your platform, as follows:

   • On RHEL platforms, type the following command:

     `# yum install perfsocket`

     When RHEL displays the download size and displays the Is this ok [y/N]: prompt, type y and press Enter.

   • On SLES platforms, type the following commands:

     `# zypper refresh`
     `# zypper install perfsocket`

     When SLES displays the download size and displays the Continue? [y/n/?] (y): prompt, type y and press Enter.

3. Type the following command to turn on the PerfSocket service:

   `# chkconfig perfsocket on`

4. Type the following command to verify that the PerfSocket service is on:

   `# chkconfig --list | grep perfsocket`

   perfsocket 0:off 1:off 2:on 3:on 4:on 5:on 6:off
5. Type the following command to start the PerfSocket daemon without a reboot:

```
# service perfsocket start
Starting the PerfSocket daemon
```

6. (Optional) Type the following command to verify that the kernel module is loaded:

```
# lsmod | grep perfsock
perfsock 71297 10
```

7. (Optional) Type the following command to display the PerfSocket processes:

```
# ps ax | grep perfsocketd
10308 ? Ss 0:00 /opt/sgi/perfsocket/sbin/perfsocketd
10319 pts/0 S+ 0:00 grep perfsocketd
```

**Running an Application With PerfSocket**

You do not need to recompile an application in order to use PerfSocket. An application that runs with PerfSocket automatically detects whether the endpoints of its communication are also using PerfSocket. All applications that use any particular socket endpoint must be run with PerfSocket in order for PerfSocket to accelerate the communication.

The following procedure explains how to invoke PerfSocket to run with your application programs.

**Procedure 10-2** To run an application with PerfSocket

1. Type the following command to load the PerfSocket environment module:

```
$ module load perfsocket
```

   The previous command adds the PerfSocket wrapper command to your PATH variable and adds the PerfSocket libraries to your LD_LIBRARY_PATH.

2. For each command that you want to run with PerfSocket, prefix the command with the `perfsocket(1)` command.

   For example, if applications `a.out` and `b.out` communicate with TCP/IP, type the following commands to enable them to communicate through PerfSocket:

   ```
   $ perfsocket ./a.out &
   $ perfsocket ./b.out
   ```
For more information, see the `perfsocket(1)` and `perfsocketd(1)` man pages.

### About Security When Using PerfSocket

The PerfSocket daemon facilitates application communication under the following conditions:

- An application that uses PerfSocket connects to another application that also uses PerfSocket
- The applications that connect run on the same host
- The user ID is identical for both the connecting process and the receiving process

Only the user who owns the applications can read the shared memory structure used for communication. No additional copies of the data are made. If a process that uses PerfSocket calls `exec()`, all PerfSocket-enabled sockets are duplicated to `/dev/null`.

### Troubleshooting

If PerfSocket detects an unsupported condition, stop using PerfSocket.

SGI implemented and tested PerfSocket with the majority of socket and file I/O APIs. It is possible that you might attempt to use PerfSocket with a rarely used API or an unsupported usage pattern. If you encounter an unsupported condition, PerfSocket logs an error message and aborts the application.

PerfSocket writes its log messages to the system log, `/var/log/messages`. 
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