Programming the Cell Broadband Engine™ Architecture
Examples and Best Practices

- Understand and apply different programming models and strategies
- Make the most of SDK 3.0 debug and performance tools
- Use practical code development and porting examples

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Preface

In this IBM® Redbooks® publication, we provide an introduction to the Cell Broadband Engine™ (Cell/B.E.™) platform. We show detailed samples from real-world application development projects and provide tips and best practices for programming Cell/B.E. applications.

We also describe the content and packaging of the IBM Software Development Kit (SDK) version 3.0 for Multicore Acceleration. This SDK provides all the tools and resources that are necessary to build applications that run IBM BladeCenter® QS21 and QS20 blade servers. We show in-depth and real-world usage of the tools and resources found in the SDK. We also provide installation, configuration, and administration tips and best practices for the IBM BladeCenter QS21. In addition, we discuss the supporting software that is provided by IBM alphaWorks®.

This book was written for developers and programmers, IBM technical specialists, Business Partners, Clients, and the Cell/B.E. community to understand how to develop applications by using the Cell/B.E. SDK 3.0.

The team that wrote this book

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Introduction to the Cell Broadband Engine Architecture

The Cell Broadband Engine (Cell/B.E.) Architecture (CBEA) is a new class of multicore processors being brought to the consumer and business market. It has a radically different design than those offered by other consumer and business chip makers in the global market. This radical departure warrants a brief discussion of the Cell/B.E. hardware and software architecture.

There is also a brief discussion of the IBM Software Development Kit (SDK) for Multicore Acceleration from a content and packaging perspective. These discussions complement the in-depth content of the remaining chapters of this book.

Specifically, this part includes the following chapters:

- Chapter 1, “Cell Broadband Engine overview” on page 3
- Chapter 2, “IBM SDK for Multicore Acceleration” on page 17
Cell Broadband Engine overview

The Cell Broadband Engine (Cell/B.E.) processor is the first implementation of a new multiprocessor family that conforms to the Cell/B.E. Architecture (CBEA). The CBEA and the Cell/B.E. processor are the result of a collaboration between Sony, Toshiba, and IBM (known as STI), which formally began in early 2001. Although the Cell/B.E. processor is initially intended for applications in media-rich consumer-electronics devices, such as game consoles and high-definition televisions, the architecture has been designed to enable fundamental advances in processor performance. These advances are expected to support a broad range of applications in both commercial and scientific fields.

In this chapter, we discuss the following topics:

- 1.1, “Motivation” on page 4
- 1.2, “Scaling the three performance-limiting walls” on page 6
- 1.3, “Hardware environment” on page 8
- 1.4, “Programming environment” on page 11
1.1 Motivation

The CBEA has been designed to support a broad range of applications. The first implementation is a single-chip multiprocessor with nine processor elements that operate on a shared memory model, as shown in Figure 1-1. In this respect, the Cell/B.E. processor extends current trends in PC and server processors. The most distinguishing feature of the Cell/B.E. processor is that, although all processor elements can share or access all available memory, their function is specialized into two types:

- Power Processor Element (PPE)
- Synergistic Processor Element (SPE)

The Cell/B.E. processor has one PPE and eight SPEs. The PPE contains a 64-bit PowerPC® Architecture™ core. It complies with the 64-bit PowerPC Architecture and can run 32-bit and 64-bit operating systems and applications. The SPE is optimized for running compute-intensive single-instruction, multiple-data (SIMD) applications. It is not optimized for running an operating system.

The SPEs are independent processor elements, each running their own individual application programs or threads. Each SPE has full access to shared memory, including the memory-mapped I/O space implemented by multiple
direct memory access (DMA) units. There is a mutual dependence between the PPE and the SPEs. The SPEs depend on the PPE to run the operating system, and, in many cases, the top-level thread control for an application. The PPE depends on the SPEs to provide the bulk of the application performance.

The SPEs are designed to be programmed in high-level languages, such as C/C++. They support a rich instruction set that includes extensive SIMD functionality. However, like conventional processors with SIMD extensions, use of SIMD data types is preferred, not mandatory. For programming convenience, the PPE also supports the standard PowerPC Architecture instructions and the vector/SIMD multimedia extensions.

To an application programmer, the Cell/B.E. processor looks like a single core, dual-threaded processor with eight additional cores each having their own local storage. The PPE is more adept than the SPEs at control-intensive tasks and quicker at task switching. The SPEs are more adept at compute-intensive tasks and slower than the PPE at task switching. However, either processor element is capable of both types of functions. This specialization is a significant factor accounting for the order-of-magnitude improvement in peak computational performance and chip-area and power efficiency that the Cell/B.E. processor achieves over conventional PC processors.

The more significant difference between the SPE and PPE lies in how they access memory. The PPE accesses main storage (the effective address space) with load and store instructions that move data between main storage and a private register file, the contents of which may be cached. PPE memory access is like that of a conventional processor technology, which is found on conventional machines.

The SPEs, in contrast, access main storage with DMA commands that move data and instructions between main storage and a private local memory, called local storage (LS). An instruction-fetches and load and store instructions of an SPE access its private LS rather than shared main storage, and the LS has no associated cache. This three-level organization of storage (register file, LS, and main storage), with asynchronous DMA transfers between LS and main storage, is a radical break from conventional architecture and programming models. The organization explicitly parallelizes computation with the transfers of data and instructions that feed computation and store the results of computation in main storage.

One of the motivations for this radical change is that memory latency, measured in processor cycles, has gone up several hundredfold from about the years 1980 to 2000. The result is that application performance is, in most cases, limited by memory latency rather than peak compute capability or peak bandwidth.
When a sequential program on a conventional architecture performs a load instruction that misses in the caches, program execution can come to a halt for several hundred cycles. (Techniques such as hardware threading can attempt to hide these stalls, but it does not help single-threaded applications.) Compared to this penalty, the few cycles it takes to set up a DMA transfer for an SPE are a much better trade-off, especially considering the fact that the DMA controllers of each of the eight SPEs can have up to 16 DMA transfers in flight simultaneously. Anticipating DMA needs efficiently can provide just-in-time delivery of data that may reduce this stall or eliminate them entirely. Conventional processors, even with deep and costly speculation, manage to get, at best, a handful of independent memory accesses in flight.

One of the DMA transfer methods of the SPE supports a list, such as a scatter-gather list, of DMA transfers that is constructed in the local storage of an SPE. Therefore, the DMA controller of the SPE can process the list asynchronously while the SPE operates on previously transferred data. In several cases, this approach to accessing memory has led to application performance exceeding that of conventional processors by almost two orders of magnitude. This result is significantly more than you expect from the peak performance ratio (approximately 10 times) between the Cell/B.E. processor and conventional PC processors. The DMA transfers can be set up and controlled by the SPE that is sourcing or receiving the data, or in some circumstances, by the PPE or another SPE.

1.2 Scaling the three performance-limiting walls

The Cell/B.E. overcomes three important limiters of contemporary microprocessor performance: power use, memory use, and processor frequency.

1.2.1 Scaling the power-limitation wall

Increasingly, microprocessor performance is limited by achievable power dissipation rather than by the number of available integrated-circuit resources (transistors and wires). Therefore, the only way to significantly increase the performance of microprocessors is to improve power efficiency at about the same rate as the performance increase.

One way to increase power efficiency is to differentiate between the following types of processors:

- Processors that are optimized to run an operating system and control-intensive code
- Processors that are optimized to run compute-intensive applications
The Cell/B.E. processor does this by providing a general-purpose PPE to run the operating system and other control-plane code, as well as eight SPEs specialized for computing data-rich (data-plane) applications. The specialized SPEs are more compute efficient because they have simpler hardware implementations. The hardware does not devote transistors to branch prediction, out-of-order execution, speculative execution, shadow registers and register renaming, extensive pipeline interlocks, and so on. By weight, more of the transistors are used for computation than in conventional processor cores.

### 1.2.2 Scaling the memory-limitation wall

On multi-gigahertz symmetric multiprocessors (SMPs), even those with integrated memory controllers, latency to DRAM memory is currently approaching 1,000 cycles. As a result, program performance is dominated by the activity of moving data between main storage (the effective-address space that includes main memory) and the processor. Increasingly, compilers and even application writers must manage this movement of data explicitly, even though the hardware cache mechanisms are supposed to relieve them of this task.

The SPEs of the Cell/B.E. processor use two mechanisms to deal with long main-memory latencies:

- Three-level memory structure (main storage, local storage in each SPE, and large register files in each SPE)
- Asynchronous DMA transfers between main storage and local storage

These features allow programmers to schedule simultaneous data and code transfers to cover long latencies effectively. Because of this organization, the Cell/B.E. processor can usefully support 128 simultaneous transfers between the eight SPE local storage and main storage. This surpasses the number of simultaneous transfers on conventional processors by a factor of almost twenty.

### 1.2.3 Scaling the frequency-limitation wall

Conventional processors require increasingly deeper instruction pipelines to achieve higher operating frequencies. This technique has reached a point of diminishing returns, and even negative returns if power is taken into account.

By specializing the PPE and the SPEs for control and compute-intensive tasks, respectively, the CBEA, on which the Cell/B.E. processor is based, allows both the PPE and the SPEs to be designed for high frequency without excessive overhead. The PPE achieves efficiency primarily by executing two threads simultaneously rather than by optimizing single-thread performance.
Each SPE achieves efficiency by using a large register file, which supports many simultaneous in-process instructions without the overhead of register-renaming or out-of-order processing. Each SPE also achieves efficiency by using asynchronous DMA transfers, which support many concurrent memory operations without the overhead of speculation.

### 1.2.4 How the Cell/B.E. processor overcomes performance limitations

By optimizing control-plane and data-plane processors individually, the Cell/B.E. processor alleviates the problems posed by power, memory, and frequency limitations. The net result is a processor that, at the power budget of a conventional PC processor, can provide approximately ten-fold the peak performance of a conventional processor.

Actual application performance varies. Some applications may benefit little from the SPEs, where others show a performance increase well in excess of ten-fold. In general, compute-intensive applications that use 32-bit or smaller data formats, such as single-precision floating-point and integer, are excellent candidates for the Cell/B.E. processor.

### 1.3 Hardware environment

In the following sections, we describe the different components of the Cell/B.E. hardware environment.

#### 1.3.1 The processor elements

Figure 1-1 on page 4 shows a high-level block diagram of the Cell/B.E. processor hardware. There is one PPE and there are eight identical SPEs. All processor elements are connected to each other and to the on-chip memory and I/O controllers by the memory-coherent element interconnect bus (EIB).

The PPE contains a 64-bit, dual-thread PowerPC Architecture RISC core and supports a PowerPC virtual-memory subsystem. It has 32 KB level-1 (L1) instruction and data caches and a 512 KB level-2 (L2) unified (instruction and data) cache. It is intended primarily for control processing, running operating systems, managing system resources, and managing SPE threads. It can run existing PowerPC Architecture software and is well-suited to executing system-control code. The instruction set for the PPE is an extended version of the PowerPC instruction set. It includes the vector/SIMD multimedia extensions and associated C/C++ intrinsic extensions.
The eight identical SPEs are SIMD processor elements that are optimized for data-rich operations allocated to them by the PPE. Each SPE contains a RISC core, 256 KB software-controlled LS for instructions and data, and a 128-bit, 128-entry unified register file. The SPEs support a special SIMD instruction set, called the Synergistic Processor Unit Instruction Set Architecture, and a unique set of commands for managing DMA transfers and interprocessor messaging and control. SPE DMA transfers access main storage by using PowerPC effective addresses. As in the PPE, SPE address translation is governed by PowerPC Architecture segment and page tables, which are loaded into the SPEs by privileged software running on the PPE. The SPEs are not intended to run an operating system.

An SPE controls DMA transfers and communicates with the system by means of channels that are implemented in and managed by the memory flow controller (MFC) of the SPE. The channels are unidirectional message-passing interfaces (MPIs). The PPE and other devices in the system, including other SPEs, can also access this MFC state through the memory-mapped I/O (MMIO) registers and queues of the MFC. These registers and queues are visible to software in the main-storage address space.

1.3.2 The element interconnect bus

The EIB is the communication path for commands and data between all processor elements on the Cell/B.E. processor and the on-chip controllers for memory and I/O. The EIB supports full memory-coherent and SMP operations. Thus, a Cell/B.E. processor is designed to be grouped coherently with other Cell/B.E. processors to produce a cluster.

The EIB consists of four 16-byte-wide data rings. Each ring transfers 128 bytes (one PPE cache line) at a time. Each processor element has one on-ramp and one off-ramp. Processor elements can drive and receive data simultaneously. Figure 1-1 on page 4 shows the unit ID numbers of each element and the order in which the elements are connected to the EIB. The connection order is important to programmers who are seeking to minimize the latency of transfers on the EIB. The latency is a function of the number of connection hops, so that transfers between adjacent elements have the shortest latencies, and transfers between elements separated by six hops have the longest latencies.

The internal maximum bandwidth of the EIB is 96 bytes per processor-clock cycle. Multiple transfers can be in-process concurrently on each ring, including more than 100 outstanding DMA memory transfer requests between main storage and the SPEs in either direction. This requests might also include SPE memory to and from the I/O space. The EIB does not support any particular quality-of-service (QoS) behavior other than to guarantee forward progress. However, a resource allocation management (RAM) facility, shown in Figure 1-1
on page 4, resides in the EIB. Privileged software can use it to regulate the rate at which resource requesters (the PPE, SPEs, and I/O devices) can use memory and I/O resources.

1.3.3 Memory interface controller

The on-chip memory interface controller (MIC) provides the interface between the EIB and physical memory. The IBM BladeCenter QS20 blade server supports one or two Rambus extreme data rate (XDR) memory interfaces, which together support between 64 MB and 64 GB of XDR DRAM memory. The IBM BladeCenter QS21 blade server uses normal double data rate (DDR) memory and additional hardware logic to implement the MIC.

Memory accesses on each interface are 1 to 8, 16, 32, 64, or 128 bytes, with coherent memory ordering. Up to 64 reads and 64 writes can be queued. The resource-allocation token manager provides feedback about queue levels.

The MIC has multiple software-controlled modes, which includes the following types:

- Fast-path mode, for improved latency when command queues are empty
- High-priority read, for prioritizing SPE reads in front of all other reads
- Early read, for starting a read before a previous write completes
- Speculative read
- Slow mode, for power management

The MIC implements a closed-page controller (bank rows are closed after being read, written, or refreshed), memory initialization, and memory scrubbing.

The XDR DRAM memory is ECC-protected, with multi-bit error detection and optional single-bit error correction. It also supports write-masking, initial and periodic timing calibration. It also supports write-masking, initial and periodic timing calibration, dynamic width control, sub-page activation, dynamic clock gating, and 4, 8, or 16 banks.

1.3.4 Cell Broadband Engine interface unit

The on-chip Cell/B.E. interface (BEI) unit supports I/O interfacing. It includes a bus interrupt controller (BIC), I/O controller (IOC), and internal interrupt controller (IIC), as defined in the Cell Broadband Engine Architecture document.\(^1\) It

\(^1\) The Cell Broadband Engine Architecture document is on the Web at the following address: http://www-01.ibm.com/chips/techlib/techlib.nsf/techdocs/1AEAE1270EA2776387257060006E61BA/$file/CBEA_v1.02_110ct2007_pub.pdf
manages data transfers between the EIB and I/O devices and provides I/O address translation and command processing.

The BEI supports two Rambus FlexIO interfaces. One of the two interfaces, IOIF1, supports only a noncoherent I/O interface (IOIF) protocol, which is suitable for I/O devices. The other interface, IOIF0 (also called BIF/IOIF0), is software-selectable between the noncoherent IOIF protocol and the memory-coherent Cell/B.E. interface (Broadband Interface (BIF)) protocol. The BIF protocol is the internal protocol of the EIB. It can be used to coherently extend the EIB, through IOIF0, to another memory-coherent device, that can be another Cell/B.E. processor.

### 1.4 Programming environment

In the following sections, we provide an overview of the programming environment.

#### 1.4.1 Instruction sets

The instruction set for the PPE is an extended version of the PowerPC Architecture instruction set. The extensions consist of the vector/SIMD multimedia extensions, a few additions and changes to PowerPC Architecture instructions, and C/C++ intrinsics for the vector/SIMD multimedia extensions.

The instruction set for the SPEs is a new SIMD instruction set, the *Synergistic Processor Unit Instruction Set Architecture*, with accompanying C/C++ intrinsics. It also has a unique set of commands for managing DMA transfer, external events, interprocessor messaging, and other functions. The instruction set for the SPEs is similar to that of the vector/SIMD multimedia extensions for the PPE, in the sense that they operate on SIMD vectors. However, the two vector instruction sets are distinct. Programs for the PPE and SPEs are often compiled by different compilers, generating code streams for two entirely different instruction sets.

Although most coding for the Cell/B.E. processor might be done by using a high-level language such as C or C++, an understanding of the PPE and SPE machine instructions adds considerably to a software developer's ability to produce efficient, optimized code. This is particularly true because most of the C/C++ intrinsics have a mnemonic that relates directly to the underlying assembly language mnemonic.
1.4.2 Storage domains and interfaces

The Cell/B.E. processor has two types of storage domains: one main-storage domain and eight SPE LS domains, as shown in Figure 1-2. In addition, each SPE has a channel interface for communication between its Synergistic Processor Unit (SPU) and its MFC.

The main-storage domain, which is the entire effective-address space, can be configured by PPE privileged software to be shared by all processor elements and memory-mapped devices in the system. The state of an MFC is accessed by its associated SPU through the channel interface. This state can also be accessed by the PPE and other devices, including other SPEs, by means of the MMIO registers of the MFC in the main-storage space. The LS of an SPU can also be accessed by the PPE and other devices through the main-storage space.
in a non-coherent manner. The PPE accesses the main-storage space through its PowerPC processor storage subsystem (PPSS).

The address-translation mechanisms used in the main-storage domain are described in Section 4, “Virtual Storage Environment,” of Cell Broadband Engine Programming Handbook, Version 1.1. In this same document, you can find a description of the channel domain in Section 19, “DMA Transfers and Interprocessor Communication.” The SPU of an SPE can fetch instructions only from its own LS, and load and store instructions executed by the SPU can only access the LS. SPU software uses LS addresses, not main storage effective addresses, to do this. The MFC of each SPE contains a DMA controller. DMA transfer requests contain both an LS address and an effective address, thereby facilitating transfers between the domains.

Data transfer between an SPE local storage and main storage is performed by the MFC that is local to the SPE. Software running on an SPE sends commands to its MFC by using the private channel interface. The MFC can also be manipulated by remote SPEs, the PPE, or I/O devices by using memory mapped I/O. Software running on the associated SPE interacts with its own MFC through its channel interface. The channels support enqueuing of DMA commands and other facilities, such as mailbox and signal-notification messages. Software running on the PPE or another SPE can interact with an MFC through MMIO registers, which are associated with the channels and visible in the main-storage space.

Each MFC maintains and processes two independent command queues for DMA and other commands: one queue for its associated SPU and another queue for other devices that access the SPE through the main-storage space. Each MFC can process multiple in-progress DMA commands. Each MFC can also autonomously manage a sequence of DMA transfers in response to a DMA list command from its associated SPU (but not from the PPE or other SPEs). Each DMA command is tagged with a tag group ID that allows software to check or wait on the completion of commands in a particular tag group.

The MFCs support naturally aligned DMA transfer sizes of 1, 2, 4, or 8 bytes, and multiples of 16 bytes, with a maximum transfer size of 16 KB per DMA transfer. DMA list commands can initiate up to 2,048 such DMA transfers. Peak transfer performance is achieved if both the effective addresses and the LS addresses are 128-byte aligned and the size of the transfer is an even multiple of 128 bytes.

Each MFC has a synergistic memory management (SMM) unit that processes address-translation and access-permission information that is supplied by the

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PPE operating system. To process an effective address provided by a DMA command, the SMM uses essentially the same address translation and protection mechanism that is used by the memory management unit (MMU) in the PPSS of the PPE. Thus, DMA transfers are coherent with respect to system storage, and the attributes of system storage are governed by the page and segment tables of the PowerPC Architecture.

### 1.4.3 Bit ordering and numbering

Storage of data and instructions in the Cell/B.E. processor uses big-endian ordering, which has the following characteristics:

- The most-significant byte is stored at the lowest address, and the least-significant byte is stored at the highest address.
- Bit numbering within a byte goes from most-significant bit (bit 0) to least-significant bit (bit n).

This differs from other big-endian processors.

Figure 1-3 shows a summary of the byte ordering and bit ordering in memory and the bit-numbering conventions.

![Bit and Byte Order for a 32-bit Word](image)

![Bit and Byte Order for a 128-bit Register](image)

*Figure 1-3  Big-endian byte and bit ordering*

Neither the PPE nor the SPEs, including their MFCs, support little-endian byte ordering. The DMA transfers of the MFC are simply byte moves, without regard to the numeric significance of any byte. Thus, the big-endian or little-endian issue becomes irrelevant to the movement of a block of data. The byte-order mapping only becomes significant when data is loaded or interpreted by a processor element or an MFC.
1.4.4 Runtime environment

The PPE runs PowerPC Architecture applications and operating systems, which can include both PowerPC Architecture instructions and vector/SIMD multimedia extension instructions. To use all of the Cell/B.E. processor’s features, the PPE requires an operating system that supports these features, such as multiprocessing with the SPEs, access to the PPE vector/SIMD multimedia extension operations, the Cell/B.E. interrupt controller, and all the other functions provided by the Cell/B.E. processor.

A main thread running on the PPE can interact directly with an SPE thread through the LS of the SPE. The thread can interact indirectly through the main-storage space. A thread can poll or sleep while waiting for SPE threads. The PPE thread can also communicate through mailbox and signal events that are implemented in the hardware.

The operating system defines the mechanism and policy for selecting an available SPE to schedule an SPU thread to run on. It must prioritize among all the Cell/B.E. applications in the system, and it must schedule SPE execution independently from regular main threads. The operating system is also responsible for runtime loading, the passing of parameters to SPE programs, notification of SPE events and errors, and debugger support.
Chapter 2. IBM SDK for Multicore Acceleration

In this chapter, we describe the software tools and libraries that are found in the Cell Broadband Engine (Cell/B.E.) Software Development Kit (SDK). We briefly discuss the following topics:

- 2.1, “Compilers” on page 18
- 2.2, “IBM Full System Simulator” on page 19
- 2.3, “Linux kernel” on page 20
- 2.4, “Cell/B.E. libraries” on page 21
- 2.5, “Code examples and example libraries” on page 23
- 2.6, “Performance tools” on page 24
- 2.7, “IBM Eclipse IDE for the SDK” on page 26
- 2.8, “Hybrid-x86 programming model” on page 26
2.1 Compilers

A number of IBM compilers are supplied as part of the SDK for Multicore Acceleration. In the following sections, we briefly describe the IBM product compilers and open source compilers in the SDK.

2.1.1 GNU toolchain

The GNU toolchain, including compilers, the assembler, the linker, and miscellaneous tools, is available for both the PowerPC Processor Unit (PPU) and Synergistic Processor Unit (SPU) instruction set architectures. On the PPU, it replaces the native GNU toolchain, which is generic for the PowerPC Architecture, with a version that is tuned for the Cell/B.E. PPU processor core. The GNU compilers are the default compilers for the SDK.

The GNU toolchains run natively on Cell/B.E. hardware or as cross-compilers on PowerPC or x86 machines.

2.1.2 IBM XLC/C++ compiler

IBM XL C/C++ for Multicore Acceleration for Linux is an advanced, high-performance cross-compiler that is tuned for the Cell Broadband Engine Architecture (CBEA). The XL C/C++ compiler, which is hosted on an x86, IBM PowerPC technology-based system, or an IBM BladeCenter QS21, generates code for the PPU or SPU. The compiler requires the GCC toolchain for the CBEA, which provides tools for cross-assembling and cross-linking applications for both the PowerPC Processor Element (PPE) and Synergistic Processor Element (SPE).

OpenMP compiler: The IBM XLC/C++ compiler that comes with SDK 3 is an OpenMP directed single source compiler that supports automatic program partitioning, data virtualization, code overlay, and more. This version of the compiler is in beta mode. Therefore, users should not base production applications on this compiler.

2.1.3 GNU ADA compiler

The GNU toolchain also contains an implementation of the GNU ADA compiler. This compiler comes in an existing Cell/B.E. processor and an x86 cross-compiler. The initial version of this compiler supports code generation for the PPU processor.
2.1.4 IBM XL Fortran for Multicore Acceleration for Linux

IBM XL Fortran for Multicore Acceleration for Linux is the latest addition to the IBM XL family of compilers. It adopts proven high-performance compiler technologies that are used in its compiler family predecessors. It also adds new features that are tailored to exploit the unique performance capabilities of processors that are compliant with the new CBEA.

This version of XL Fortran is a cross-compiler. First, you compile your applications on an IBM System p™ compilation host running Red Hat Enterprise Linux (RHEL) 5.1. Then you move the executable application produced by the compiler onto a Cell/B.E. system that is also running the RHEL 5.1 Linux distribution. The Cell/B.E. system is the execution host where you run your compiled application.

2.2 IBM Full System Simulator

The IBM Full System Simulator (referred to as the simulator in this chapter) is a software application that emulates the behavior of a full system that contains a Cell/B.E. processor. You can start a Linux operating system on the simulator, simulate a two chip Cell/B.E. environment, and run applications on the simulated operating system. The simulator also supports the loading and running of statically-linked executable programs and stand-alone tests without an underlying operating system. Other functions, such as debug output, are not available on the hardware.

The simulator infrastructure is designed for modeling the processor and system-level architecture at levels of abstraction, which vary from functional to performance simulation models with several hybrid fidelity points in between:

- Functional-only simulation

  This type of simulation models the program-visible effects of instructions without modeling the time it takes to run these instructions. Functional-only simulation assumes that each instruction can be run in a constant number of cycles. Memory accesses are synchronous and are performed in a constant number of cycles.

  This simulation model is useful for software development and debugging when a precise measure of execution time is not significant. Functional simulation proceeds much more rapidly than performance simulation, and therefore, is useful for fast-forwarding to a specific point of interest.
Performance simulation

For system and application performance analysis, the simulator provides performance simulation (also referred to as timing simulation). A performance simulation model represents internal policies and mechanisms for system components, such as arbiters, queues, and pipelines.

Operation latencies are modeled dynamically to account for both processing time and resource constraints. Performance simulation models have been correlated against hardware or other references to acceptable levels of tolerance.

The simulator for the Cell/B.E. processor provides a cycle-accurate SPU core model that can be used for performance analysis of computationally-intense applications.

2.2.1 System root image for the simulator

The system root image for the simulator is a file that contains a disk image of Fedora files, libraries, and binaries that can be used within the system simulator. This disk image file is preloaded with a full range of Fedora utilities and includes all of the Cell/B.E. Linux support libraries.

2.3 Linux kernel

For the IBM BladeCenter QS21 blade server, the kernel is installed into the /boot directory, yaboot.conf is modified, and a reboot is required to activate this kernel. The cellsdk installation task is documented in the Cell Broadband Engine Software Development Kit 2.1 Installation Guide Version 2.1, SC33-8323-02.¹

2.4 Cell/B.E. libraries

In the following sections, we describe various the programming libraries.

2.4.1 SPE Runtime Management Library

The SPE Runtime Management Library (libspe) constitutes the standardized low-level application programming interface (API) for application access to the Cell/B.E. SPEs. This library provides an API to manage SPEs that are neutral with respect to the underlying operating system and its methods. Implementations of this library can provide additional functionality that allows for access to operating system or implementation-dependent aspects of SPE runtime management. These capabilities are not subject to standardization. Their use may lead to non-portable code and dependencies on certain implemented versions of the library.

2.4.2 SIMD Math Library

The traditional math functions are scalar instructions and do not take advantage of the powerful single instruction, multiple data (SIMD) vector instructions that are available in both the PPU and SPU in the CBEA. SIMD instructions perform computations on short vectors of data in parallel, instead of on individual scalar data elements. They often provide significant increases in program speed because more computation can be done with fewer instructions.

2.4.3 Mathematical Acceleration Subsystem libraries

The Mathematical Acceleration Subsystem (MASS) consists of libraries of mathematical intrinsic functions, which are tuned specifically for optimum performance on the Cell/B.E. processor. Currently the 32-bit, 64-bit PPU, and SPU libraries are supported.

These libraries offer the following advantages:

- Include both scalar and vector functions, are thread-safe, and support both 32-bit and 64-bit compilations
- Offer improved performance over the corresponding standard system library routines
- Are intended for use in applications where slight differences in accuracy or handling of exceptional values can be tolerated
2.4.4 Basic Linear Algebra Subprograms

The Basic Linear Algebra Subprograms (BLAS) library is based upon a published standard interface for commonly used linear algebra operations in high-performance computing (HPC) and other scientific domains. It is widely used as the basis for other high quality linear algebra software, for example LAPACK and ScaLAPACK. The Linpack (HPL) benchmark largely depends on a single BLAS routine (DGEMM) for good performance.

The BLAS API is available as standard ANSI C and standard FORTRAN 77/90 interfaces. BLAS implementations are also available in open source (netlib.org).

The BLAS library in IBM SDK for Multicore Acceleration supports only real single-precision (SP) and real double-precision (DP) versions. All SP and DP routines in the three levels of standard BLAS are supported on the PPE. These routines are available as PPE APIs and conform to the standard BLAS interface.

Some of these routines are optimized by using the SPEs and show a marked increase in performance compared to the corresponding versions that are implemented solely on the PPE. These optimized routines have an SPE interface in addition to the PPE interface. The SPE interface does not conform to, yet provides a restricted version of, the standard BLAS interface. Moreover, the single precision versions of these routines have been further optimized for maximum performance by using various features of the SPE.

2.4.5 ALF library

The Accelerated Library Framework (ALF) provides a programming environment for data and task parallel applications and libraries. The ALF API provides library developers with a set of interfaces to simplify library development on heterogenous multi-core systems. Library developers can use the provided framework to offload computationally intensive work to the accelerators. More complex applications can be developed by combining the several function offload libraries. Application programmers can also choose to implement their applications directly to the ALF interface.

The ALF supports the multiple-program-multiple-data (MPMD) programming module where multiple programs can be scheduled to run on multiple accelerator elements at the same time.

The ALF library includes the following functionality:

- Data transfer management
- Parallel task management
- Double buffering and dynamic load balancing
2.4.6 Data Communication and Synchronization library

The Data Communication and Synchronization (DaCS) library provides a set of services for handling process-to-process communication in a heterogeneous multi-core system. In addition to the basic message passing service, the library includes the following services:

- Mailbox services
- Resource reservation
- Process and process group management
- Process and data synchronization
- Remote memory services
- Error handling

The DaCS services are implemented as a set of APIs that provide an architecturally neutral layer for application developers. They structure the processing elements, referred to as *DaCS elements* (DE), into a hierarchical topology. This includes general purpose elements, referred to as *host elements* (HE), and special processing elements, referred to as *accelerator elements* (AE). HEs usually run a full operating system and submit work to the specialized processes that run in the AEs.

2.5 Code examples and example libraries

The example libraries package provides a set of optimized library routines that greatly reduce the development cost and enhance the performance of Cell/B.E. programs. To demonstrate the versatility of the CBEA, a variety of application-oriented libraries are included, such as the following libraries:

- Fast Fourier Transform (FFT)
- Image processing
- Software managed cache
- Game math
- Matrix operation
- Multi-precision math
- Synchronization
- Vector

Additional examples and demonstrations show how you can exploit the on-chip computational capacity.
2.6 Performance tools

The Cell/B.E. SDK supports many of the traditional Linux-based performance tools that are available. The performance tools, such as gprof, pertain specifically to the PPE execution environment and do not support the SPE environment. The tools in the following sections are special tools that support the PPE, SPE, or both types of environments.

2.6.1 SPU timing tool

The SPU static timing tool, spu_timing, annotates an SPU assembly file with scheduling, timing, and instruction issue estimates assuming a straight, linear execution of the program, which is useful for analyzing basic code blocks. The tool generates a textual output of the execution pipeline of the SPE instruction stream from this input assembly file. The output generated can show pipeline stalls, which can be explained by looking at the subsequent instructions. Data dependencies are pipeline hazards that can be readily identified by using this tool.

Lastly, this is a static analysis tool. It does not identify branch behavior or memory transfer delays.

2.6.2 OProfile

OProfile is a Linux tool that exists on other architectures besides the Cell/B.E. It has been extended to support the unique hardware on the PPU and SPUs. It is a sampling based tool that does not require special source compile flags to produce useful data reports.

The opreport tool produces the output report. Reports can be generated based on the file names that correspond to the samples, symbol names, or annotated source code listings. Special source compiler flags are required in this case.

2.6.3 Cell-perf-counter tool

The cell-perf-counter (cpc) tool is used for setting up and using the hardware performance counters in the Cell/B.E. processor. These counters allow you to see how many times certain hardware events are occurring, which is useful if you are analyzing the performance of software running on a Cell/B.E. system. Hardware events are available from all of the logical units within the Cell/B.E. processor, including the PPE, SPEs, interface bus, and memory and I/O controllers. Four 32-bit counters, which can also be configured as pairs of 16-bit
counters, are provided in the Cell/B.E. performance monitoring unit (PMU) for counting these events.

### 2.6.4 Performance Debug Tool

The Cell/B.E. Performance Debug Tool (PDT) provides programmers with a means of analyzing the execution of such a system and tracking problems in order to optimize execution time and utilization of resources. The PDT addresses performance debugging of one Cell/B.E. board with two PPEs that share the main memory, run under the same (Linux) operating system, and share up to 16 SPEs. The PDT also enables event tracing on the Hybrid-x86.

### 2.6.5 Feedback Directed Program Restructuring tool

The Feedback Directed Program Restructuring (FDPR-Pro or fdprpro) tool for the Linux on Power is a performance tuning utility that reduces the execution time and the real memory utilization of user space application programs. It optimizes the executable image of a program by collecting information about the behavior of the program under a workload. It then creates a new version of that program that is optimized for that workload. The new program typically runs faster and uses less real memory than the original program and supports the Cell/B.E. environment.

### 2.6.6 Visual Performance Analyzer

Visual Performance Analyzer (VPA) is an Eclipse-based performance visualization toolkit. It consists of six major components:

- Profile Analyzer
- Code Analyzer
- Pipeline Analyzer
- Counter Analyzer
- Trace Analyzer
- Control Flow Analyzer

Profile Analyzer provides a powerful set of graphical and text-based views. By using these views, users can narrow down performance problems to a particular process, thread, module, symbol, offset, instruction, or source line. Profile Analyzer supports time-based system profiles (Tprofs) that are collected from a number of IBM platforms and the Linux profile tool oprofile. The Cell/B.E. platform is now a fully supported environment for VPA.
2.7 IBM Eclipse IDE for the SDK

IBM Eclipse IDE for the SDK is built upon the Eclipse and C Development Tools (CDT) platform. It integrates the GNU toolchain, compilers, the Full System Simulator, and other development components to provide a comprehensive, Eclipse-based development platform that simplifies development. Eclipse IDE for the SDK includes the following key features:

- A C/C++ editor that supports syntax highlighting, a customizable template, and an outline window view for procedures, variables, declarations, and functions that appear in source code
- A visual interface for the PPE and SPE combined GDB (GNU debugger) and seamless integration of the simulator into Eclipse
- Automatic builder, performance tools, and several other enhancements; remote launching, running and debugging on a BladeCenter QS21; and ALF source code templates for programming models within IDE
- An ALF Code Generator to produce an ALF template package with C source code and a readme.txt file
- A configuration option for both the Local Simulator and Remote Simulator target environments so that you can choose between launching a simulation machine with the Cell/B.E. processor or an enhanced CBEA-compliant processor with a fully pipelined, double precision SPE processor
- Remote Cell/B.E. and simulator BladeCenter support
- SPU timing integration and automatic makefile generation for both GCC and XLC projects

2.8 Hybrid-x86 programming model

The CBEA is an example of a multi-core hybrid system on a chip. Heterogeneous cores are integrated on a single processor with an inherent memory hierarchy. Specifically, the SPEs can be thought of as computational accelerators for a more general purpose PPE core. These concepts of hybrid systems, memory hierarchies, and accelerators can be extended more generally to coupled I/O devices. Such systems exist today, such as GPUs in Peripheral Component Interconnect Express (PCIe) slots for workstations and desktops.

Similarly, the Cell/B.E. processor is used in systems as an accelerator, where computationally intensive workloads that are well suited for the CBEA are off-loaded from a more standard processing node. There are potentially many
ways to move data and functions from a host processor to an accelerator processor and vice versa.

The SDK has implementations of the Cell/B.E. multi-core data communication and programming model libraries, DaCS and ALF, which can be used on x86 or Linux host process systems with Cell/B.E.-based accelerators. They provide a consistent methodology and set of APIs for a variety of hybrid systems, including the Cell/B.E. SoC hybrid system. The current implementation is over TCP/IP sockets and is provided so that you can gain experience with this programming style and focus on how to manage the distribution of processing and data decomposition. For example, in the case of hybrid programming when moving data point to point over a network, use care to maximize the computational work done on accelerator nodes potentially with asynchronous or overlapping communication, given the potential cost in communicating input and results.
Part 2

Programming environment

In this part, we provide in-depth coverage of various programming methods, tools, strategies, and adaptions to different computing workloads. Specifically this part includes the following chapters:

- Chapter 3, “Enabling applications on the Cell Broadband Engine hardware” on page 31
- Chapter 4, “Cell Broadband Engine programming” on page 75
- Chapter 5, “Programming tools and debugging techniques” on page 329
- Chapter 6, “The performance tools” on page 417
- Chapter 7, “Programming in distributed environments” on page 445
Enabling applications on the Cell Broadband Engine hardware

In this chapter, we describe the process of enabling an existing application on the Cell Broadband Engine (Cell/B.E.) hardware. The goal is to provide guidance for choosing the best programming model and framework for a given application. This is valuable information for people who develop applications and for IT specialists who must manage a Cell/B.E. application enablement project.

We include an example of a completely new application that is written from scratch. You can consider this example as a case of application enablement, where the starting point is not the actual code, but the algorithms with no initial data layout decisions. In a sense, this is an easier way because the options are completely open and not biased by the current state of the code.

In this chapter, we answer the following questions:

- Should we enable this application on the Cell/B.E. hardware? Is it a good fit?
- If the answer to the previous question is “yes,” then which parallel programming model should we use? The Cell Broadband Engine Architecture (CBEA), with its heterogenous design and software controlled memory
hierarchy, offers new parallel programming paradigms to complement the well established ones.

- Which Cell/B.E. programming framework will best support the programming model that was chosen for the application under study?

After we answer these questions, we make the necessary code changes to exploit the CBEA. We describe this process and show that it can be iterative and incremental, which are two interesting features. Considering these features, we can use a step-by-step approach, inserting checkpoints during the course of the porting project to track the progress. Finally, we present scenarios and make a first attempt at creating a set of design patterns for Cell/B.E. programming.

In this chapter, we begin by defining the concepts and terminology. We introduce the following concepts:

- The computational kernels frequently found in applications
- The distinctive features of the CBEA, which are covered in detail in Chapter 4, “Cell Broadband Engine programming” on page 75
- The parallel programming models
- The Cell/B.E. programming frameworks, which are described in Chapter 4, “Cell Broadband Engine programming” on page 75, and Chapter 7, “Programming in distributed environments” on page 445

Next we describe the relationship between the computational kernels and the Cell/B.E. features and between parallel programming models and Cell/B.E. programming frameworks. We give examples of the most common parallel programming models and contrast them in terms of control parallelism and data transfers. We also present design patterns for Cell/B.E. programming following a formalism used in other areas of computer sciences.

Specifically, this chapter includes the following sections:

- 3.1, “Concepts and terminology”
- 3.2, “Determining whether the Cell/B.E. system fits the application requirements” on page 48
- 3.3, “Deciding which parallel programming model to use” on page 53
- 3.4, “Deciding which Cell/B.E. programming framework to use” on page 61
- 3.5, “The application enablement process” on page 63
- 3.6, “A few scenarios” on page 66
- 3.7, “Design patterns for Cell/B.E. programming” on page 70
3.1 Concepts and terminology

Figure 3-1 shows the concepts and how they are related. As described in this section, the figure shows an application as having one or more computational kernels and one or more potential parallel programming models. The computational kernels exercise or stress one or more of the Cell/B.E. features (the Q1 connection). The different Cell/B.E. features can either strongly or weakly support the different parallel programming model choices (the Q2 connection). The chosen parallel programming model can be implemented on the CBEA by using various programming frameworks (the Q3 connection).

To answer questions Q1 and Q2, the programmer must be able to match the characteristics of the computational kernel and parallel programming model to the strengths of the CBEA. Many programming frameworks are available for the CBEA. Which one is best suited to implement the parallel programming model that is chosen for the application? We provide advice to the programmer for question Q3 in 3.4, “Deciding which Cell/B.E. programming framework to use” on page 61.

Figure 3-1   Overview of programming considerations and relations
3.1.1 The computation kernels

A study by David Patterson et al. in *The Landscape of Parallel Computing Research: A View from Berkeley* [1 on page 623], complements earlier work from Phillip Colella [8 on page 624]. Patterson et al. establish that the computational characteristics and data movement patterns of all applications in scientific computing, embedded computing, and desktop and server computing can be captured by no more than thirteen different kernels. They call them the “13 dwarfs” in their paper.\(^1\)

This work is based on a careful examination of the most popular benchmark suites:

- Embedded Microprocessor Benchmark Consortium (EEMBC) for the embedded computing
- Standard Performance Evaluation Consortium (SPEC) int and fp for desktop and server computing
- High Performance Computing Challenge (HPCC) and NASA Advanced Supercomputing (NAS) parallel benchmarks for scientific computing,

This work is also based on input from other domains including machine learning, database, computer graphics, and games. The first seven dwarfs are the ones that were initially found by Phillip Colella. The six remaining ones were identified by Patterson et al. The intent of the study is to help the parallel computing research community, in both academia and the industry, by providing a limited set of patterns against which new ideas for hardware and software can be evaluated.

We describe the “13 dwarfs” in Table 3-1, with example applications or benchmarks. This table is adapted from *The Landscape of Parallel Computing Research: A View from Berkeley* [1 on page 623).

<table>
<thead>
<tr>
<th>Dwarf name</th>
<th>Description</th>
<th>Example, application, benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dense matrices</td>
<td>Basic Linear Algebra Subprograms (BLAS), matrix-matrix operations</td>
<td>HPCC:HPL, ScaLAPACK, NAS:LU</td>
</tr>
<tr>
<td>Sparse matrices</td>
<td>Matrix-vector operations with sparse matrices</td>
<td>SuperLU, SpMV, NAS:CG</td>
</tr>
<tr>
<td>Spectral methods</td>
<td>Fast Fourier Transform (FFT)</td>
<td>HPCC:FFT, NAS:FT, FFTW</td>
</tr>
</tbody>
</table>

\(^1\) Intel also classifies applications in three categories named RMS for Recognition, Mining and Synthesis to direct its research in computer architecture. See 9 on page 624.
During the course of writing *The Landscape of Parallel Computing Research: A View from Berkeley* [1 on page 623], IBM provided an additional classification for the 13 dwarfs. IBM did this by evaluating that factor that was often limiting its performance, whether it was the processor, memory latency, or memory bandwidth. Table 3-2 is extracted from that paper.

### Table 3-2 Performance bottlenecks of the 13 dwarfs

<table>
<thead>
<tr>
<th>Dwarf name</th>
<th>Description</th>
<th>Example, application, benchmark</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-body methods</td>
<td>Interactions between particles, external, near and far</td>
<td>NAMD, GROMACS</td>
</tr>
<tr>
<td>Structured grids</td>
<td>Regular grids, can be automatically refined</td>
<td>WRF, Cactus, NAS:MG</td>
</tr>
<tr>
<td>Unstructured grids</td>
<td>Irregular grids, finite elements and nodes</td>
<td>ABAQUS, FIDAP (Fluent)</td>
</tr>
<tr>
<td>Map-reduce</td>
<td>Independent data sets, simple reduction at the end</td>
<td>Monte-Carlo, NAS:EP, Ray tracing</td>
</tr>
<tr>
<td>Combinatorial logic</td>
<td>Logical functions on large data sets, encryption</td>
<td>AES, DES</td>
</tr>
<tr>
<td>Graph traversal</td>
<td>Decision tree, searching</td>
<td>XML parsing, Quicksort</td>
</tr>
<tr>
<td>Dynamic programming</td>
<td>Hidden Markov models, sequence alignment</td>
<td>BLAST</td>
</tr>
<tr>
<td>Back-track and Branch+Bound</td>
<td>Constraint optimization</td>
<td>Simplex algorithm</td>
</tr>
<tr>
<td>Graphical models</td>
<td>Hidden Markov models, Bayesian networks</td>
<td>HMMER, bioinformatics, genomics</td>
</tr>
<tr>
<td>Finite state machine</td>
<td>XML transformation, Huffman decoding</td>
<td>SPECInt:gcc</td>
</tr>
</tbody>
</table>

During the course of writing *The Landscape of Parallel Computing Research: A View from Berkeley* [1 on page 623], IBM provided an additional classification for the 13 dwarfs. IBM did this by evaluating that factor that was often limiting its performance, whether it was the processor, memory latency, or memory bandwidth. Table 3-2 is extracted from that paper.
The Cell/B.E. platform brings improvements in all three directions:

- Nine cores on a chip represent a lot of processor power.
- The extreme data rate (XDR) memory subsystem is extremely capable.
- The software managed memory hierarchy (direct memory access (DMA)) is a new way of dealing with the memory latency problem, quoted in the paper as the most critical one.

### 3.1.2 Important Cell/B.E. features

In this section, we consider features that are meaningful from an application programmer’s point of view. Referring to Table 3-3 on page 37, the column labelled “Not so good” only means that the feature is probably going to cause problems for the programmer or that code that exercises this feature a lot will not perform as fast as expected. The difference between the “Good” and “Not so good” columns is also related to the relative performance advantage of the Cell/B.E. platform over contemporary processors from IBM or others. This analysis is based on current hardware implementation at the time this book was written.

This table is likely to change with future products. Most of the items in the table are described in detail in Chapter 4, “Cell Broadband Engine programming” on page 75.
Table 3-3  Important Cell/B.E. features as seen from a programmer's perspective

<table>
<thead>
<tr>
<th>Good</th>
<th>Not so good</th>
</tr>
</thead>
<tbody>
<tr>
<td>Large register file</td>
<td></td>
</tr>
<tr>
<td>DMA (memory latency hiding)^a</td>
<td>DMA latency</td>
</tr>
<tr>
<td>Element interconnect bus (EIB) bandwidth ^b</td>
<td></td>
</tr>
<tr>
<td>Memory performance</td>
<td>Memory size</td>
</tr>
<tr>
<td>Single-instruction, multiple-data (SIMD)^c</td>
<td>Scalar performance (scalar on vector)</td>
</tr>
<tr>
<td>Local storage (latency/bandwidth)</td>
<td>Local storage (limited size)</td>
</tr>
<tr>
<td>Eight Synergistic Processor Element (SPE) per processor (high level of achievable parallelism)</td>
<td>Power Processor Element (PPE) performance</td>
</tr>
<tr>
<td>Nonuniform memory access (NUMA; good scaling)</td>
<td>Symmetric multiprocessor (SMP) scaling</td>
</tr>
<tr>
<td></td>
<td>Branching</td>
</tr>
<tr>
<td>Single precision floating point</td>
<td>Double precision floating point^d</td>
</tr>
</tbody>
</table>

^b. See 4.3.4, “Direct problem state access and LS-to-LS transfer” on page 144.  
^c. See 4.6.4, “SIMD programming” on page 258.  
^d. This is expected to improve with the introduction of the BladeCenter QS-22.

**Peak performance:** In general, the percentage of peak performance that can be achieved on the Cell/B.E. platform can be higher than for most general purpose processors. This percentage is a result of the large register file, the short local storage latency, the software managed memory hierarchy, the high EIB bandwidth, and the capable memory subsystem.

### 3.1.3 The parallel programming models

The parallel programming models abstract the hardware for the application programmers. The purpose is to offer an idealized view of the current system architectures, a view onto which applications can be mapped as shown in Figure 3-2 on page 38.
A parallel application tries to bind multiple resources for its own use, which may be in regard to memory and processors. The purpose is either to speed up the whole computation (more processors) or to treat bigger problems (more memory). The work of parallelizing an application involves the following tasks:

- Distributing the work across processors
- Distributing the data if the memory is distributed
- Synchronizing the subtasks, possibly through shared data access if the memory is shared
- Communicating the data if the memory is distributed

Let us look at the options for each of these components in the following sections.
**Work distribution**
The first task is to find concurrency in the application by exposing multiple independent tasks and grouping them together inside execution threads. The following options are possible:

- Independent tasks operating on largely independent data
- Domain decomposition, where the whole data can be split in multiple subdomains, each of which is assigned to a single task
- Streaming, where the same piece of data undergoes successive transformations, each of which is performed by a single task

All tasks are arranged in a string and pass data in a producer-consumer mode. The amount of concurrency here is the number of different steps in the computation.

Now each parallel task can perform the work itself, or it can call other processing resources for assistance. This process is completely transparent to the rest of the participating tasks. We find the following techniques:

- Function offload, where the compute-intensive part of the job is offloaded to a supposedly faster processor
- Accelerator mode, which is a variant of the previous technique, where multiple processors can be called to collectively speed up the computation

**Data distribution**
The data model is an important part of the parallelization work. Currently, the choices are available:

- Shared memory, in which every execution thread has direct access to other threads’ memory contents
- Distributed memory, which is the opposite of shared memory in that each execution thread can only access its own memory space

Specialized functions are required to import other threads’ data into its own memory space.

- Partitioned Global Address Space (PGAS), where each piece of data must be explicitly declared as either shared or local, but within a unified address space

**Task synchronization**
Sometimes, during program execution, the parallel tasks must synchronize. Synchronization can be realized in the following ways:

- Messages or other asynchronous events emitted by other tasks
- Locks or other synchronization primitives, for example, to access a queue
- Transactional memory mechanisms
**Data communication**

In the case where the data is distributed, tasks exchange information through one of the following two mechanisms:

- Message passing, using send and receive primitives
- Remote DMA (rDMA), which is sometimes defined as one-sided communication

The programming models can further be classified according to the way each of these tasks is taken care of, either explicitly by the programmer or implicitly by the underlying runtime environment. Table 3-4 lists common parallel programming models and shows how they can be described according to what was exposed previously.

<table>
<thead>
<tr>
<th>Programming model(^a)</th>
<th>Task distribution</th>
<th>Data distribution</th>
<th>Task synchronization</th>
<th>Data communication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Message passing interface (MPI)</td>
<td>Explicit</td>
<td>Explicit</td>
<td>Messages</td>
<td>Messages, can do rDMA too</td>
</tr>
<tr>
<td>pthreads</td>
<td>Explicit</td>
<td>N/A</td>
<td>Mutexes, condition variables</td>
<td>N/A</td>
</tr>
<tr>
<td>OpenMP</td>
<td>Implicit (directives)</td>
<td>N/A</td>
<td>Implicit (directives)</td>
<td>N/A</td>
</tr>
<tr>
<td>UPC, CAF (PGAS)</td>
<td>Explicit</td>
<td>Explicit</td>
<td>Implicit</td>
<td>Implicit</td>
</tr>
<tr>
<td>X10 (PGAS)</td>
<td>Explicit</td>
<td>Explicit</td>
<td>Future, clocks</td>
<td>Implicit</td>
</tr>
<tr>
<td>StreamIt</td>
<td>Explicit</td>
<td>Explicit</td>
<td>Explicit</td>
<td>Implicit</td>
</tr>
</tbody>
</table>

\(^a\) Not all of these programming models are available or appropriate for the Cell/B.E. platform.

**Programming model composition**

An application can use multiple programming models, which incurs additional efforts but may be dictated by the hardware on which it is to be run. For example, the MPI and OpenMP combination is quite common today for high-performance computing (HPC) applications because it matches the Beowulf\(^2\) type of clusters, interconnecting small SMP nodes (4- and 8-way) with a high-speed interconnection network.

\(^2\) Clusters based on commodity hardware. See [http://www.beowulf.org/overview/index.html](http://www.beowulf.org/overview/index.html)
In this discussion about the parallel programming models, we ignore the instruction level (multiple execution units) and word level (SIMD) parallelisms. They are to be considered as well to maximize the application performance, but they usually do not interfere with the high level tasks of data and work distribution.

### 3.1.4 The Cell/B.E. programming frameworks

The CBEA supports a wide range of programming frameworks, from the most basic ones, close to the hardware, to the most abstract ones.

At the lowest level, the Cell/B.E. chip appears to the programmer as a distributed memory cluster of 8+1 computational cores, with an ultra high-speed interconnect and a remote DMA engine on every core. On a single blade server, two Cell/B.E. chips can be viewed as either a single 16+2 cores compute resource (SMP mode) or a NUMA machine with two NUMA nodes.

Multiple blade servers can then be gathered in a distributed cluster, by using Beowulf, with a high-speed interconnect network such as a 10G Ethernet or InfiniBand®. Such a cluster is not any different, in regard to programming, from a cluster of Intel, AMD™, or IBM POWER™ technology-based SMP servers. It is likely that the programming model will be based on distributed memory programming that uses MPI as the communication layer across the blades.

An alternative arrangement is to have Cell/B.E. blade servers serve only as accelerator nodes for other systems. In this configuration, the application logic is not managed at the Cell/B.E. level but at the accelerated system level. The dialog that we are interested in is between the Cell/B.E. level and the system for which it provides acceleration.

The frameworks that are part of the IBM SDK for Multicore Acceleration are described in greater detail in 4.7, “Frameworks and domain-specific libraries” on page 289, and 7.1, “Hybrid programming models in SDK 3.0” on page 446. We only give a brief overview in the following sections.

#### libspe2 and newlib

libspe2 and newlib are the lowest possible levels for application programmers. By using the libspe2 and newlib libraries, programmers deal with each feature of the Cell/B.E. architecture with full control. libspe2 and newlib offer the following features:
- SPE context management for creating, running, scheduling, and deleting SPE contexts
- DMA primitives for accessing remote memory locations from the PPE and the SPEs
- Mailboxes, signal, events, and synchronization functions for PPE-SPE and SPE-SPE dialogs and control
- PPE-assisted calls, which is a mechanism to have the PPE service requests from the SPEs

**Important:** Libspe2 is a framework for obtaining access to the SPEs. Mailbox, DMAs, signals, and so on are much faster when using direct problem state. High performance programs should avoid making frequent libspe2 calls since they often use kernel services. Therefore, it is best to use libspe2 to get parallel tasks started and then to use the memory flow controller (MFC) hardware facilities for application task management, communications, and synchronization.

By using these libraries, any kind of parallel programming model can be implemented. Libspe2 is described in detail in 4.1, “Task parallelism and PPE programming” on page 77.

### Software cache

A software cache can help implement a shared memory parallel programming model when the data that the SPEs reference cannot be easily predicted. See 4.3.6, “Automatic software caching on SPE” on page 157, for more details.

### Data Communication and Synchronization

Data Communication and Synchronization (DaCS) provides services to multi-tier applications by using a hierarchy of processing elements. A DaCS program can be a host element (HE), an accelerator element (AE), or both if multiple levels of acceleration are needed. An AE can only communicate within the realm of the HE. An HE does not need to be of the same type as its AEs. This is the hybrid model. DaCS handles the necessary byte swapping if the data flows from a little endian machine to a big endian.

DaCS offers the following typical services:

- Resource and process management, where an HE manipulates its AEs
- Group management, for defining groups within which synchronization events like barriers can happen
- Message passing by using send and receive primitives
Mailboxes
Remote DMA operations
Process synchronization using barriers
Data synchronization using mutexes to protect memory accesses

The DaCS services are implemented as an API for the Cell/B.E.-only version and are complemented by a runtime daemon for the hybrid case. For a complete discussion, see 4.7.1, “Data Communication and Synchronization” on page 291, and 7.2, “Hybrid Data Communication and Synchronization” on page 449.

MPI
MPI is not part of the IBM SDK for Multicore Acceleration. However, any implementation for Linux on Power can run on the Cell/B.E., leveraging the PPE only. The following implementations are the most common:

- MPICH/MPICH2, from Argonne National Laboratory
- MVAPICH/MVAPICH2, from Ohio State University
- OpenMPI, from a large consortium involving IBM and Los Alamos National Laboratory, among others

There is no difference from a functional point of view between these MPI implementations running on the Cell/B.E. platform and running on other platforms. MPI is a widespread standard for writing distributed memory applications. As opposed to DaCS, MPI treats all tasks as equal. The programmer can decide later if some tasks are to play a particular role in the computation. MPI is implemented as an API and a runtime environment to support all sorts of interconnection mechanisms between the MPI tasks, including shared memory, sockets for TCP/IP networks, or OpenIB (OFED) for InfiniBand networks.

Accelerated Library Framework
An Accelerated Library Framework (ALF) program uses multiple ALF accelerator tasks to perform the compute-intensive part of the work. The idea is to have the host program split the work into multiple independent pieces, which are called work blocks. They are described by a computational kernel, the input data they need, and the output data they produce.

On the accelerator side, the programmer only has to code the computational kernel, unwrap the input data, and pack the output data when the kernel finishes processing. In between, the runtime system manages the work blocks queue on the accelerated side and gives control to the computational kernel upon receiving a new work block on the accelerator side.
The ALF imposes a clear separation between the application logic and control running on the host task from the computational kernels that run on the accelerator nodes, acting as service providers that are fed with input data and echo output data in return. The ALF run time provides the following services “for free” from the application programmer perspective:

- Work blocks queue management
- Load balancing between accelerators
- Transparent DMA transfers, exploiting the data transfer list used to describe the input and output data

Table 3-5 summarizes the various duties.

<table>
<thead>
<tr>
<th>Who</th>
<th>Does what</th>
</tr>
</thead>
<tbody>
<tr>
<td>Host code writer</td>
<td>▶ Program flow logic&lt;br&gt;▶ Manage accelerators&lt;br&gt;▶ Work blocks creation, input and output data specified as a series of address-type-length entries&lt;br&gt;▶ Manage communication and synchronization with peer host tasks</td>
</tr>
<tr>
<td>Accelerator code writer</td>
<td>Computational kernel</td>
</tr>
<tr>
<td>ALF run time</td>
<td>▶ Schedule work blocks to accelerators&lt;br&gt;▶ Data transfer</td>
</tr>
</tbody>
</table>

The ALF also offers more sophisticated mechanisms for managing multiple computational kernels or express dependencies, or to further tune the data movement. As with DaCS, the ALF can operate inside a Cell/B.E. server or in hybrid mode.


**IBM Dynamic Application Virtualization**

With Dynamic Application Virtualization (DAV), an IBM alphaWorks offering, an application can benefit from Cell/B.E. acceleration without any source code changes. The original, untouched application is directed to use a stub library that is dynamically loaded and offloads the compute intense functions to a Cell/B.E system. DAV currently supports C/C++ or Visual Basic® Applications, such as Microsoft® Excel® 2007 spreadsheets, that run under the Microsoft Windows® operating system.
DAV comes with tools to generate ad-hoc stub libraries based on the prototypes of the offloaded functions on the client side and similar information about the server side (the Cell/B.E. system) where the functions are implemented. For the main application, the Cell/B.E. system is completely hidden. The actual implementation of the function on the Cell/B.E. system uses the existing programming frameworks to maximize the application performance.

See 7.4, “Dynamic Application Virtualization” on page 475, for a more complete description.

**Workload specific libraries**

The IBM SDK for Multicore Acceleration contains workload specialized libraries. The libraries are the BLAS library for linear algebra, libFFT for FFT in 1D and 2D, and libmc for random number generations.

**OpenMP**

The IBM SDK for Multicore Acceleration contains a technology preview of the XL C/C++ single source compiler. Usage of this compiler completely hides the Cell/B.E. system to the application programmer who can continue by using OpenMP, which is a familiar shared memory parallel programming model. The compiler runtime library takes care of spawning threads of execution on the SPEs and manages the data movement and synchronization of the PPE threads to SPE threads.

Other groups or companies are working on providing programming frameworks for the Cell/B.E. platform as discussed in the following sections.

**Mercury Computer Systems**

Mercury has two main offerings for the Cell/B.E. platform:

- The MultiCore Framework (MCF), which implements the manager/worker model with an input/output tile management similar to the ALF work blocks
- Multicore Plus SDK, which bundles MCF with additional signal processing and FFT libraries (SAL, Vector Signal and Image Processing Library (VSIPL)), a trace library (TATL), and open source tools for MPI communications (OpenMPI) and debugging (gdb)

**PeakStream**

The PeakStream platform offers an API, a generalize array type, and a virtual machine environment that abstracts the programmer from the real hardware. Data is moved back and forth between the application and the virtual machine that accesses the Cell/B.E. resources by using an I/O interface. All the work in the virtual machine is asynchronous from the main application perspective, which can keep doing work before reading the data from the virtual machine. The
PeakStream platform currently runs on the Cell/B.E. platform, GPUs, and traditional homogeneous multi-core processors.

**CodeSourcery**
CodeSourcery offers Sourcery VSIPL++, a C++ implementation of the open standard VSIPL++ library that is used in signal and image processing. The programmer is freed from accessing the low level mechanisms of the Cell/B.E. platform. This is handled by the CodeSourcery runtime library.

The VSIPL contains routines for the following items:
- Linear algebra for real and complex values
- Random numbers
- Signal and image processing (FFT, convolutions, and filters)

CodeSourcery also runs on GPU and a multi-core general purpose processor.

**Gedae**
Gedae tries to automate the software development by using a model-driven approach. The algorithm is captured in a flow diagram that is then used by the multiprocessor compiler to generate a code that will match both the target architecture and the data movements required by the application.

**RapidMind**
RapidMind works with standard C++ language constructs and augments the language by using specialized macro language functions. The whole integration involves the following steps:

1. Replace float or int arrays by RapidMind equivalent types (Array, Value).
2. Capture the computations that are enclosed between the RapidMind keywords Program BEGIN and END and convert them into object modules.
3. Stream the recorded computations to the underlying hardware by using platform-specific constructs, such as Cell/B.E., processor, or GPU, when the modules are invoked.

There are also research groups working on implementing other frameworks onto the Cell/B.E. platform. It is worth noting the efforts of the Barcelona Supercomputing teams with CellSs (Cell Superscalar) and derivatives, such as SMPSs (SMP Superscalar), and from Los Alamos National Laboratory with CellFS, based on concepts taken from the Plan9 operating system.
Summary
In Figure 3-3, we plot these frameworks on a scale ranging from the closest to
the hardware to the most abstract.

IBM DAV is of particular note here. On the accelerated program side (the client
side in DAV terminology), the Cell/B.E. system is completely hidden by using the
stub dynamic link library (DLL) mechanism. On the accelerator side (the server
side for DAV), any Cell/B.E. programming model can be used to implement the
functions that have been offloaded from the client application.
3.2 Determining whether the Cell/B.E. system fits the application requirements

We use the decision tree in Figure 3-4 to help us determine whether the Cell/B.E. system fits the application requirements.

![Decision Tree Diagram]

Figure 3-4 Determining whether the Cell/B.E. system is a good fit for this application
3.2.1 Higher performance/watt

The main driver for enabling applications on the Cell/B.E. system is the need for a higher level of performance per watt. This is a concern shared by many customers as reported by the IDC study referenced in Solutions for the Datacenter’s Thermal Challenges [14 on page 624]. Customers may be willing to make the following accommodations:

- Lower their electricity bills.
- Overcome computer rooms limits in space, power, and cooling.
- Adopt a green strategy for their IT, a green “ITtude.”
- Allow for more computing power for a given space and electrical power budget as is often the case in embedded computing.

The design choices for the Cell/B.E. system exactly match these new requirements with a power efficiency (expressed in peak GFlops per Watt) that is over two times better than conventional general purpose processors.

3.2.2 Opportunities for parallelism

The Cell/B.E. server offers parallelism at four levels:

- Across multiple IBM System x™ servers in a hybrid environment
  This is expressed using MPI at the cluster level or some sort of grid computing middleware.
- Across multiple Cell/B.E. chips or servers
  Here we use MPI communication between the Cell/B.E. servers in the case of a homogeneous cluster of stand-alone Cell/B.E. servers or possibly ALF or DaCS for hybrid clusters.
- Across multiple SPE inside the Cell/B.E. chip or server by using libspe2, ALF, DaCS, or a single source compiler
- At the word level with SIMD instructions on each SPE, by using SIMD intrinsics or the “auto-SIMDization” capabilities of the compilers

The more parallel processing opportunities the application can leverage, the better.
3.2.3 Algorithm match

For the algorithm match, we are looking for a match between the main computational kernels of the application and the Cell/B.E. strengths as listed in Table 3-3 on page 37. As we have seen in 3.1.1, “The computation kernels” on page 34, most applications can be characterized by a composition of the “13 dwarfs” of Patterson et al. in *The Landscape of Parallel Computing Research: A View from Berkeley* [1 on page 623]. Therefore, it is important to know with which kernels a given application is built. This is usually easy to do because it is related to the numerical methods that are used in the applications.

In the paper *Scientific Computing Kernels on the Cell Processor*, by Williams et al. [2 on page 623], the authors studied how the Cell/B.E. system performs on four of the 13 dwarfs, specifically dense matrices algebra, sparse matrices algebra, spectral methods, and structured grids. They compared the performance of these kernels on a Cell/B.E. server with what they obtained on a superscalar processor (AMD Opteron™), a VLIW processor (Intel Itanium2), and a vector processor (Cray X1E). The Cell/B.E. system had good results because these kernels are extremely common in many HPC applications.

Other authors have reported successes for graphical models (bioinformatics, HMMer [15 on page 624]), dynamic programming (genomics, BLAST [16 on page 624]), unstructured grids (Finite Element Solvers [17 on page 624], and combinatorial logic (AES, DES [18 on page 624]).

The map-reduce dwarf is parallel and, therefore, a perfect fit for the Cell/B.E. system. Examples can be found in ray-tracing or Monte-Carlo simulations.

The graph traversal dwarf is a more difficult target for the Cell/B.E. system due to random memory accesses although some new sorting algorithms (AA-sort in [5 on page 623]) have been shown to exploit the CBEA.

The N-Body simulation does not seem yet ready for Cell/B.E. exploitation although research efforts are providing good early results [19 on page 624].

Table 3-6 on page 51 summarizes the results of these studies. We present each of the “13 dwarfs”, their Cell/B.E. affinity (from 1 (poor) to 5 (excellent)), and the Cell/B.E. features that are of most value for each kernel.

The algorithm match also depends on the data types that are being used. The current Cell/B.E. system has a single precision floating point affinity. There will be much larger memory and enhanced double-precision floating point capabilities in later versions of the Cell/B.E. system.
<table>
<thead>
<tr>
<th>Dwarf name</th>
<th>Cell/B.E. affinity (1 = poor to 5 = excellent)</th>
<th>Main Cell/B.E. features</th>
</tr>
</thead>
</table>
| Dense matrices     | 5                                             | ▶ Eight SPE per Cell/B.E. system  
▶ SIMD  
▶ Large register file for deep unrolling  
▶ Fused multiply-add |
| Sparse matrices    | 4                                             | ▶ Eight SPE per Cell/B.E. system  
▶ Memory latency hiding with DMA  
▶ High memory sustainable load |
| Spectral methods   | 5                                             | ▶ Eight SPE per Cell/B.E. system  
▶ Large register file  
▶ Six cycles local storage latency  
▶ Memory latency hiding with DMA |
| N-body methods     | To be determined (TBD)                         | TBD                                                                                      |
| Structured grids   | 5                                             | ▶ Eight SPE per Cell/B.E. system  
▶ SIMD  
▶ High memory bandwidth  
▶ Memory latency hiding with DMA |
| Unstructured grids | 3                                             | ▶ Eight SPE per Cell/B.E. system  
▶ High memory thruput |
| Map-reduce         | 5                                             | ▶ Eight SPE per Cell/B.E. system |
| Combinatorial logic| 4                                             | ▶ Large register file |
| Graph traversal    | 2                                             | ▶ Memory latency hiding |
| Dynamic programming| 4                                             | ▶ SIMD |
| Back-track and Branch-Bound | TBD | TBD |
| Graphical models   | 5                                             | ▶ Eight SPE per Cell/B.E. system  
▶ SIMD |
| Finite state machine| TBD                                           | TBD                                                                                      |
As can be derived from Table 3-6 on page 51, the Cell/B.E. system is a good match for many of the common computational kernels. This result is based on the design decisions that were made to address the following main bottlenecks:

- Memory latency and throughput
- Very high computational density with eight SPEs per Cell/B.E. system, each with a large register file and an extremely low local storage latency (6 cycles compared to 15 for current general purpose processors from Intel or AMD)

### 3.2.4 Deciding whether you are ready to make the effort

The Cell/B.E. system may be easy on the electricity bill, but it can be difficult on the programmer. The enablement of an application on the Cell/B.E. system can result in substantial algorithmic and coding work, which is usually worth the effort.

What are the alternatives? The parallelization effort may have already been done by using OpenMP at the process level. In this case, using the prototype of the XLC single source compiler might be the only viable alternative. Despite high usability, these compilers are still far from providing the level of performance that can be attained with native SPE programming. The portability of the code is maintained, which for some customers, might be a key requirement.

For new developments, it might be a good idea to use the higher level of abstraction provided by such technologies from PeakStream, RapidMind, or StreamIt. The portability of the code is maintained between the Cell/B.E., GPU, and general multi-core processors. However, the application is tied to the development environment, which is a different form of lock-in.

In the long run, new standardized languages may emerge. Such projects as X10 from IBM or Chapel from Cray, Inc. might become the preferred language for writing applications to run on massively multi-core systems. Adopting new languages has historically been a slow process. Even if we get a new language, that still does not help the millions of lines of code written in C/C++ and Fortran. The standard API for the host-accelerator model may be closer. ALF is a good candidate. The fast adoption of MPI in the mid-1990s has proven that an API can be what we need to enable a wide range of applications.

Can we wait for these languages and standards to emerge? If the answer is “no” and the decision has been made to enable the application on the Cell/B.E. system now, there is a list of items to consider and possible workarounds when problems are encountered as highlighted in Table 3-7 on page 53.

---

3 http://www.cag.csail.mit.edu/streamit
5 http://chapel.cs.washington.edu
Table 3-7  Considerations when enabling an application on the Cell/B.E. system

<table>
<thead>
<tr>
<th>Topic</th>
<th>Potential problem or problems</th>
<th>Workaround</th>
</tr>
</thead>
</table>
| Source code changes | ▶ Portability concerns  
▶ Limits the scope of code changes                                                              | The Cell/B.E. APIs are standard C. Approaches such as a host-accelerator can limit the amount of source code changes. |
| Operating systems   | ▶ Windows applications                                                                         | The Cell/B.E. system runs Linux only. If the application runs on Windows, we may want to use DAV to offload only the computational part to the Cell/B.E. system. |
| Languages           | ▶ C/C++ fully supported  
▶ Fortran and ADA supported  
▶ Other languages not supported                                                              | Rewrite the compute-intensive part in C. Use some type of offloading for Java™ or VBA applications running on Windows with DAV. |
| Libraries           | ▶ Not many libraries supported yet  
▶ Little ISV support                                                                             | Use the workload libraries that are provided by the IBM SDK for Multicore Acceleration. |
| Data types          | ▶ 8-bit, 16-bit, and 32-bit data well supported  
▶ 64-bit float point supported                                                                | Full speed double-precision support is soon to be available. |
| Memory requirements | ▶ Maximum of 2 GB per blade server                                                              | Use more smaller MPI tasks. On an IBM Blade Server, use a single MPI task with 16 SPEs rather than two MPI tasks with eight SPEs. (This is subject to change because a much larger memory configuration per blade is due in future product releases.) |
| Memory requirements | ▶ Local storage (LS) size of 256k                                                              | Large functions need to be split. We will have to use overlay. Limit recursions (stack space). |
| I/O                 | ▶ I/O intensive tasks                                                                           | The Cell/B.E. system does not help I/O bound workloads. |

### 3.3 Deciding which parallel programming model to use

Large homogeneous compute clusters can be built by collecting stand-alone Cell/B.E. blade servers with an InfiniBand interconnect. At this cluster level, the usual distributed memory programming models, such as MPI, can be used. An application that is already programmed by using MPI is a good start because we only need to add Cell/B.E. parallelism incrementally to fully exploit the Cell/B.E. potential.
Hybrid clusters are becoming increasingly popular as a means of building powerful configurations by incrementally upgrading existing clusters built with off-the-shelf components. In this model, the MPI parallelism at the cluster level is maintained, but each task is now accelerated by one or more Cell/B.E. systems.

We first describe the parallel programming models that are found in the literature and then focus on the Cell/B.E. chip or board-level parallelism and the host-accelerator model.

### 3.3.1 Parallel programming models basics

Mattson et al. in *Patterns for Parallel Programming* [4 on page 623] define a taxonomy of parallel programming models. First they define four “spaces” (described in Table 3-8) that the application programmer must visit.

**Table 3-8** Four design spaces from Mattson et al.

<table>
<thead>
<tr>
<th>Space</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Finding concurrency</td>
<td>Find parallel tasks and group and order them</td>
</tr>
<tr>
<td>Algorithm structure</td>
<td>Organize the tasks in processes</td>
</tr>
<tr>
<td>Supporting structure</td>
<td>Code structures for tasks and data</td>
</tr>
<tr>
<td>Implementation mechanisms</td>
<td>Low level mechanisms for managing and synchronizing execution threads as well as data communication</td>
</tr>
</tbody>
</table>

The first space is application dependent. The implementation mechanisms are described in more detail in Chapter 4, “Cell Broadband Engine programming” on page 75, and Chapter 7, “Programming in distributed environments” on page 445.

In the algorithm space, Mattson et al. propose to look at three different ways of decomposing the work, each with two modalities. This leads to six major algorithm structures, which are described in Table 3-9.

**Table 3-9** Algorithm structures

<table>
<thead>
<tr>
<th>Organization principle</th>
<th>Organization subtype</th>
<th>Algorithm structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>By tasks</td>
<td>Linear</td>
<td>Task parallelism</td>
</tr>
<tr>
<td></td>
<td>Recursive</td>
<td>Divide and conquer</td>
</tr>
<tr>
<td>By data decomposition</td>
<td>Linear</td>
<td>Geometric decomposition</td>
</tr>
<tr>
<td></td>
<td>Recursive</td>
<td>Tree</td>
</tr>
</tbody>
</table>
Task parallelism occurs when multiple independent tasks can be scheduled in parallel. The divide and conquer structure is applied when a problem can be recursively treated by solving smaller subproblems. Geometric decomposition is common when we try to solve a partial differential equation that has been made discrete on a 2-D or 3-D grid, and grid regions are assigned to processors.

As for the supporting structures, Mattson et al. identified four structures for organizing tasks and three for organizing data. They are given side by side in Table 3-10.6

<table>
<thead>
<tr>
<th>Code structures</th>
<th>Data structures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single Program Multiple Data (SPMD)</td>
<td>Shared data</td>
</tr>
<tr>
<td>Master/worker</td>
<td>Shared queue</td>
</tr>
<tr>
<td>Loop parallelism</td>
<td>Distributed array</td>
</tr>
<tr>
<td>Fork/join</td>
<td></td>
</tr>
</tbody>
</table>

SPMD is a code structure that is well-known to MPI programmers. Although MPI does not impose the use of SPMD, this is a frequent construct. Master/worker is sometimes called “bag of tasks” when a master task distributes work elements independently of each other to a pool of workers. Loop parallelism is a low-level structure where the iterations of a loop are shared between execution threads. Fork/join is a model where a master execution thread calls (fork) multiple parallel execution threads and waits for their completion (join) before continuing with the sequential execution.

Shared data refers to the constructs that are necessary to share data between execution threads. Shared queue is the coordination among tasks to process a queue of work items. Distributed array addresses the decomposition of multi-dimensional arrays into smaller sub-arrays that are spread across multiple execution units.

---

We now look at how these constructs map to the Cell/B.E. system and what we must examine to determine the best parallel programming model for the application. The following forces (provided in no particular order of importance) are specific to the Cell/B.E. system that influences the choice:

- Heterogenous PPE/SPE
- Distributed memory between the SPE, shared memory view possible by memory mapping the local storage
- SIMD
- PPE slow compared to SPE
- Software managed memory hierarchy
- Limited size of the LS
- Dynamic code loading (overlay)
- High bandwidth on the EIB
- Coherent shared memory
- Large SPE context, startup time

### 3.3.2 Chip or board level parallelism

The Cell/B.E. processor is heterogenous and multi-core, with distributed memory. It offers many opportunities for parallel processing at the chip or board level. Figure 3-5 shows the reach of multiple programming models. The models can sometimes be classified as PPE-centric or SPE-centric. Although this is a somewhat artificial distinction, it indicates that the application control is either run more on the PPE side or on the SPE side.

*Figure 3-5 Various models on the Cell/B.E. processor*
Figure 3-5 shows the following different models:

- A small single SPE program, where the whole code holds in the local storage of a single SPE
- A large single SPE program, one SPE program accessing system memory
- A small multi-SPE program
- A general Cell/B.E. program with multiple SPE and PPE threads

When multiple SPEs are used, they can be arranged in a data parallel, streaming mode, as illustrated in Figure 3-6.

![Diagram](image)

Figure 3-6 The streaming model

Each piece of input data (I0, I1, ...) is streamed through one SPE to produce a piece of output data (O0, O1, and so on). The exact same code runs on all SPEs. Sophisticated load balancing mechanisms can be applied here to account for differing compute time per data chunk.

Chapter 3. Enabling applications on the Cell Broadband Engine hardware
The SPE can also be arranged in a pipeline fashion, where the same piece of data undergoes various transformations as it moves from one SPE to the other. Figure 3-7 shows a general pipeline.

![Diagram of pipeline model]

The main benefit is that we aggregate the code size of all the SPEs that are participating in the pipeline. We also benefit from the huge EIB bandwidth to transfer the data. A possible variation is to move the code instead of the data, whichever is the easiest or smallest to move around. Good load balancing is much more challenging because it relies on constant per stage computational time.

3.3.3 More on the host-accelerator model

A common approach with Cell/B.E. parallel programming is to use a function offload mechanism similar to the RPC model. The application flows on the PPE, and only for selected, highly computational kernels do we call upon SPE
acceleration. This method is the easiest from a program development perspective because it limits the scope of source code changes and does not require much re-engineering at the application logic level. It is much a fork/join model and, therefore, requires care in giving enough work to the accelerator threads to compensate for the startup time. This method is typically implemented with specialized workload libraries such as BLAS, FFT, or RNG for which exists a Cell/B.E.-tuned version. BLAS is the only library that can be considered a “drop in replacement” at this time.

A variation of this model is to have general purpose accelerator programs running on the SPE, sitting in a loop, awaiting to be asked to provide services for the PPE threads. The existence of persistent threads on each SPE eliminates the startup time of SPE threads but requires that the accelerator programs be able to service various requests for various functions, possibly incurring the use of dynamic code uploading techniques. The ALF framework described in 4.7.2, “Accelerated Library Framework” on page 298, and 7.3, “Hybrid ALF” on page 463, is one implementation.

Figure 3-8 shows a general accelerator.

![Memory structure of an accelerator](image)

An accelerator can implement multiple services (functions f1 to f4). Each function may require some “state data” that is persistent across multiple invocations. The “live data” is the data in and out of the accelerator for each invocation. It is important to understand which is read, written, or both to optimize the data transfers.
### 3.3.4 Summary

Of the various programming models and structures listed in Table 3-9 on page 54 and Table 3-10 on page 55, some are easier to implement on the Cell/B.E. platform than others. Table 3-11, Table 3-12, and Table 3-13 on page 61 summarize the Cell/B.E. specific issues.

**Table 3-11**  
**Cell/B.E. suitability of algorithm structures and specific issues**

<table>
<thead>
<tr>
<th>Algorithm structure</th>
<th>Cell/B.E. suitability (1 = poor and 5 = excellent)</th>
<th>Things to look at</th>
</tr>
</thead>
<tbody>
<tr>
<td>Task parallelism</td>
<td>5</td>
<td>Load balancing; synchronization required to access the queue of work items</td>
</tr>
<tr>
<td>Divide and conquer</td>
<td>TBD</td>
<td>TBD</td>
</tr>
<tr>
<td>Geometric decomposition</td>
<td>5</td>
<td>DMA and double buffering are required; code size</td>
</tr>
<tr>
<td>Tree</td>
<td>3</td>
<td>Random memory access.</td>
</tr>
<tr>
<td>Pipeline</td>
<td>5</td>
<td>▶ Load balancing</td>
</tr>
<tr>
<td></td>
<td></td>
<td>▶ EIB exploitation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>▶ Whether to move code or data</td>
</tr>
<tr>
<td>Event-based coordination</td>
<td>3</td>
<td>Code size; the resulting code might be inefficient because the operation that is</td>
</tr>
<tr>
<td></td>
<td></td>
<td>required is not known until we get the event data to process.</td>
</tr>
</tbody>
</table>

**Table 3-12**  
**Cell/B.E. suitability of code structures and specific issues**

<table>
<thead>
<tr>
<th>Code structure</th>
<th>Cell/B.E. suitability (1 = poor and 5 = excellent)</th>
<th>Things to look at</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPMD</td>
<td>3</td>
<td>Code size; the whole application control may not fit in local storage and more PPE</td>
</tr>
<tr>
<td></td>
<td></td>
<td>intervention may be required.</td>
</tr>
<tr>
<td>Master/worker</td>
<td>5</td>
<td>Load balancing; synchronization is required to access the queue of work items.</td>
</tr>
</tbody>
</table>
3.4 Deciding which Cell/B.E. programming framework to use

We have now found the parallel programming model (work distribution, data distribution, and synchronization) that we want to apply to the application. We can draw on more than one model for a given application. We must implement these models by using the available Cell/B.E. frameworks. Some frameworks are general. The most versatile is libspe2 with MFC services accessed through direct problem state (PPE) and channels (SPE). In addition, some frameworks are specialized, such as workload libraries.

Choosing a framework is a matter of weighing the features, the ease of implementation, and the performance as well as application area specifics. For example, for radar applications, Gedae seems to be almost mandatory.

<table>
<thead>
<tr>
<th>Code structure</th>
<th>Cell/B.E. suitability (1 = poor and 5 = excellent)</th>
<th>Things to look at</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loop parallelism</td>
<td>3</td>
<td>▶ PPE centric</td>
</tr>
<tr>
<td></td>
<td></td>
<td>▶ Task granularity</td>
</tr>
<tr>
<td></td>
<td></td>
<td>▶ Shared memory synchronization</td>
</tr>
<tr>
<td>Fork/join</td>
<td>5</td>
<td>Fits the accelerator model; weigh the thread startup time with the amount of work</td>
</tr>
<tr>
<td></td>
<td></td>
<td>and data transfer that are needed.</td>
</tr>
</tbody>
</table>

Table 3-13  Data structures and Cell/B.E. specific issues

<table>
<thead>
<tr>
<th>Data structure</th>
<th>Things to look at</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shared data</td>
<td>Synchronization for accessing shared data, memory ordering, and locks</td>
</tr>
<tr>
<td>Shared queue</td>
<td>From PPE managed to SPE self-managed work queues</td>
</tr>
<tr>
<td>Distributed array</td>
<td>Data partitioning and DMA</td>
</tr>
</tbody>
</table>
In Table 3-14, we list the various parallel programming constructs. For each one, we indicate the most appropriate primary and secondary choices of the frameworks.

<table>
<thead>
<tr>
<th>Programming construct</th>
<th>Primary</th>
<th>Secondary, comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>MPI</td>
<td>OpenMPI, nothing specific to the Cell/B.E. server. This is a cluster/PPE level construct.</td>
<td>MVAPICH, MPICH</td>
</tr>
<tr>
<td>pthreads</td>
<td>pthreads is supported on the PPE. No direct support for a mapping between PPE pthreads and SPE pthreads. Must be implemented by using libspe.</td>
<td></td>
</tr>
<tr>
<td>OpenMP</td>
<td>XLC single source compiler</td>
<td></td>
</tr>
<tr>
<td>UPC, CAF</td>
<td>Not supported</td>
<td></td>
</tr>
<tr>
<td>X10</td>
<td>Not supported</td>
<td></td>
</tr>
<tr>
<td>Task parallelism</td>
<td>libspe</td>
<td>This is function offload; see Fork/join too</td>
</tr>
<tr>
<td>Divide and conquer</td>
<td>TBD</td>
<td>TBD</td>
</tr>
<tr>
<td>Geometric decomposition</td>
<td>ALF if data blocks can be processed independently</td>
<td>DaCS for more general data decomposition</td>
</tr>
<tr>
<td>Tree</td>
<td>TBD; software cache may be a good choice</td>
<td></td>
</tr>
<tr>
<td>Pipeline</td>
<td>libspe</td>
<td>DaCS, Streamit</td>
</tr>
<tr>
<td>Event-based coordination</td>
<td>libspe</td>
<td></td>
</tr>
<tr>
<td>SPMD</td>
<td>This is a PPE level construct.</td>
<td></td>
</tr>
<tr>
<td>Master/worker</td>
<td>ALF</td>
<td>libspe</td>
</tr>
<tr>
<td>Loop parallelism</td>
<td>XLC single source compiler, with OpenMP support</td>
<td></td>
</tr>
<tr>
<td>Fork/join</td>
<td>Workload specific libraries if they exist and ALF otherwise. ALF can be used to create new workload specific libraries.</td>
<td>This is the accelerator model. Use DAV if necessary to accelerate a Windows application.</td>
</tr>
<tr>
<td>Shared data</td>
<td>DaCS</td>
<td>MFC intrinsics, libsync</td>
</tr>
<tr>
<td>Shared queue</td>
<td>DaCS</td>
<td>MFC intrinsics, libsync</td>
</tr>
<tr>
<td>Distributed array</td>
<td>ALF</td>
<td>DaCS</td>
</tr>
</tbody>
</table>
### 3.5 The application enablement process

The process of enabling an application on the Cell/B.E. system (Figure 3-9) can be incremental and iterative. It is incremental in the sense that the hotspots of the application should be moved progressively off the PPE to the SPE. It is iterative for each hotspot. The optimization can be refined at the SIMD, synchronization, and data movement levels until satisfactory levels of performance are obtained.

As for the starting point, a thorough profiling of the application on a general purpose system (PPE is acceptable) gives all the hotspots that need to be examined. Then, for each hotspot, we can write a multi-SPE implementation with all the data transfer and synchronization between the PPE and the SPE. After this first implementation is working, we then turn to the SIMDization and tuning of the SPE code. The last two steps can be repeated in a tight loop until we obtain good performance. We can repeat the same process for all the major hotspots.

![Figure 3-9 General flow for enabling an application on the Cell/B.E. system](image-url)
The flow in Figure 3-9 on page 63 will change a bit depending on the framework that is chosen. If we are fortunate to have an application whose execution time is dominated by a function that happens to be part of a workload specific library that is ported to the Cell/B.E. system, then we follow the process shown in Figure 3-10.

Figure 3-10  Implementing the Cell/B.E. system-tuned, workload-specific libraries
Figure 3-11 shows the process for ALF.

Figure 3-11  Enabling a Cell/B.E. application with ALF
### 3.5.1 Performance tuning for Cell/B.E. programs

The goal of enabling applications on the Cell/B.E. system is to achieve the best performance, which does not come for free. Performance tuning is an integral part of the application enablement. Chapter 4, “Cell Broadband Engine programming” on page 75, and Chapter 7, “Programming in distributed environments” on page 445, provide detailed performance tips for writing good SPE and PPE code. Mike Acton in “Developing Technology for Ratchet and Clank Future: Tools of Destruction” [24 on page 625] gives valuable advice about using the experience he and his team at Insomniac Games gathered in the process of developing video games for the Sony Playstation 3. He makes the following recommendations:

- Let us not hide the CBEA but exploit it instead.
- For a successful port to the CBEA, we must do as follows:
  - Understand the architecture.
  - Understand the data, including the movement, dependencies, generation, and usage (read, write, or read-write).
  - Do the hard work.
- Put more work on the SPE, and do less on the PPE.
- Do not to view the SPE as co-processors but rather view the PPE as a service provider for the SPE.
- Ban scalar code on the SPE.
- With less PPE/SPE synchronization, use deferred updates, lock-free synchronization (see 4.5, “Shared storage synchronizing and data ordering” on page 218) and perform dataflow management as much as possible from the SPE.

### 3.6 A few scenarios

In this section, we review examples of Cell/B.E. programs. Figure 3-12 on page 67 shows the program flow for a typical function offload to a workload library. We illustrate the PPE thread as running useful work until it calls a function that is part of the Cell/B.E.-tuned library. Then the library starts the SPE context and executes the library code. The library on the SPE might be doing any kind of inter-SPE communication, DMA accesses, and so on as indicated by the cloud. After the function finishes executing, the SPE contexts are terminated, and the PPE thread resumes execution on the PPE.

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7 [http://well.cellperformance.com](http://well.cellperformance.com)
Starting and terminating SPE contexts takes time. We must ensure that the time spent in the library far exceeds the SPE context startup time.

A variation of this scheme is when the application calls the library repeatedly. In this case, it might be interesting to keep the library contexts running on the SPE and set them to work with a lightweight mailbox operation, for example, as shown in Figure 3-13 on page 68.

Figure 3-12  A typical workflow for Cell/B.E. tuned workload libraries
Figure 3-13  Successive invocations of the same library
Figure 3-13 on page 68 shows three successive invocations of the library with data1, data2 and data3. The dotted lines indicate an SPE context that is active but waiting. This arrangement minimizes the impact of the SPE contexts creation but can only work if the application has a single computational kernel that is called repeatedly.

Figure 3-14 shows the typical workflow of an ALF application. The PPE thread prepares the work blocks (numbered wb0 to wb12 here), which execute on the SPE accelerators.
Typically no communication exists between accelerators when they are processing a work block and the ALF run time takes care of balancing the work among the accelerators. Also, the PPE thread might do useful work while the accelerators are crunching though the work blocks.

### 3.7 Design patterns for Cell/B.E. programming

Design patterns were first introduced by Christopher Alexander in the field of town and building architecture. Gamma et al. in *Design Patterns. Elements of Reusable Object-Oriented Software* [3 on page 623] applied the same principles to computer programming, and they have proven to be a useful tool since then. Multiple definitions can be found for a pattern. In *Patterns for Parallel Programming* [4 on page 623], Mattson et al. define a pattern as a “good solution to a recurring problem in a particular context”. Marc Snir, in “Programming Design Patterns, Patterns for High Performance Computing” [6 on page 623], describes them as a “way to encode expertise”. Patterns are usually characterized by the following items:

- A name
- A problem
- The forces that are shaping the solution
- A solution to the problem

By using the same formalism, we started to build a list of design patterns that apply to Cell/B.E. programming. This is clearly only a start, and we hope that new patterns will emerge as we gain more expertise in porting code to the Cell/B.E. environment.

We look at the following design patterns:

- A shared queue
- Indirect addressing
- A pipeline
- Multi-SPE software cache
- Plug-in

#### 3.7.1 Shared queue

We want to distribute work elements between multiple SPEs. They are arranged in a first in, first out (FIFO) queue in PPE memory.
Forces
The two main forces are the need for a good load balance between the SPE and minimal contention.

Solution
We can envision three solutions for dividing work between SPEs. They are described in the following sections.

Fixed work assignment
Each SPE is statically assigned the same amount of work elements. This incurs no contention but may be a weak scheme if the time taken to process a work element is not constant.

Master/worker
The PPE assigns the work elements to the SPE. The PPE can give the pieces of work one at a time to the SPE. When an SPE is finished with its piece of work, it signals the PPE, which then feeds the calling SPE with a new item, automatically balancing the work. This scheme is good for the load balance, but may lead to contention if many SPE are being used because the PPE might be overwhelmed by the task of assigning the work blocks.

Self-managed by the SPE
The SPE synchronizes between themselves without PPE intervention. After an SPE is finished with its work item, it grabs the next piece of work from the queue and processes it. This is the best scheme because it ensures good load balance and does not put any load on the PPE. The critical part of the scheme is to ensure that the SPE removes work items off the queue atomically, by using either of the following methods:

- The MFC atomic operations
- The features from the sync library that are provided with the IBM SDK for Multicore Acceleration

3.7.2 Indirect addressing
We want to load in SPE memory a vector that is addressed through an index array. This is common in a sparse matrix-vector product that arises, for example, when solving linear systems with conjugate gradient methods. Example 3-1 on page 72 shows the typical construct in which the matrix is stored in the Compressed Sparse Row (CSR) format. This storage is described in Optimization of Sparse Matrix-vector Multiplication on Emerging Multicore Platforms [25 on page 625].
Example 3-1  Matrix-vector product with a sparse matrix

```c
float A[], x[], y[];
int ja[], idx[];
for(i=0; i<N; i++) {
    for(j=ja[i]; j<ja[i+1]; j++) {
    }
}
```

**Forces**
The index array by which the x vector is accessed leads to random memory accesses.

**Solution**
The index array is known in advance, and we can exploit this information by using software pipelining with a multi-buffering scheme and DMA lists as shown in Figure 3-15.

![Software pipeling and multi-buffering](image)

Get IDXi : DMA the ith block of the IDX array
Pr DMALi : Prepare the DMA list for the ith access to the X array
Gt DMALi : DMA the list for the ith access to the X array
Work X i : work with the ith access to the X array

We do not to show the accesses to matrix A and array y. They are accessed sequentially, and a simple multi-buffering scheme can also be applied.
3.7.3 Pipeline

We want to arrange SPE contexts in a multi-stage pipeline manner.

Forces
We want to minimize the time it takes for the data to move from one pipeline stage to the other.

Solution
By using the affinity functions as described in 4.3.4, “Direct problem state access and LS-to-LS transfer” on page 144, we can make sure that successive SPE contexts are ideally placed on the EIB to maximize the LS-to-LS transfer speed. An alternative arrangement is to move the code instead of the data, whichever is the fastest. Often when the programming model is a pipeline, some state data must reside on each stage, and moving the function also requires moving the state data.

3.7.4 Multi-SPE software cache

We want to define a large software cache that gathers LS space from multiple participating SPEs.

Forces
We want to push the software cache a bit further by allowing data to be cached not necessarily in the SPE that encounters a “miss” but also in the LS of another SPE. The idea is to exploit the high EIB bandwidth.

Solution
We do not have a solution for this yet. The first step is to look at the cache coherency protocols (MESI, MOESI, and MESIF)\(^8\) that are in use today on multiprocessor systems and try to adapt them to the Cell/B.E. system.

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\(^8\) M=Modified, O=Owner, E=Exclusive, S=Shared, I=Invalid, F=Forwarding
3.7.5 Plug-in

We want to process data whose contents, and therefore its associated treatment, are discovered on the go.

Forces
The forces are similar to what happens with a Web browser when the flow of data coming from the Web server contains data that requires the loading of external plug-ins to be displayed. The challenge is to load on the SPE both the data and the code to process it when the data is discovered.

Solution
The Cell/B.E. system has overlay support already, which is one solution. However, there might be a better solution to this particular problem by using dynamically loaded code. We can imagine loading code together with data by using exactly the same DMA functions. Nothing in the SPE memory differentiates code from data. This has been implemented successfully by Eric Christensen et al. in *Dynamic Code Uploading on the SPU* [26 on page 625] by using the following process:

1. Compile the code.
2. Dump the object code as binary.
3. Load the binary code as data.
4. Use DMA on the data (containing the code) just as with regular data to the SPE. Actual data can also be loaded during the same DMA operation.
5. On the SPE, jump to the address location where the code has been loaded to pass the control to the plug-in that has just been loaded.
The goal of this chapter is to provide an introductory guide for how to program an application on Cell Broadband Engine (Cell/B.E.). In this chapter, we cover aspects of Cell/B.E. programming, from low-level programming by using intrinsics to higher-level programming by using frameworks that hide the processor-unique architecture. We also discuss issues that are related to programming a single Cell/B.E. processor or base blade system, such as the QS21, that contains two Cell/B.E. processors but shares the same operating system and memory map.

When describing the programming techniques, we try to balance between two opposite and complementary approaches:

- Use as high-level programming as possible to reduce development time and produce code that is readable and simple. You can do this type of programming, for example, by using the C functions of the IBM Software Development Kit (SDK) version 3.0 for Multicore Acceleration to access the different Cell/B.E. hardware mechanisms. Such mechanisms include direct memory access (DMA), mailboxes, and signals. You can also use the abstract high-level libraries of the SDK to manage the work with the Cell/B.E. processor, for example, Data Communication and Synchronization (DaCS), Accelerated Library Framework (ALF), and software managed cache.
Use low-level programming in sections of the code where performance is critical. You can do this, for example, by using the low-level intrinsics that are mapped to a single or small number of assembly instructions.

When describing these techniques, we usually emphasize the cases in which each of the approaches is suitable.

In addition, we tried to include a wide range of important issues related to Cell/B.E. programming, that until this point were described in several different documents.

The programming techniques and libraries in this chapter are divided into sections according to the functionality within the program that they perform. We hope this approach is useful for the program developer in helping you to find the corresponding subject according to the current stage in development or according to the specific part of the program that is currently implemented.

We divide this chapter into the following sections:

- In 4.1, “Task parallelism and PPE programming” on page 77, we explain how to program the Power Processor Element (PPE) and how to exploit task parallelism by distributing the work between the Synergistic Processor Elements (SPEs).

- In 4.2, “Storage domains, channels, and MMIO interfaces” on page 97, we discuss the different storage domains on the Cell/B.E. processor and how either a PPE or SPE program can access them. We also explain how to use the memory flow controller (MFC), which is the main component for communicating between the processors and transferring data using DMA. You should be familiar with this subject when deciding on the program’s data partitioning or when you need to use the MFC, as any Cell/B.E. program does.

- In 4.3, “Data transfer” on page 110, we discuss the various methods for performing data transfers in the Cell/B.E. processor between the different available memories. Obviously this is a key issue in any program development.

- In 4.4, “Inter-processor communication” on page 178, we explain how to implement communication mechanisms, such as mailbox, signals, and events, between the different processors in the Cell/B.E. system in parallel.

- In 4.5, “Shared storage synchronizing and data ordering” on page 218, we discuss how that data transfer of the different processors can be ordered and synchronized. The unique memory architecture of the Cell/B.E. processor requires the programmer to be aware of this issue, which in many cases must be handled explicitly by the program by using dedicated instructions.
In 4.6, “SPU programming” on page 244, we show how to write an optimized synergistic processor unit (SPU) program. The intention is for programming issues related only to programming the SPU itself and without interacting with external components, such as the PPE, other SPEs, or main storage.

In 4.7, “Frameworks and domain-specific libraries” on page 289, we discuss high-level programming frameworks and libraries, such as DaCS, ALF and domain specific libraries, that can reduce development efforts and hide the Cell/B.E. specific architecture. In some case, the usage of such frameworks provides similar performance as programming by using low-level libraries.

Finally, in 4.8, “Programming guidelines” on page 319, we provide a collection of programming guidelines and tips. In this section, we discuss:

- Information gathered from various resources and new items that we added
- Issues that are described in detail in other chapters and references to those chapters

We recommend that programmers read this chapter before starting to develop a new application in order to understand the different considerations that must be made.

4.1 Task parallelism and PPE programming

The Cell/B.E. processor has a single PPE that is intended primarily for running the operating system, controlling the application process, managing system resources, and managing SPE threads. The execution of any user program starts on this processor, which the PPE may later offload some of its functionality to run on one or more of the SPEs.

From a programmer’s point of view, managing the work with the SPEs is similar to working with Linux threads. Also, the SDK contains libraries that assist in managing the code that runs on the SPE and communicate with this code during execution.

The PPE itself conforms to the PowerPC Architecture so that programs written for the PowerPC 970 processor, for example, should run on the Cell/B.E. processor without modification. In addition, most programs that run on a Linux-based Power system and use the operating system (OS) facilities should work properly on a Cell/B.E.-based system. Such facilities include accessing the file system, using sockets and message passing interface (MPI) for communication with remote nodes, and managing memory allocation.

The programmer should know that usage of the operating system facilities in any Cell/B.E. application always take place on the PPE. While SPE code might use...
those facilities, such usage causes the blocking of SPU code and lets the PPE handle the system request. Only when the PPE completes handling the request, the SPE execution continues.

In this section, we cover the following topics:

- In 4.1.1, “PPE architecture and PPU programming” on page 78, we describe the PPE architecture and instruction set and general issues regarding programming code that runs on the PowerPC Processor Unit (PPU).
- In 4.1.2, “Task parallelism and managing SPE threads” on page 83, we discuss how PPU code can implement task parallelism by using SPE threads. In this section, we also explain how to create and execute those threads and how to create affinity between groups of threads.
- In 4.1.3, “Creating SPEs affinity by using a gang” on page 94, we discuss how to create affinity between SPE threads that are meant to run together.

In this section, we include the issues related to PPE programming that we found are the most important when running most Cell/B.E. applications. However, in case you are interested in learning more about this subject or need specific details that are not covered in this section, refer to the “PowerPC Processor Element” chapter in the Cell Broadband Engine Programming Handbook, Version 1.1 as a good starting point.¹

4.1.1 PPE architecture and PPU programming

Programming the PPU is similar to programming any Linux-based program that runs on a PowerPC processor system. The PPE and its PPU instructions set include the following features among others:

- A general-purpose, dual-threaded, 64-bit RISC processor
- Conformance to the PowerPC Architecture with Vector/single-instruction, multiple-data (SIMD) multimedia extensions
- Usage of 32-bit instructions that are word aligned
- Dual-threaded
- Support for Vector/SIMD Multimedia Extension (VMX) 32-bits and word-aligned instructions that work on 128-bit-wide operands
- 32 KB L1 instruction and data caches

512 KB L2 unified (instruction and data) cache
- Cache line of 128 bytes
- Instructions executed in order

The PPU supports two instruction sets:
- The PowerPC instruction set
- The Vector/SIMD Multimedia Extension instruction set

In most cases, programmers prefer to use the eight SPEs to perform the massive SIMD operations and let the PPU program manage the application flow. However, it may be useful in some cases to add some SIMD computation on the PPU.

Most of the coding for the Cell/B.E. processor is in a high-level language, such as C or C++. However, an understanding of the PPE architecture and PPU instruction sets considerably helps a developer to produce efficient, optimized code. This is particularly true because C-language internals are provided for some instruction set of the PPU. In the following section, we discuss the PPU intrinsics (C/C++ language extensions) and how to use them. We also discuss intrinsics that operate both on scalars and on vector data type.

C/C++ language extensions (intrinsics)

Intrinsics are essentially inline assembly-language instructions, in the form of function calls, that have a syntax familiar to high-level programmers who use the C language. Intrinsics provide explicit control of the PPU instructions without directly managing registers and scheduling instructions, as assembly-language programming requires. The compilers that come with the SDK package support these C-language extensions.

We discuss the two main types of PPU intrinsics in the following sections.

Scalar intrinsics

Scalar intrinsics provide a minimal set of specific intrinsics to make the PPU instruction set accessible from the C programming language. Except for __setflm, each of these intrinsics has a one-to-one assembly language mapping, unless compiled for a 32-bit application binary interface (ABI) in which the high and low halves of a 64-bit double word are maintained in separate registers.

The most useful intrinsics under this category are those related to shared memory access and synchronization and those related to cache management. Efficient use of those intrinsic can assist in improving the overall performance of the application.
Refer to “PPE ordering instructions” on page 221, which discusses some of the more important intrinsic instructions, such as `sync`, `lwsync`, `eieio`, and `isync`, that are related to shared memory access and synchronization. In addition, some of those scalar instructions provide access to the PPE registers and internal data structures, which enables the programmer to use some of the PPE facilities.

All such intrinsics are declared in the `ppu_intrinsics.h` header file that must be included in order to use the intrinsics. They may be either defined within the header as macros or implemented internally within the compiler.

By default, a call to an intrinsic with an out-of-range literal is reported by the compiler as an error. Compilers may provide an option to issue a warning for out-of-range literal values and use only the specified number of least significant bits for the out-of-range argument.

Intrinsics do not have a specific ordering unless otherwise noted. They can be optimized by the compiler and be scheduled like any other instruction.

You can find additional information about PPU scalar intrinsics in the following resources:

- The “PPU specific intrinsics” chapter of *C/C++ Language Extensions for Cell Broadband Engine Architecture* document, which contains a list of the available intrinsics and their meaning.

- The “PPE instruction sets” chapter of the *Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial* document, which provides a useful table that summarizes those intrinsics.

### Vector data types intrinsics

The vector data types intrinsics is a set of intrinsics that is provided to support the VMX instructions, which follow the AltiVec standard. The VMX model adds a set of fundamental data types, called *vector types*. The vector registers are 128 bits and can contain either sixteen 8-bit values (signed or unsigned), eight 16-bit values (signed or unsigned), four 32-bit values (signed or unsigned), or four single-precision IEEE-754 floating-point values.

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2 *C/C++ Language Extensions for Cell Broadband Engine Architecture* is available on the Web at the following address: [http://www-01.ibm.com/chips/techlib/techlib.nsf/techdocs/30B3520C93F437AB887257060006FFE5E/$file/Language_Extensions_for_CBEA_2.5.pdf](http://www-01.ibm.com/chips/techlib/techlib.nsf/techdocs/30B3520C93F437AB887257060006FFE5E/$file/Language_Extensions_for_CBEA_2.5.pdf)

The vector instructions include a reach set of operations that can be performed on those vectors. Such operations include arithmetic operations, rounding and conversion, floating-point estimate intrinsics, compare intrinsics, logical intrinsics, rotate and shift Intrinsics, load and store intrinsics, and pack and unpack intrinsics.

VMX data types and Vector/SIMD Multimedia Extension intrinsics can be used in a seamless way throughout a C-language program. The programmer does not need to set up to enter a special mode. The intrinsics may be either defined as macros within the system header file or implemented internally within the compiler.

To use the Vector/SIMD intrinsics of the PPU, the programmer should ensure the following settings:

- Include the system header file altivec.h, which defines the intrinsics.
- Set the -qaltivec and -qenablevmx flags in case XLC compilation is used.
- Set the -mabi=altivec and -maltivec flags in case GCC compilation is used.

Example 4-1 demonstrates simple PPU code that initiates two unsigned integer vectors and adds them while placing the results in a third similar vector.

```c
#include <stdio.h>
#include <altivec.h>

typedef union {
  int i[4];
  vector unsigned int v;
} vec_u;

int main()
{
  vec_u a, b, d;

  a.v = (vector unsigned int){1,2,3,4};
  b.v = (vector unsigned int){5,6,7,8};

  d.v = vec_add(a.v,b.v);
```
For additional information about PPU vector data type intrinsics, refer to the following resources:

- *Altivec Technology Programming Interface Manual*, which provides a detailed description of VMX intrinsics
- The “Vector Multimedia Extension intrinsics” chapter of the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document, which includes a list of the available intrinsics and their meaning
- The “PPE instruction sets” chapter of the *Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial* document, which includes a useful table that summarizes the intrinsics

In most cases, programmers prefer to use the eight SPEs to perform the massive SIMD operations and let the PPU program manage the application flow. For this practical reason, we do not discuss the issue of PPU Vector/SIMD operations in detail when we discuss the SPU SIMD instructions. See 4.6.4, “SIMD programming” on page 258.

However, it might be useful in some applications to add SIMD computation on the PPU. Another case when a SIMD operation may take place on the PPU side is when a programmer starts the application development on the PPU, optimizes it to use SIMD instructions, and only later ports the application to the SPU. We do not recommend the use of this approach in most cases because it consumes more development time. One reason for the additional time is that, despite the strong similarity between the Vector/SIMD instructions set of the PPU and the instruction set of the SPU, the instruction sets are different. Most of the PPU Vector/SIMD instructions have equivalent SPU SIMD instructions and vice versa.

The SDK also provides a set of header files that aim to minimize the effort when porting PPU program to the SPU and vice versa:

- **vmx2spu.h**
  The macros and inline functions to map PPU Vector/SIMD intrinsics to generic SPU intrinsics
- **spu2vmx.h**
  The macros and inline functions to map generic SPU intrinsics to PPU Vector/SIMD intrinsics

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4 See note 2 on page 80.
5 See 3 on page 80.
vec_types.h

An SPU header file that defines a set of single token vector data types that are available on both the PPU and SPU. The SDK 3.0 provides both GCC and XLC versions of this header file.

To learn more about this issue, we recommend that you read the “SPU and PPU Vector Multimedia Extension Intrinsics” chapter and “Header files” chapter in the C/C++ Language Extensions for Cell Broadband Engine Architecture document.6

While the Vector/SIMD intrinsics contains various basic mathematical functions that are implemented by corresponding SIMD assembly instructions, more complex mathematical functions are not supported by these intrinsics. The SIMD Math Library is provided in the SDK and addresses this issue by providing a set of functions that extend the SIMD intrinsics and support additional common mathematical functions. Similar to SIMD intrinsics, the library operates on short 128-bit vectors from different types.

The SIMD Math Library is supported both by the SPU and PPU. The SPU version of this library is discussed in “SIMD Math Library” on page 262. The PPU version is similar, but the location of the library files is different:

- The simdmath.h file is in the /usr/spu/include directory.
- The Inline headers are in the /usr/spu/include/simdmath directory.
- The libsimdmath.a library is in the /usr/spu/lib directory.

4.1.2 Task parallelism and managing SPE threads

Programs that run on the Cell/B.E. processor typically partition the work among the eight available SPEs since each SPE is assigned with a different task and data to work on. In 3.3, “Deciding which parallel programming model to use” on page 53, we suggest several programming models to partition the work between the SPEs.

Regardless of the programming model, the main thread of the program is executed on the PPE, which creates sub-threads that run on the SPEs and off-load some function of the main program (to be run on the SPEs). It depends on the programming model how the threads and tasks are managed later, how the data is transferred, and how the different processors communicate.

Managing the code running on the SPEs on a Cell/B.E.-based system can be done by using the libspe library (SPE runtime management library) that is part of the SDK package. This library provides a standardized low-level application programming interface (API) that enables application access for the SPEs and runs some of the program threads on those SPEs.

6 See note 2 on page 80.
In general, applications that run on the Cell/B.E. processor do not have control over the physical SPE system resources because the operating system manages those resources. Instead, applications manage and use software constructs called *SPE contexts*. These SPE contexts are a logical representation of an SPE and hold all persistent information about a logical SPE. The libspe library operates on those contexts to manage the SPEs, but the programmer should not access those objects directly.

The operating system schedules SPE contexts from all running applications onto the physical SPE resources in the system for execution according to the scheduling priorities and policies that are associated with the runnable SPE contexts.

**Rescheduling an SPE and context switching:** Rescheduling an SPE and performing the context switching usually requires time because you must store most of the 256 KB of the local storage (LS) in memory and reload it with the code and data of the new thread. Therefore, ensure that the application does not allocate more SPE threads than the number of physical SPEs that are currently available (8 for a single Cell/B.E. processor and 16 for a QS20 or QS21 blade server).

The programmer must run the SPE contexts on a separate Linux thread, which enables the operating system to run them in parallel compared to the PPE threads and compared to other SPEs.

Refer to the *SPE Runtime Management Library* document, which contains a detailed description of the API for managing the SPE threads. The library also implements an API, which provides the means for communication and data transfer between PPE threads and SPEs. For more information, see 4.3, “Data transfer” on page 110, and 4.4, “Inter-processor communication” on page 178.

When creating an SPE thread, similar to Linux threads, the PPE program might pass up to three parameters to this function. The parameters may be either 64-bit parameters or 128-bit vectors. These parameters may be used later by the code that is running on the SPE. One common use, in the parameters, is to place an effective address of a control block that might be larger and contains additional information. The SPE can use this address to fetch this control block into its local storage memory.

---

There are two main methods to load SPE programs:

- **Static loading of SPE object**

  Statically compile the SPE object within the PPE program. At run time, the object is accessed as an external pointer that can be used by the programmer to load the program into local storage. The loading itself is implemented internally by the library API by using DMA.

- **Dynamic loading of SPE executable**

  Compile the SPE as stand-alone application. At run time, open the executable file, map it into the main memory, and then load it into the local storage of the SPE. This method is more flexible because you can decide, at run time, which program to load, for example, depending on the run time parameters. By using this method, you save linking the SPE program with the PPE program at the cost of lost encapsulation, so that the program is now a set of files, and not just a single executable.

We discuss the following topics next:

- In “Running a single SPE program” on page 85, we explain how to run code on a single SPE by using the static loading of an SPE object.

- In “Producing a multi-threaded program by using the SPEs” on page 89, we explain how to run code on multiple SPEs concurrently by using the dynamic loading of the SPE executable.

**Running a single SPE program**

In this section, we show how the user can run code on a single SPE. In our example, no Linux threads are used. Therefore, the PPE program blocks until the SPE stops executing, and the operating system returns from the system call that invoked the SPE execution.

Example 4-2 on page 86 through Example 4-4 on page 89 include the following actions for the PPU code, which are ordered according to how they are executed in the code:

1. Initiate a control structure to point to input and output data buffers and initiate the SPU executable’s parameter to point to this structure (step 1 in the code).

2. Create the SPE context by using `spe_context_create` function.

3. Statically load the SPE object into the SPE context local storage by using the `spe_program_load` function.

4. Run the SPE context by using `spe_context_run` function.

5. (Optional) Print the reason why the SPE stopped. In this example, obviously the end of its main function with return code 0 is the preferred one.

6. Destroy the SPE context by using the `spe_context_destroy` function.
For the SPU code, Example 4-2 on page 86 uses the parameters that the PPU code initiates in order to get the address control block and to get the control block from main storage to local storage.

Example 4-2 on page 86 shows the common header file. Be sure that the libspe2.h header file is included in order to run the SPE program.

**Source code:** The code in Example 4-2 through Example 4-4 on page 89 is included in the additional material for this book. See “Running a single SPE” on page 618 for more information.

---

**Example 4-2 Running a single SPE - Shared header file**

```c
// ==================================================================
// common.h file
// ==================================================================
#ifndef _COMMON_H_
#define _COMMON_H_

#define BUFF_SIZE 256

// the context that PPE forward to SPE
typedef struct{
    uint64_t ea_in; // effective address of input buffer
    uint64_t ea_out; // effective address of output buffer
} parm_context; // aligned to 16B

#endif // _COMMON_H_
```

Example 4-3 shows the PPU code.

**Example 4-3 Running a single SPE - PPU code**

```c
#include <libspe2.h>

#include "common.h"

spe_program_handle_t spu_main; // a pointer to SPE object
spe_context_ptr_t spe_ctx; // SPE context

// data structures to work with the SPE
#include <libspe2.h>  

volatile parm_context ctx __attribute__((aligned(16)));
volatile char in_data[BUFF_SIZE] __attribute__((aligned(128)));
```

---
volatile char out_data[BUFF_SIZE] __attribute__((aligned(128))); // function for printing the reason for SPE thread to stop // ==============================================================
void print_stop_reason( spe_stop_info_t *stop_info ){

    // result is a union that holds the SPE output result
    int result=stop_info->result.spe_exit_code;

    switch (stop_info->stop_reason) {
    case SPE_EXIT:
        printf("PPE: SPE stop_reason=SPE_EXIT, exit_code=");
        break;
    case SPE_STOP_AND_SIGNAL:
        printf("PPE: SPE stop_reason=SPE_STOP_AND_SIGNAL, signal_code=");
        break;
    case SPE_RUNTIME_ERROR:
        printf("PPE: SPE stop_reason=SPE_RUNTIME_ERROR, runtime_error=");
        break;
    case SPE_RUNTIME_EXCEPTION:
        printf("PPE: SPE stop_reason=SPE_RUNTIME_EXCEPTION, runtime_exception=");
        break;
    case SPE_RUNTIME_FATAL:
        printf("PPE: SPE stop_reason=SPE_RUNTIME_FATAL, runtime_fatal=");
        break;
    case SPE_CALLBACK_ERROR:
        printf("PPE: SPE stop_reason=SPE_CALLBACK_ERROR callback_error=");
        break;
    default:
        printf("PPE: SPE stop_reason=UNKNOWN, result=
");
        break;
    }
    printf("%d, status=%d\n",result,stop_info->spu_status);
}

// main
//===============================================================
int main( )
{
    spe_stop_info_t stop_info;
uint32_t entry = SPE_DEFAULT_ENTRY;

// STEP 1: initiate SPE control structure
cxt.ea_in = (uint64_t)in_data;
cxt.ea_out = (uint64_t)out_data;

// STEP 2: create SPE context
if ((spe_ctx = spe_context_create(0, NULL)) == NULL) {
    perror("Failed creating context"); exit(1);
}

// STEP 3: Load SPE object into SPE context local store
// (SPU’s executable file name is ‘spu_main’.
if (spe_program_load(spe_ctx, &spu_main)) {
    perror("Failed loading program"); exit(1);
}

// STEP 4: Run the SPE context (see ‘spu_pthread’ function above
// Note: this a synchronous call to the operating system
// which blocks until the SPE stops executing and the
// operating system returns from the system call that
// invoked the SPE execution.
if (spe_context_run(spe_ctx, &entry, 0, (void*)&cxt, NULL, &stop_info) < 0) {
    perror("Failed running context"); exit(1);
}

// STEP 5: Optionally print the SPE thread stop reason
print_stop_reason(&stop_info);

// STEP 6: destroy the SPE context
if (spe_context_destroy(spe_ctx)) {
    perror("Failed spe_context_destroy"); exit(1);
}
return (0);
Example 4-4 shows the SPU code.

**Example 4-4 Running a single SPE - SPU code**

```c
#include <spu_intrinsics.h>
#include <spu_mfcio.h>

#include "common.h"

static parm_context ctx __attribute__ ((aligned (128)));
volatile char in_data[BUFF_SIZE]  __attribute__ ((aligned(128)));
volatile char out_data[BUFF_SIZE] __attribute__ ((aligned(128)));

int main(int speid , uint64_t argp)
{
    uint32_t tag_id;

    //STEP 1: reserve tag IDs
    if((tag_id=mfc_tag_reserve())==MFC_TAG_INVALID){ // allocate tag
        printf("SPE: ERROR - can't reserve a tag ID\n"); return 1;
    }

    //STEP 2: get context information from system memory.
    mfc_get((void*) &ctx, argp, sizeof(ctx), tag_id, 0, 0);
    mfc_write_tag_mask(1<<tag_id);
    mfc_read_tag_status_all();

    //STEP 3: get input buffer, process it, and put results in output buffer

    //STEP 4: release tag IDs
    mfc_tag_release(tag_id); // release tag ID before exiting
    return 0;
}
```

**Producing a multi-threaded program by using the SPEs**

To obtain the best performance from an application running on the Cell/B.E. processor, use multiple SPEs concurrently. In this case, the application must create at least as many threads as concurrent SPE contexts are required. Each of these threads can run a single SPE context at a time. If \( n \) concurrent SPE contexts are needed, it is common to have a main application thread plus \( n \) threads dedicated to the SPE context execution.
In this section, we explain how the user can run code on multiple SPEs concurrently by using Linux threads. We use a specific scheme that is the most common one for Cell/B.E. programming. However, depending on the specific application, the programmer can use any other scheme.

In the code example in this section, we execute two SPE threads and include the following actions:

- Initiate the SPE control structures.
- Dynamically load the SPE executable into several SPEs:
  - Create SPE contexts.
  - Open images of SPE programs and map them into main storage.
  - Load SPE objects into SPE context local storage.
- Initiate the Linux threads and run the SPE executable concurrently on those threads. The PPU forwards parameters to the SPU programs.

In this example, the common header file is the same as Example 4-2 on page 86 in “Running a single SPE program” on page 85. You must include the libspe2.h header file to run the SPE programs and include the pthread.h file to use the Linux threads.

Source code: The code shown in Example 4-5 and Example 4-6 on page 93 is included in the additional material for this book. See “Running multiple SPEs concurrently” on page 618 for more information.

Example 4-5 shows the PPU code.

Example 4-5  Running multiple SPEs concurrently - PPU code

```c
// ppu_main.c file
#include <libspe2.h>
#include <cbe_mfc.h>
#include <pthread.h>

#include "common.h"

#define NUM_SPES 2

// input and output data buffers
volatile char in_data[BUFF_SIZE] __attribute__((aligned(128)));
volatile char out_data[BUFF_SIZE] __attribute__((aligned(128)));

// Data structures to work with the SPE
volatile parm_context ctx[NUM_SPES] __attribute__((aligned(16)));
```
spe_program_handle_t *program[BUFF_SIZE];

// data structure for running SPE thread ==============================
typedef struct spu_data {
   spe_context_ptr_t spe_ctx;
   pthread_t pthread;
   void *argp;
} spu_data_t;

spu_data_t data[NUM_SPES];

// create and run one SPE thread ==============================
void *spu_pthread(void *arg) {

   spu_data_t *datp = (spu_data_t *)arg;
   uint32_t entry = SPE_DEFAULT_ENTRY;

   if(spe_context_run(datp->spe_ctx,&entry,0,datp->argp,NULL,NULL)<0){
      perror ("Failed running context"); exit (1);
   };

   pthread_exit(NULL);
}

// main ===============================================================
int main( )
{
   int num;

   // names of the two SPU executable file names
   char spe_names[2][20] = {"spu1/spu_main1","spu2/spu_main2"};

   // STEP 1: initiate SPEs control structures
   for( num=0; num<NUM_SPES; num++){
      ctx[num].ea_in = (uint64_t)in_data  + num*(BUFF_SIZE/NUM_SPES);
      ctx[num].ea_out= (uint64_t)out_data + num*(BUFF_SIZE/NUM_SPES);
      data[num].argp = &ctx;
   }

   // Loop on all SPEs and for each perform two steps:
   // STEP 2: create SPE context
   // STEP 3: open images of SPE programs into main storage
   // 'spe_names' variable store the executable name
   // STEP 4: Load SPEs objects into SPE context local store
   for( num=0; num<NUM_SPES; num++){

if ((data[num].spe_ctx = spe_context_create(0, NULL)) == NULL) {
    perror("Failed creating context"); exit(1);
}
if (!(program[num] = spe_image_open(&spe_names[num][0]))) {
    perror("Fail opening image"); exit(1);
}
if (spe_program_load (data[num].spe_ctx, program[num])) {
    perror("Failed loading program"); exit(1);
}

// STEP 5: create SPE pthreads
for (num=0; num<NUM_SPES; num++){
    if(pthread_create(&data[num].pthread,NULL,&spu_pthread, &data[num])){
        perror("Failed creating thread"); exit(1);
    }
}

// Loop on all SPEs and for each perform two steps:
//   STEP 6: wait for all the SPE pthread to complete
//   STEP 7: destroy the SPE contexts
for (num=0; num<NUM_SPES; num++){
    if (pthread_join (data[num].pthread, NULL)) {
        perror("Failed joining thread"); exit (1);
    }
    if (spe_context_destroy(data[num].spe_ctx)) {
        perror("Failed spe_context_destroy"); exit(1);
    }
}
printf(")PPE:) Complete running all super-fast SPEs\n");
return (0);
Example 4-6 and Example 4-7 on page 94 show the SPU code.

**Example 4-6  Running multiple SPEs concurrently - SPU code version 1**

```c
// spu_main1.c file ==================================================
#include <spu_intrinsics.h>
#include <spu_mfcio.h>
#include "common.h"

static parm_context ctx __attribute__ ((aligned (128)));
volatile char in_data[BUFF_SIZE] __attribute__ ((aligned(128)));
volatile char out_data[BUFF_SIZE] __attribute__ ((aligned(128)));

int main(int speid , uint64_t argp)
{
    uint32_t tag_id;

    if((tag_id=mfc_tag_reserve())==MFC_TAG_INVALID){ // allocate tag
        printf("SPE: ERROR - can't reserve a tag ID\n"); return 1;
    }

    // get context information from system memory.
    mfc_get((void*) &ctx, argp, sizeof(ctx), tag_id, 0, 0);
    mfc_write_tag_mask(1<<tag_id);
    mfc_read_tag_status_all();

    printf("<SPE: Harel Rauch joyfully sleeps on the coach\n");

    // get input, process it using method A, and put results in output
    mfc_tag_release(tag_id); // release tag ID before exiting
    return 0;
}
```
Example 4-7   Running multiple SPEs concurrently - SPU code version 2

// spu_main2.c file ==================================================

// same variables and include as Example 4-6

int main(int speid , uint64_t argp)
{
    // same prefix as Example 4-4 on page 89

    printf("<SPE: Addie Dvir would like to fly here.\n"");
    // get input, process it using method A, and put results in output

    mfc_tag_release(tag_id); // release tag ID before exiting
    return 0;
}

4.1.3 Creating SPEs affinity by using a gang

The libspe library enables the programmer to create a gang, which is group of SPE contexts that should be executed together with certain properties. The mechanism enables you to create SPE-to-SPE affinity. This type of affinity means that certain SPE contexts can be created and placed next to another previously created SPE context. Affinity is always specified for pairs.

The SPE scheduler, which is responsible for mapping the SPE logical context to the physical SPE, honors this relationship by trying schedule the SPE contexts on physically adjacent SPUs. It depends on the current status of the system and whether it is able to do so. If the PPE program tries to create such affinity when no other code is running on the SPEs (in this program or other program), the schedule should succeed in doing so.
The usage of SPE-to-SPE affinity can create performance advantages in some cases. The performance gain is based mainly on the following characteristics of the Cell Broadband Engine Architecture (CBEA) and systems:

- On a Cell/B.E.-based symmetric multiprocessor (SMP) system, such as a BladeCenter QS21, communication between SPEs that are located on the same Cell/B.E. system are more efficient than data transfer between SPEs that are located on different Cell/B.E. chips. This includes both data transfer, for example LS to LS, and other types of communication such as mailbox and signals.

- Similarly to the previous characteristic, but on the same chip, communication between SPEs that are adjacent on the local element interconnect bus (EIB) is more efficient than between SPEs that are not adjacent.

Given these characteristics, in the case of massive SPE-to-SPE communication, you must physically locate specific SPEs next to each other.

Example 4-8 shows PPU code that creates such a chain of SPEs. This example is inspired by the SDK code example named `dmabench` that is in the `/opt/cell/sdk/src/benchmarks/dma` directory.

### Chain of SPEs:

This example demonstrates only how to create a chain of SPEs that are physically located one next to the other. The SPE pipeline, which is based on this structure, for example where each SPE executes DMA transfers from the local storage of the previous SPE on the chain, does not provide the optimal results since only half of the EIB rings are used. Therefore, half of the bandwidth is lost. When the physical location of the SPEs is known by using the affinity methods, the programmer can use this information to locate the SPEs elsewhere on the SPE pipeline.

The article “Cell Broadband Engine Architecture and its first implementation: A performance view” [18 on page 624] provides information about the bandwidth that was measured for some SPE-to-SPE DMA transfers. This information might be useful when deciding how to locate the SPEs related to each other on a given algorithm.

**Example 4-8  PPU code for creating SPE physical chain using affinity**

```
// take include files, ‘spu_data_t’ structure and the ‘spu_pthread’
// function from Example 4-5 on page 90

spe_gang_context_ptr_t gang;
spu_context_ptr_t ctx[NUM_SPEES];

int main( )
```
{
    int i;

    gang = NULL;

    // create a gang
    if ((gang = spe_gang_context_create(0))==NULL) {
        perror("Failed spe_gang_context_create"); exit(1);
    }

    // create SPE contexts as part of the gang which preserve affinity
    // between each SPE pair.
    // SPEs' affinity is based on a chain architecture such as SPE[i]
    // and SPE[i+1] are physically adjacent.
    for (i=0; i<NUM_SPES; i++) {
        ctx[i]=spe_context_create_affinity(0,(i==0)?NULL:ctx[i-1],gang));
        if(ctx[i]==NULL){
            perror("Failed spe_context_create_affinity"); exit(1);
        }
    }

    // ... Omitted section:
    // creates SPE contexts, load the program to the local stores,
    // run the SPE threads, and waits for SPE threads to complete.

    // (the entire source code for this example is comes with the book's
    // additional material).

    // See also 4.1.2, “Task parallelism and managing SPE threads” on
    page 83
}

4.2 Storage domains, channels, and MMIO interfaces

In this section, we describe the main storage domains of the CBEA. The CBEA has a unique memory architecture. Your understanding of those domains is a key issue in order to know how to program the Cell/B.E. application and how the data can be partitioned and transferred in such an application. We discuss the storage domain in 4.2.1, “Storage domains” on page 97.

The MFC is a hardware component that implements most of the Cell/B.E. inter-processor communication mechanism. It includes the most significant means to initiate data transfer, including DMA data transfers. While they are in each of the SPEs, the MFC interfaces can be accessed by both a program running on an SPU or a program running on the PPU. We discuss the MFC in the following sections:

- In 4.2.2, “MFC channels and MMIO interfaces and queues” on page 99, we discuss the main features of the MFC and the two main interfaces, the channel interface and memory-mapped I/O (MMIO) interface, that it has with the programs.

- In 4.2.3, “SPU programming methods to access the MFC channel interface” on page 101, we discuss the programming methods for accessing the MFC interfaces and initiating its mechanisms from an SPU program.

- In 4.2.4, “PPU programming methods to access the MFC MMIO interface” on page 105, we discuss the programming methods for accessing the MFC interfaces and initiating its mechanisms from a PPU program.

In addition to discussing the MFC interfaces and programming methods to program it, we explain how to use the MFC mechanisms in the following sections:

- DMA data transfers and synchronization of data transfers in 4.3, “Data transfer” on page 110

- Communication mechanisms, such as mailbox, signals and events, between the different processors (PPE and SPEs) in 4.4, “Inter-processor communication” on page 178

4.2.1 Storage domains

The CBEA defines three types of storage domains that are defined in the Cell/B.E. chip: one main-storage domain, eight SPE LS domains, and eight SPE channel domains. Figure 4-1 on page 98 illustrates the storage domains and interfaces in the CBEA.
The main-storage domain, which is the entire effective address space, can be configured by the PPE operating system to be shared by all processors in the system. Alternatively, the local-storage and channel problem-state (user-state) domains are private to the SPE components. The main components in each SPE are the SPU, the LS and the MFC, which handles the DMA data transfer.

**Main storage:** In this document, we use the term *main storage* to describe any component that has an effective address mapping on the main storage domain.
An SPE program references its own LS by using a *local store address* (*LSA*). The LS of each SPE is also assigned a *real address* (*RA*) range within the system’s memory map. As a result, privileged software on the PPE can map LS areas into the *effective address* (*EA*) space, where the PPE, other SPEs, and other devices that generate EAs can access the LS like any regular component on the main storage.

Code that runs on an SPU can only fetch instructions from its own LS, and loads and stores can only access that LS.

Data transfers between the LS and main storage of the SPE are primarily executed by using DMA transfers that are controlled by the MFC DMA controller for that SPE. The MFC of each SPE serves as a data-transfer engine. DMA transfer requests contain both an LSA and an EA. Therefore, they can address both the LS and main storage of an SPE and thereby initiate DMA transfers between the domains. The MFC accomplishes this by maintaining and processing an MFC command queue.

Because the local storage can be mapped to the main storage, SPEs can use DMA operations to directly transfer data between their LS to the LS of another SPE. This mode of data transfer is efficient, because the DMA transfers go directly from SPE to SPE on the high performance local bus without involving the system memory.

### 4.2.2 MFC channels and MMIO interfaces and queues

Each MFC has two main interfaces through which MFC commands can be initiated:

- **Channel interface**

  The SPU can use this interface to interact with the associated MFC by executing a series of writes or reads to the various channels, which in response enqueue MFC commands. Since accessing the channel remains local within a certain SPE, it has low latency (for nonblocking commands, about six cycles if the channel is not full). It also does not have any negative influence EIB bandwidth.

- **MMIO interface**

  PPE or other SPUs can use this interface to interact with any MFC by accessing the Command-Parameter Registers of the MFC. The registers can be mapped to the system’s real-address space so that the PPE or SPUs can access them by executing MMIO reads and writes to the corresponding effective address.
For a detailed description of the channels and MMIO interfaces, see the “SPE Channel and related MMIO interface” chapter in the *Cell Broadband Engine Programming Handbook.*

When accessing the two interfaces, commands are inserted into one of the two MFC-independent command queues:

- The channel interface is associated with the MFC SPU command queue.
- The MMIO interface is associated with the MFC Proxy command queue.

In regard to the channel interface, each channel can be defined as either *blocking* or *nonblocking*. When the SPE reads or writes a nonblocking channel, the operation executes without delay. However, when the SPE software reads or writes a blocking channel, the SPE might stall for an arbitrary length of time if the associated channel count (which is its remaining capacity) is 0. In this case, the SPE remains stalled until the channel count becomes 1 or more.

The stalling mechanism reduces SPE software complexity and allows an SPE to minimize the power consumed by message-based synchronization. To avoid stalling on access to a blocking channel, SPE software can read the channel count to determine the available channel capacity. In addition, many of the channels have a corresponding and independent event that can be enabled to cause an asynchronous interrupt.

Alternatively, accessing the MMIO interface is always nonblocking. If a PPE (or other SPE) writes a command while the queue is full, then the last entry in the queue is overridden with no indication to the software. Therefore, the PPE (or other SPE) should first verify if there is available space in the queue by reading the queue status register. Then only if it is not full, then the PPE should write a command to it. Be aware that waiting for available space by continuously reading this register in a loop has a negative effect on the performance of the entire chip because it involves transactions on the local EIB.

Similarly, reading from an MMIO register when a queue is empty returns invalid data. Therefore, the PPE (or other SPE) should first read the corresponding status register. Only if there is a valid entry (queue is not empty), the MMIO register itself should be read.

---

8 See note 1 on page 78.
Table 4-1 summarizes the main attributes of the two main interfaces of the MFC.

<table>
<thead>
<tr>
<th>Interface</th>
<th>Queue</th>
<th>Initiator</th>
<th>Blocking</th>
<th>Full</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Channels</td>
<td>MFC SPU command queue</td>
<td>Local SPU</td>
<td>Blocking or nonblocking</td>
<td>Wait until the queue has an available entry</td>
<td>For MFC commands sent from the SPU through the channel interface</td>
</tr>
<tr>
<td>MMIO</td>
<td>MFC proxy command queue</td>
<td>PPE or other SPEs</td>
<td>Always non blocking</td>
<td>Overwrite the last entry</td>
<td>For MFC commands sent from the PPE, other SPUs, or other devices through the MMIO registers</td>
</tr>
</tbody>
</table>

4.2.3 SPU programming methods to access the MFC channel interface

Software running on an SPU can access the MFC facilities through the channel interface. In this section, we discuss the four programming methods to access this interface. We list them as follows from the most abstract to the lowest level:

1. MFC functions
2. Composite intrinsics
3. Low-level intrinsics
4. Assembly-language instructions

**MFC functions:** The simplest method, from a *programming point of view*, is to access the DMA mechanism through the MFC functions. Therefore, most of the examples in this chapter, besides the examples in this section, are written by using MFC functions. However, from a *performance point of view*, use of the MFC functions does not always provide the best results, especially when invoked from a PPE program.

Many code examples in the SDK package also use this method. However, the examples in the SDK documentation rely mostly on composite intrinsics and low-level intrinsics. Such examples are available in the *Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial* document.a

a. See note 3 on page 80.

In this section, we illustrate the differences between the four methods by using the DMA *get* command, which moves data from a component in main storage to local storage. This is done only for demonstration purposes. Similar implementation can be performed for using each of the other MFC facilities, such as mailboxes, signals, and events. If you are not familiar with the MFC DMA
commands, refer to 4.3, “Data transfer” on page 110, before continuing with this section.

Some parameters, which are listed in Table 4-2, are common to all DMA transfer commands. For all alternatives, we assume that the DMA transfer parameters, described in Table 4-2 are defined previously to executing the DMA command.

Table 4-2  DMA transfer parameters

<table>
<thead>
<tr>
<th>Name</th>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isa</td>
<td>void*</td>
<td>local-storage address</td>
</tr>
<tr>
<td>ea</td>
<td>uint64_t or eah</td>
<td>Effective address in main storage&lt;sup&gt;a&lt;/sup&gt; or Effective address higher bits in main storage&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>eah</td>
<td>uint32_t</td>
<td>Effectve address higher bits in main storage&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>eal</td>
<td>uint32_t</td>
<td>Effective address lower bits in main storage&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>size</td>
<td>uint32_t</td>
<td>DMA transfer size in bytes</td>
</tr>
<tr>
<td>tag</td>
<td>uint32_t</td>
<td>DMA group tag ID</td>
</tr>
<tr>
<td>tid</td>
<td>uint32_t</td>
<td>Transfer class identifier&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>rid</td>
<td>uint32_t</td>
<td>Replacement&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> Used for MFC functions only.
<sup>b</sup> Used for methods other than MFC functions.

In the following sections, we describe the four methods to access the MFC facilities.

**MFC functions**

MFC functions are a set of convenience functions. Each perform a single DMA command (for example get, put, barrier). The functions are implemented either as macros or as built-in functions within the compiler, causing the compiler to map each of those functions to a certain composite intrinsic (similar to those discussed in “Composite intrinsics” on page 103) with the corresponding operands.

Table 4-3 on page 113 provides a list and descriptions of all the available MFC functions. For a more detailed description, see the “Programming support for MFC input and output” chapter in the C/C++ Language Extensions for Cell Broadband Engine Architecture document.<sup>9</sup>

To use the intrinsics, the programmer must include the spu_mfcio.h header file. Example 4-9 on page 103 shows the initiation of a single get command by using the MFC functions.

<sup>9</sup> See note 2 on page 80.
Example 4-9  SPU MFC function get command example

```c
#include "spu_mfcio.h"

mfc_get(lsa, ea, size, tag, tid, rid);

// Implemented as the following composite intrinsic:
// spu_mfcdma64(lsa, mfc_ea2h(ea), mfc_ea2l(ea), size, tag,
// ((tid<<24)|(rid<<16)|MFC_GET_CMD));

// wait until DMA transfer is complete (or do other things before that)
```

**Composite intrinsics**

The SDK 3.0 defines a small number of composite intrinsics to handle DMA commands. Each composite intrinsic handles one DMA command and is constructed from a series of low-level intrinsics (similar to those discussed in “Low-level intrinsics” on page 104). These intrinsics are further described in the “Composite intrinsics” chapter in the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document\(^{10}\) and in the *Cell Broadband Engine Architecture* document\(^{11}\).

To use the intrinsics, the programmer must include the `spu_intrinsics.h` header file. In addition, the `spu_mfcio.h` header file includes useful predefined values of the DMA commands (for example, MFC_GET_CMD in Example 4-10). The programmer can include this file and use its predefined values instead of explicitly writing the corresponding values. Example 4-10 shows the initiation of a single get command by using composite intrinsics.

Example 4-10  SPU composite intrinsics get command example

```c
#include <spu_intrinsics.h>
#include "spu_mfcio.h"

spu_mfcdma64(lsa, eah, eal, size, tag, MFC_GET_CMD);

// Implemented using the six low level intrinsics in Example 4-11 on page 104

// MFC_GET_CMD is defined as 0x0040 in spu_mfcio.h
```

---

\(^{10}\) See note 2 on page 80.

\(^{11}\) The *Cell Broadband Engine Architecture* document is on the Web at the following address: http://www-01.ibm.com/chips/techlib/techlib.nsf/techdocs/1AEEE1270EA2776387257060006E61BA/$file/CBEA_v1.02_11oct2007_pub.pdf
Low-level intrinsics
A series of a few low-level intrinsics (meaning generic or specific intrinsics) should be executed in order to run a single DMA transfer. Each intrinsic is mapped to a single assembly instruction.

The relevant low-level intrinsic are described in the “Channel control intrinsics” chapter in the C/C++ Language Extensions for Cell Broadband Engine Architecture document.12

To use the intrinsics, the programmer must include the spu_intrinsics.h header file. Example 4-11 shows the initiation of a single get command by using low-level intrinsics.

Example 4-11  SPU low-level intrinsics get command example

```
spu_writech(MFC_LSA, lsa);
spu_writech(MFC_EAH, eah);
spu_writech(MFC_EAL, eal);
spu_writech(MFC_Size, size);
spu_writech(MFC_TagID, tag);
spu_writech(MFC_CMD, 0x0040);
```

Assembly-language instructions
Assembly-language instructions are similar to low-level intrinsics. The intrinsics are a series of ABI-compliant assembly language instructions that are executed for a single DMA transfer. Each of the low-level intrinsics represents one assembly instruction. From practical point of view, the only case where we can recommend using this method instead the low-level intrinsics is when the program is written in assembly.

Example 4-12 illustrates the initiation of a single get command by using assembly-language instructions.

Example 4-12  SPU assembly-language instructions get command example

```
.text
.globl dma_transfer
dma_transfer:

wrch$MFC_LSA, $3
wrch$MFC_EAH, $4
wrch $MFC_EAL, $5
wrch $MFC_Size, $6
```

12 See note 2 on page 80.
4.2.4 PPU programming methods to access the MFC MMIO interface

Software running on a PPU may access the MFC facilities through the MMIO interface. There are two main methods to access this interface as discussed in this section. We list these methods from the most abstract to the lowest level:

1. MFC functions
2. Direct problem state access (or direct SPE access)

Unlike the SPU case when using the channel interface, in the PPU case, it is not always recommended to use the MFC functions. The following list summarizes the differences between the two methods and recommendations for using them:

- MFC functions are simpler from a programmer's point of view. Therefore, usage of this method can reduce development time and make the code more readable.
- Direct problem state access gives the programmer more flexibility, especially when non-standard mechanism must be implemented.
- Direct problem state access has significantly better performance in many cases, such as when writing to the inbound mailbox. Two reasons for the reduction in performance for the MFC functions is the call overhead and the mutex locking associated with the library functions being thread safe. Therefore, in cases where the performance, for example latency, of the PPE access to the MFC is important, use the direct SPE access, which may have significantly better performance over the MFC functions.

Most of the examples in this document, as well as many code examples in the SDK package, use the MFC functions method. However, the examples in the SDK documentation rely mostly on the direct SPE access method. Many such examples are available in the Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial.\(^\text{13}\)

In this section, we illustrate the differences between the two methods by using the DMA **get** command to move data from a component on the main storage to the local storage. Similar implementation may be performed for using each of the other MFC facilities, including mailboxes, signals, events, and so on. We used the same parameters that are defined in Table 4-2 on page 102, but the additional `spe_context_ptr_t spe_ctx` parameter is added in the PPU case. This

---

\(^{13}\) See note 3 on page 80.
parameter provides a pointer to the context of the relevant SPE. This context is created when the SPE thread is created.

In the following sections, we describe the two main methods for a PPE to access the MFC facilities.

**MFC functions**

MFC functions are a set of convenience functions. Each set implements a single DMA command, for example a **get**, **put**, or **barrier**). Table 4-3 on page 113 provides a list and description of all the available functions. For a more detailed description, see the “SPE MFC problem state facilities” chapter in the *SPE Runtime Management Library* document.\(^{14}\)

Unlike the SPE implementation, the implementation of the MFC functions for the PPE usually involves accessing the operating system kernel, which adds a non-negligible number of cycles and increases the latency of those functions.

To use the intrinsics, the programmer must include the *libspe2.h* header file. Example 4-13 illustrates the initiation of a single **get** command by using MFC functions.

*Example 4-13  PPU MFC functions get example*

```c
#include “libspe2.h”

spe_mfcio_get ( spe_ctx, lsa, ea, size, tag, tid, rid);

// wait till data was transferred to LS, or do other things...
```

**Direct problem state access**

The second option for PPE software to access the MFC facilities is to explicitly interact with the relevant MMIO interface of the relevant SPE. To do so, the programmer should use the software to perform the following actions:

1. Map the corresponding problem state area of the relevant SPE to the PPE thread address space. The programmer can do this by using the `spe_ps_area_get` function in the *libspe* library. (Include the *libspe2.h* file to use this function.)

2. After the corresponding problem state area is mapped, the programmer can access the area by using one of the following methods:
   - Use one of the inline functions for direct problem state access that are defined in the *cbe_mfc.h* header file. This header file makes using the

\(^{14}\) See note 7 on page 84.
direct problem state as easy as using the libspe functions. For example, the function _spe_sig_notify_1_read reads the SPU_Sig.Notify_1 register, the _spe_out_mbox_read function reads a value from the SPU_Out_Mbox mailbox register, and the _spe_mfc_dma function enqueues a DMA request.

- Use direct memory load or store instruction to access the relevant MMIO registers. The easiest way to do so is by using enums and structs that describe the problem state areas and the offset of the MMIO registers. The enums and structs are defined in the libspe2_types.h and cbea_map.h header files (see Example 4-15 on page 109). However, to use them, the programmer should include only the libspe2.h file.

After the problem state area is mapped, direct access to this area by the application does not involve the kernel, and therefore, has a smaller latency than the corresponding MFC function.

**SPE_MAP_PS flag:** The PPE programmer must set the SPE_MAP_PS flag when creating the SPE context (in the spe_context_create function) of the SPE whose problem state area the programmer later will try to map by using the spe_ps_area_get function. See Example 4-14.

Example 4-14 shows the PPU code for mapping an SPE problem state to the thread address space and initiating a single get command by using direct SPE access.

**Source code:** The code of Example 4-14 is included in the additional material for this book. See “Simple PPU vector/SIMD code” on page 617 for more information.

**Example 4-14  PPU direct SPE access get example**

```c
#include <libspe2.h>
#include <cbe_mfc.h>
#include <pthread.h>

spe_context_ptr_t spe_ctx;
uint32_tlsa, eah, eal, tag, size, ret, status;
volatile spe_mfc_command_area_t* mfc_cmd;
volatile char data[BUFF_SIZE] __attribute__((aligned (128)));

// create SPE context: must set SPE_MAP_PS flag to access problem state
spe_ctx = spe_context_create (SPE_MAP_PS, NULL);
```
// - open an SPE executable and map using ‘spe_image_open’ function
// - load SPU program into LS using ‘spe_program_load’ function
// - create SPE pthread using ‘pthread_create’ function

// map SPE problem state using spe_ps_area_get
if ((mfc_cmd = spe_ps_area_get(data.spe_ctx, SPE_MFC_COMMAND_AREA)) == NULL) {
    perror("Failed mapping MFC command area"); exit (1);
}

// lsa = LS space address that SPU code provide
// eal = ((uintptr_t)&data) & 0xffffffff;
// eah = ((uint64_t)(uintptr_t)&data)>>32;
// tag = number from 0 to 15 (as 16-31 are used by the kernel)
// size= .....  
while( (mfc_cmd->MFC_QStatus & 0x0000FFFF) == 0);

do{
    mfc_cmd->MFC_LSA = lsa;
    mfc_cmd->MFC_EAH = eah;
    mfc_cmd->MFC_EAL = eal;
    mfc_cmd->MFC_Size_Tag = (size<<16) | tag;
    mfc_cmd->MFC_ClassID_CMD = MFC_PUT_CMD;
    ret = mfc_cmd->MFC_CMDStatus;
} while(ret&0x3); //enqueuing until success

//following 2 lines are commented in order to be similar to Example 4-13 on page 106
//ret=spe_mfcio_tag_status_read(spe_ctx, 1<<tag, SPE_TAG_ALL, &status);
//if( ret !=0) printf("error in GET command");

The SDK 3.0 header files libspe2_types.h and cbea_map.h contain several enums and structs that define the problem state areas and registers. Therefore, the programming is made more convenient when accessing the MMIO interface from the PPE. Example 4-15 on page 109 shows the enums and structs.
Example 4-15  Enums and structs for defining problem state areas and registers

// From libspe2_types.h header file
// ==============================================================
enum ps_area { SPE_MSSYNC_AREA, SPE_MFC_COMMAND_AREA, SPE_CONTROL_AREA,
SPE_SIG_NOTIFY_1_AREA, SPE_SIG_NOTIFY_2_AREA };  

// From cbea_map.h header file
// ==============================================================
SPE_MSSYNC_AREA: MFC multisource synchronization register area
typedef struct spe_mssync_area {
    unsigned int MFC_MSSync;
} spe_mssync_area_t;

// SPE_MFC_COMMAND_AREA: MFC command parameter queue control area
typedef struct spe_mfc_command_area {
    unsigned char reserved_0_3[4];
    unsigned int MFC_LSA;
    unsigned int MFC_EAH;
    unsigned int MFC_EAL;
    unsigned int MFC_Size_Tag;
    union {
        unsigned int MFC_ClassID_CMD;
        unsigned int MFC_CMDStatus;
    };
    unsigned char reserved_18_103[236];
    unsigned int MFC_QStatus;
    unsigned char reserved_108_203[252];
    unsigned int Prxy_QueryType;
    unsigned char reserved_208_21B[20];
    unsigned int Prxy_QueryMask;
    unsigned char reserved_220_22B[12];
    unsigned int Prxy_TagStatus;
} spe_mfc_command_area_t;

// SPE_CONTROL_AREA: SPU control area
typedef struct spe_spu_control_area {
    unsigned char reserved_0_3[4];
    unsigned int SPU_Out_Mbox;
    unsigned char reserved_8_B[4];
    unsigned int SPU_In_Mbox;
    unsigned char reserved_10_13[4];
    unsigned int SPU_Mbox_Stat;
    unsigned char reserved_18_1B[4];
    unsigned int SPU_RunCntl;
4.3 Data transfer

The Cell/B.E. processor has a radical organization of storage and asynchronous DMA transfers between LS and main storage. While this architecture enables high performance, it requires the application programmer to explicitly handle the data transfers between LS and main memory or other local storage. Programming efficient data transfers is a key issue for preventing errors, such as synchronization errors, which are difficult to debug, and for having optimized out of a program running on a Cell/B.E.-based system.

Programming the DMA data transfer can be done by using either an SPU program with the channel interface or by using the a PPU program with the MMIO interface. Refer to 4.2, “Storage domains, channels, and MMIO interfaces” on page 97, about the usage of these interfaces.

Regarding the issue of DMA commands to the MFC command, the channel interface has 16 entries in its corresponding MFC SPU command queue, which stands for up to 16 DMA commands that can be handled simultaneously by the MFC. The corresponding MMIO interface has only eight entries in its corresponding MFC proxy command queue. For this reason and for other reasons, such as smaller latency in issuing the DMA commands, less overhead on the internal EIB, and so on, the programmer should run DMA commands from the SPU program rather than from the PPU.
In this section, we explain the DMA data transfer methods as well as other data transfer methods, such as direct load and store, that may be used to transfer data between LS and main memory or between one LS to another LS.

In this section, we present the following topics:

- In 4.3.1, “DMA commands” on page 112, we provide an overview over the DMA commands that are supported by the MFC, whether they are initiated by the SPE of the PPE.

In the next three sections, we explain how to initiate various data transfers by using the SDK 3.0 core libraries:

- In 4.3.2, “SPE-initiated DMA transfer between the LS and main storage” on page 120, we discuss how a program running on an SPU can initiate DMA commands between its LS and the main memory by using the associated MFC.

- In 4.3.3, “PPU initiated DMA transfer between LS and main storage” on page 138, we explain how a program running on a PPU can initiate DMA commands between the LS of some SPEs and main memory by using the MFC that is associated with this SPE.

- In 4.3.4, “Direct problem state access and LS-to-LS transfer” on page 144, we discuss two different issues. We explain how an LS of some SPEs can be accessed directly by the PPU or by an SPU program running on another SPE.

In the next two sections, we discuss two alternatives (other the core libraries) that come with the SDK 3.0 and can be used for simpler initiation of data transfer between the LS and main storage:

- In 4.3.5, “Facilitating random data access by using the SPU software cache” on page 148, we explain how to use the SPU software managed cache and in which cases to use it.

- In 4.3.6, “Automatic software caching on SPE” on page 157, we discuss an automated version of the SPU software cache that provides an even simpler programming method but with the potential for reduced performance.

In the next three sections, we describe several fundamental techniques for programming performance of efficient data transfers:

- In 4.3.7, “Efficient data transfers by overlapping DMA and computation” on page 159, we discuss the double buffering and multibuffering techniques that enable overlap between DMA transfers and computation. The overlapping often provides a significant performance improvement.

- In 4.3.8, “Improving the page hit ratio by using huge pages” on page 166, we explain how to configure large pages in the system and when it can be useful to do this.
In 4.3.9, “Improving memory access using NUMA” on page 171, we explain how to use the nonuniform memory access (NUMA) features on a Cell/B.E.-based system.

Another topic that is relevant to data transfer and that is not covered in the following sections is the ordering between different data turnovers and synchronization techniques. Refer to 4.5, “Shared storage synchronizing and data ordering” on page 218, to learn more about this topic.

### 4.3.1 DMA commands

MFC supports a set of DMA commands that provide the main mechanism that enables data transfer between the LS and main storage. It also supports a set of synchronization commands that are used to control the order in which storage accesses are performed and maintain synchronization with other processors and devices in the system.

Each MFC has an associated memory management unit (MMU) that holds and processes address-translation and access-permission information that is supplied by the PPE operating system. While this MMU is distinct from the one used by the PPE, to process an effective address provided by a DMA command, the MMU uses the same method as the PPE memory-management functions. Thus, DMA transfers are coherent with respect to system storage. Attributes of system storage are governed by the page and segment tables of the PowerPC Architecture.

In the following sections, we discuss several issues related to the supported DMA commands.

MFC supports a set of DMA commands. DMA commands can initiate or monitor the status of data transfers.

- Each MFC can maintain and process up to 16 in-progress DMA command requests and DMA transfers, which are executed asynchronously to the code execution.

- The MFC can autonomously manage a sequence of DMA transfers in response to a DMA list command from its associated SPU. DMA lists are a sequence of eight-byte list elements, stored in the LS of an SPE, each of which describes a single DMA transfer.

- Each DMA command is tagged with a 5-bit Tag ID, which defines up to 32 IDs. The software can use this identifier to check or wait on the completion of all queued commands in one or more tag groups.

Refer to “Supported and recommended values for DMA parameters” on page 116 for the supported and recommended values for the DMA parameters.
Refer to “Supported and recommended values for DMA-list parameters” on page 117 for the supported and recommended parameters of a DMA list.

Table 4-3 summarizes all the DMA commands that are supported by the MFC. For each command, we mention the SPU and the PPE MFC functions that implement it, if any. (A blank cell indicates that this command is not supported by either the SPE or PPE.) For detailed information about the MFC commands, see the “DMA transfers and inter-processor communication” chapter in the Cell Broadband Engine Programming Handbook.\(^\text{15}\)

The SPU functions are defined in the `spu_mfcio.h` header file and are described in the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document.\(^\text{16}\) The PPE functions are defined in the `libspe2.h` header file and are described in the *SPE Runtime Management Library* document.\(^\text{17}\)

SDK 3.0 defines another set of PPE inline functions for handling the DMA data transfer in the `cbe_mfc.h` file, which is preferred from a performance point of view over the `libspe2.h` functions. While the `cbe_mfc.h` functions are not well described in the official SDK documents, they are straight forward and easy to use. To enqueue a DMA command, the programmer can issue the `_spe_mfc_dma` function with the `cmd` parameter indicating that the DMA command should be enqueued. For example, set the `cmd` parameter to MFC_PUT_CMD for the put command or set it to MFC_GETS_CMD for the gets command, and so on.

<table>
<thead>
<tr>
<th>Command</th>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><strong>SPU</strong></td>
<td><strong>PPE</strong></td>
</tr>
<tr>
<td>Put commands</td>
<td></td>
<td></td>
</tr>
<tr>
<td>put</td>
<td>mfc_put</td>
<td>spe_mfcio_put</td>
</tr>
<tr>
<td>puts</td>
<td>Unsupported</td>
<td>None(^\text{a})</td>
</tr>
<tr>
<td>putf</td>
<td>mfc_putchar</td>
<td>spe_mfcio_putchar</td>
</tr>
</tbody>
</table>

\(^{15}\) See note 1 on page 78.  
\(^{16}\) See note 2 on page 80.  
\(^{17}\) See note 7 on page 84.
<table>
<thead>
<tr>
<th>Command</th>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>putb</td>
<td>mfc_putb</td>
<td>Moves data from the LS to the effective address with the barrier option. This command and all subsequent commands with the same tag ID as this command are locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
<tr>
<td>putfs</td>
<td>Unsupported</td>
<td>Moves data from the LS to the effective address with the fence option. This command is locally ordered with respect to all previously issued commands within the same tag group and command queue. Starts the SPU after the DMA operation completes.</td>
</tr>
<tr>
<td>putbs</td>
<td>Unsupported</td>
<td>Moves data from the LS to the effective address with the barrier option. This command and all subsequent commands with the same tag ID as this command are locally ordered with respect to all previously issued commands within the same tag group and command queue. Starts the SPU after the DMA operation completes.</td>
</tr>
<tr>
<td>putl</td>
<td>mfc_putl</td>
<td>Moves data from the LS to the effective address by using an MFC list.</td>
</tr>
<tr>
<td>putlf</td>
<td>mfc_putlf</td>
<td>Moves data from the LS to the effective address by using an MFC list with the fence option. This command is locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
<tr>
<td>putlb</td>
<td>mfc_putlb</td>
<td>Moves data from the LS to the effective address by using an MFC list with the barrier option. This command and all subsequent commands with the same tag ID as this command are locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
</tbody>
</table>

**get commands**

<table>
<thead>
<tr>
<th>Command</th>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>get</td>
<td>mfc_get</td>
<td>Moves data from the effective address to the LS.</td>
</tr>
<tr>
<td>gets</td>
<td>Unsupported</td>
<td>Moves data from the effective address to the LS and starts the SPU after the DMA operation completes.</td>
</tr>
<tr>
<td>Command</td>
<td>Function</td>
<td>Description</td>
</tr>
<tr>
<td>---------</td>
<td>-----------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>getf</strong></td>
<td>mfc_getf</td>
<td>Moves data from the effective address to the LS with the fence option. This command is locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
<tr>
<td></td>
<td>spe_mfcio_getf</td>
<td></td>
</tr>
<tr>
<td><strong>getb</strong></td>
<td>mfc_getb</td>
<td>Moves data from the effective address to the LS with the barrier option. This command and all subsequent commands with the same tag ID as this command are locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
<tr>
<td></td>
<td>spe_mfcio_getb</td>
<td></td>
</tr>
<tr>
<td><strong>getfs</strong></td>
<td>Unsupported</td>
<td>Moves data from the effective address to the LS with the fence option. This command is locally ordered with respect to all previously issued commands within the same tag group. Starts the SPU after the DMA operation completes.</td>
</tr>
<tr>
<td></td>
<td>None&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td><strong>getbs</strong></td>
<td>Unsupported</td>
<td>Moves data from the effective address to LS with the barrier option. This command and all subsequent commands with the same tag ID as this command are locally ordered with respect to all previously issued commands within the same tag group and command queue. Starts the SPU after the DMA operation completes.</td>
</tr>
<tr>
<td></td>
<td>None&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td><strong>getl</strong></td>
<td>mfc_getl</td>
<td>Moves data from the effective address to the LS by using an MFC list.</td>
</tr>
<tr>
<td></td>
<td>Unsupported</td>
<td></td>
</tr>
<tr>
<td><strong>getlf</strong></td>
<td>mfc_getlf</td>
<td>Moves data from the effective address to the LS by using an MFC list with the fence option. This command is locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
<tr>
<td></td>
<td>Unsupported</td>
<td></td>
</tr>
<tr>
<td><strong>getlb</strong></td>
<td>mfc_getb</td>
<td>Moves data from the effective address to the LS by using an MFC list with the barrier option. This command and all subsequent commands with the same tag ID as this command are locally ordered with respect to all previously issued commands within the same tag group and command queue.</td>
</tr>
<tr>
<td></td>
<td>Unsupported</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> While this command can be issued by the PPE, no MFC function supports it.
The suffixes in Table 4-4 are associated with the DMA commands and extend or refine the function of a command. For example, a *putb command moves data from the LS to the effective address similar to the *put command, but also adds a barrier.

**Table 4-4 MFC commands suffixes**

<table>
<thead>
<tr>
<th>Mnemonic</th>
<th>Possible initiator</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>s</td>
<td>F o r SPU Yes</td>
<td>Starts the SPU. Starts the SPU running at the address in the SPU Next Program Counter Register (SPU_NPC) after the MFC command completes.</td>
</tr>
<tr>
<td>f</td>
<td>F o r SPU Yes</td>
<td>Tag-specific fence. The command is locally ordered with respect to all previously issued commands in the same tag group and command queue.</td>
</tr>
<tr>
<td>b</td>
<td>F o r SPU Yes</td>
<td>Tag-specific barrier. The command is locally ordered with respect to all previously issued and all subsequently issued commands in the same tag group and command queue.</td>
</tr>
<tr>
<td>l</td>
<td>F o r SPU Yes</td>
<td>List command. The command processes a list of DMA list elements that are in the LS. There are up to 2048 elements in a list. Each list element specifies a transfer of up to 16 KB.</td>
</tr>
</tbody>
</table>

**Supported and recommended values for DMA parameters**

In the following list, we summarize the supported or recommended values of the MFC for the parameters of the DMA commands:

- **Direction**
  
  Data transfer can be in any of the following two directions as referenced from the perspective of an SPE:
  
  - For *get commands, transfer data to an LA from the main storage.
  - For *put commands, transfers data out of the LS to the main storage.

- **Size**
  
  The transfer size should obey the following guidelines:
  
  - Supported transfer sizes are 1, 2, 4, 8, or 16 bytes, and multiples of 16-bytes.
  - The maximum transfer size is 16 KB.
  - The peak performance is achieved when the transfer size is a multiple of 128 bytes.
Alignment

Alignment of the LSA and the EA should obey the following guidelines:

- The source and destination addresses must have the same four least significant bits.
- For transfer sizes less than 16 bytes, the address must be naturally aligned. Bits 28 through 31 must provide natural alignment based on the transfer size.
- For transfer sizes of 16 bytes or greater, the address must be aligned to at least a 16-byte boundary. Bits 28 through 31 must be 0.
- The peak performance is achieved when both the source and destination are aligned on a 128-byte boundary. Bits 25 through 31 must be cleared to 0.

**Header file definitions:** The `spu_mfcio.h` header file contains useful definitions, for example `MFC_MAX_DMA_SIZE`, of the supported parameter of the DMA command.

If a transaction has an illegal size or the address is invalid, due to a segment fault, a mapping fault, or other address violation, no errors will occur during compilation. Instead, during run time, the corresponding DMA command queue processing is suspended and an interrupt is raised to the PPE. The application is usually terminated in this case and a “Bus error” message is printed.

The MFC checks the validity of the effective address during the transfers. Partial transfers can be performed before the MFC encounters an invalid address and raises the interrupt to the PPE.

**Supported and recommended values for DMA-list parameters**

The following list summarizes the supported or recommended values of the MFC for the parameters of the DMA list commands:

- The parameters of each transfer (for example size, alignment) should be as described in “Supported and recommended values for DMA parameters” on page 116.
- All the data transfers that are issued in a single DMA list command have the same high 32 bits of a 64-bit effective address.
- All the data transfers that are issued in a single DMA list command share the same tag ID.
In addition, use the following supported parameters of the DMA list:

- **Length**
  A DMA list command can specify up to 2048 DMA transfers, defining up to 16 KB of memory in the LS to maintain the list. Since each transfer of this type has up to 16 KB in length, a DMA list command can transfer up to 32 MB, which is 128 times the size of the 256 KB LS.

- **Continuity**
  A DMA list can move data between a contiguous area in an LS and possibly a non-contiguous area in the effective address space.

- **Alignment**
  The local storage address of the DMA list must be aligned on an 8-byte boundary. Bits 29 through 31 must be 0.

**Header file definitions**: The `spu_mfcio.h` header file contains useful definitions, for example MFC_MAX_DMA_LIST_SIZE, of the supported parameter of the DMA list command.

**Synchronization and atomic commands**

The MFC also supports a set of synchronization and atomic commands to control the order in which DMA storage accesses are performed. The commands include four atomic commands, three send-signal commands, and three barrier commands. Synchronization can be performed for all the transactions in a queue or only to a group of them as explained in “DMA-command tag groups” on page 120. While we provide a brief overview of those commands in this section, you can find a more detailed description in 4.5, “Shared storage synchronizing and data ordering” on page 218.

Table 4-5 on page 119 shows the synchronization and atomic commands that are supported by the MFC. For each command, we mention the SPU and the PPE MFC functions that implement it, if any. A blank cell indicates that this command is not supported by neither the SPE nor PPE. For detailed information about the MFC commands, see the “DMA transfers and inter-processor communication” chapter in the *Cell Broadband Engine Programming Handbook*.18

The SPU MFC functions are defined in the `spu_mfcio.h` header file and are described in the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document.19

---

18 See note 1 on page 78.
19 See note 2 on page 80.
The PPEs are defined in the `libspe2.h` header file and are described in the *SPE Runtime Management Library* document.20

Table 4-5  Synchronization commands supported by the MFC

<table>
<thead>
<tr>
<th>Command</th>
<th>Possible Initiator</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>SPU</strong></td>
<td><strong>PPE</strong></td>
<td></td>
</tr>
</tbody>
</table>

**Synchronization commands**

<table>
<thead>
<tr>
<th>Command</th>
<th>Initiator</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>barrier</td>
<td>mfc_barrier</td>
<td>Barrier type ordering. Ensures ordering of all preceding DMA commands with respect to all commands following the <code>barrier</code> command in the same command queue. The <code>barrier</code> command has no effect on the immediate DMA commands of <code>getllar</code>, <code>putllc</code>, and <code>putlluc</code>.</td>
</tr>
<tr>
<td>mfceieio</td>
<td>mfc_eieio</td>
<td>Controls the ordering of <code>get</code> commands with respect to <code>put</code> commands, and of <code>get</code> commands with respect to <code>get</code> commands accessing storage that is caching inhibited and guarded. Also controls the ordering of <code>put</code> commands with respect to <code>put</code> commands accessing storage that is memory coherence required and not caching inhibited.</td>
</tr>
<tr>
<td>mfcsync</td>
<td>mfc_sync</td>
<td>Controls the ordering of DMA <code>put</code> and <code>get</code> operations within the specified tag group with respect to other processing units and mechanisms in the system.</td>
</tr>
<tr>
<td>sndsig</td>
<td>mfc_sndsig</td>
<td>Writes to the SPU Signal Notification Register in another device.</td>
</tr>
<tr>
<td>sndsigf</td>
<td>mfc_sndsigf</td>
<td>Writes to the SPU Signal Notification Register in another device, with the fence option.</td>
</tr>
<tr>
<td>sndsigb</td>
<td>mfc_sndsigb</td>
<td>Writes to the SPU Signal Notification Register in another device, with the barrier option.</td>
</tr>
</tbody>
</table>

**Atomic commands**

<table>
<thead>
<tr>
<th>Command</th>
<th>Initiator</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>getllar</td>
<td>mfc_getllar</td>
<td>Gets a lock line and reserve.</td>
</tr>
<tr>
<td>putllc</td>
<td>mfc_putllc</td>
<td>Puts a lock line conditional.</td>
</tr>
<tr>
<td>putlluc</td>
<td>mfc_putlluc</td>
<td>Puts lock line unconditional.</td>
</tr>
<tr>
<td>putqlluc</td>
<td>mfc_putqlluc</td>
<td>Puts a queued lock line unconditional.</td>
</tr>
</tbody>
</table>

\[a.~\text{There is no function call for implementing this command. Instead it is implemented as an intrinsic as defined in the `ppu_intrinsics.h` file.}\]

20 See note 7 on page 84.
DMA-command tag groups
All DMA commands, except the atomic ones, can be tagged with a 5-bit tag group ID. By assigning a DMA command or group of commands to different tag groups, the status of the entire tag group can be determined within a single command queue. The software can use this identifier to check or wait on the completion of all queued commands in one or more tag groups.

Tag groups can be formed separately within any of the two MFC command queues. Therefore, tags that are assigned to commands in the SPU command queue are independent of the tags that are assigned to commands in the proxy command queue of the MFC.

Tagging is useful when using barriers to control the ordering of MFC commands within a single command queue. DMA commands within a tag group can be synchronized with a fence or barrier option by appending an “f” or “b,” respectively, to the command mnemonic:

- The execution of a fenced command option is delayed until all previously issued commands within the same tag group have been performed.
- The execution of a barrier command option and all subsequent commands is delayed until all previously issued commands in the same tag group have been performed.

4.3.2 SPE-initiated DMA transfer between the LS and main storage

Software running on an SPU initiates a DMA data transfer by accessing the local MFC facilities through the channel interface. In this section, we explain how such SPU code can initiate basic data transfers between the main storage and the LS. We illustrate this concept by using the get command, which transfers data from the main storage to the LS, and by using the put command, which transfers data in the opposite direction. We also describe the getl and putl commands, which transfer data by using a DMA list.

The MFC supports additional data transfer commands, such as putf, putlb, getlf, and getb, that guarantee ordering between the data transfer. These commands are initiated similar to the way in which the basic get and put commands are initiated, although their behavior is different.

For detailed information about the channel interface and the MFC commands, see the “SPE channel and related MMIO interface” chapter and the “DMA transfers and interprocessor communication” chapter in the *Cell Broadband Engine Programming Handbook*.²¹

²¹ See note 1 on page 78.
Tag manager
The tag manager facilitates the management of tag identifiers that are used for DMA operations in an SPU application. It is implemented through a set of functions that the programmer must use to reserve tag IDs before initializing DMA transactions and release them upon completion.

The functions are defined in the `spu_mfcio.h` header file and are described in the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document. The following functions are the main ones:

- `mfc_tag_reserve`, which reserves a single tag ID
- `mfc_tag_release`, which releases a single tag ID

Some tags may be pre-allocated and used by the operating environment, for example by the software managed cache or PDT, Performance Analysis Tool. Therefore, the implementation of the tag manager does not guarantee making all 32 architected tag IDs available for user allocation. If the programmer uses a fixed value of tag IDs instead of using the tag manager, possible inefficiencies can result that are caused by waiting for DMA completions on tag groups that contain DMAs that are issued by other software components.

**Reserving a tag ID or a set of IDs:** When programming an SPU application that initiates DMAs, use the tag manager’s functions to reserve a tag ID or a set of IDs and not random or fixed values. Although it is not required, consider using tag allocation services to ensure that the other software component’s use of tag IDs does not overlap with the application’s use of tags.

Example 4-16 on page 124 shows usage of the tag manager.

Basic DMA transfer between the LS and main storage
In this section, we describe how SPU software can transfer data between the LS and main storage by using basic DMA commands. That term *basic commands* implies to commands that should be explicitly issued for each DMA transaction separately. Another alternative is to use the DMA list commands that can initialize a sequence of DMA transfers as explained in “DMA list data transfer” on page 126.

In the following sections, we explain how to initialize basic *get* and *put* DMA commands. We illustrate this concept by using a code example that also includes the use of the tag manager.

---

22 See note 2 on page 80.
Initiating a DMA transfer

To initialize a DMA transfer, the SPE programmer can call one of the corresponding functions of the `spu_mfcio.h` header file. Each function implements a single command, for example:

- The `mfc_get` function implements a `get` command.
- The `mfc_put` function implements a `put` command.

These functions are nonblocking in terms of issuing the DMA command. The software continues its execution after enqueuing the commands into the MFC SPU command queue but does not block until the DMA commands are issued on the EIB. However, these functions will block if the command queue is full and then wait until space is available in that queue. Table 4-3 on page 113 shows the full list of the supported commands.

The programmer should be aware of the fact that the implementation of these functions involves a sequence of the following six channel writes:

1. Write the LSA parameter to the MFC_LSA channel.
2. Write the effective address higher (EAH) bits parameter to the MFC_EAH channel.
3. Write the effective address lower (EAL) bits parameter to the MFC_EAL channel.
4. Write the transfer size parameter to the MFC_Size channel.
5. Write the tag ID parameter to the MFC_TagID channel.
6. Write the class ID and command opcode to the MFC_Cmd channel. The opcode command defines the transfer type, for example `get` or `put`.

**DMA parameter values:** The supported and recommended values of the different DMA parameters are discussed in “Supported and recommended values for DMA parameters” on page 116.

Waiting for completion of a DMA transfer

After the DMA command is initiated, the software might wait for completion of the DMA transaction. The programmer can call to one of the following functions that are implemented in the `spu_mfcio.h` header file:

- The `mfc_write_tag_mask` function writes the tag mask that determines to which tag IDs a completion notification is needed. This is done by using the following two functions.
- The `mfc_read_tag_status_any` function waits until any of the specified tagged DMA commands is completed.
The mfc_read_tag_status_all function waits until all of the specified tagged DMA commands are completed.

The last two functions are blocking and, therefore, will cause the software to halt until all DMA transfers that are related to the tag ID are complete. Refer to Table 4-3 on page 113 for a full list of the supported commands.

The implementation of the first function generates the channel operation to set the bit that represents the tag ID by writing the corresponding value to the MFC_WrTagMask channel. All bits are 0 beside the bit number tag ID.

The implementation of the next two functions involves a sequence of the following two channel operations:

1. Write an MFC_TAG_UPDATE_ALL or MFC_TAG_UPDATE_ANY mask to the MFC_WrTagUpdate channel.
2. Read the MFC_RdTagStat channel.

Basic DMA get and put transfers (code example)

In this section, we show how the SPU code can perform basic get and put commands as well as other relevant issues. The example demonstrates the following techniques:

- The SPU code uses the tag manager to reserve and release the tag ID.
- The SPU code uses the get command to transfer data from the main storage to the LS.
- The SPU code uses the put command to transfer data from the LS to the main storage.
- The SPU code waits for completion of the get and put commands.
- The SPU macro waits for completion of DMA group related to input tag.
- The PPU macro rounds the input value to the next higher multiple of either 16 or 128 to fulfill the DMA requirements of the MFC.

As mentioned in 4.2.3, “SPU programming methods to access the MFC channel interface” on page 101, we use the MFC functions method to access the DMA mechanism. Each of the functions implements few of the steps mentioned previously, resulting in simpler code. From a programmer’s point of view, you must be familiar with the number of commands that are involved to understand the impact on the application execution.

Example 4-16 on page 124 and Example 4-17 on page 125 contain the corresponding SPU and PPU code respectively.
Example 4-16  SPU initiated basic DMA between LS and main storage - SPU code

```
#include <spu_mfcio.h>

// Macro for waiting to completion of DMA group related to input tag:
// 1. Write tag mask
// 2. Read status which is blocked untill all tag's DMA are completed
#define waitag(t) mfc_write_tag_mask(1<<t); mfc_read_tag_status_all();

// Local store buffer: DMA address and size alignment:
// - MUST be 16B aligned otherwise a bus error is generated
// - may be 128B aligned to get better performance
// In this case we use 16B because we don’t care about performance
volatile char str[256] __attribute__ ((aligned(16)));

// argp - effective address pointer to the string in main storage
// envp - size of string in main memory in bytes
int main( uint64_t spuid , uint64_t argp, uint64_t envp ){
  uint32_t tag_id = mfc_tag_reserve();

  // reserve a tag from the tag manager
  if (tag_id==MFC_TAG_INVALID){
    printf("SPE: ERROR can't allocate tag ID\n"); return -1;
  }

  // get data from main storage to local store
  mfc_get((void *)(str), argp, (uint32_t)envp, tag_id, 0, 0);

  // wait for 'get' command to complete. wait only on this tag_id.
  waitag(tag_id);

  printf("SPE: %s\n", str);
  strcpy(str, "Am I there? No! I'm still here! I will go there again....");

  // put data to main storage from local store
  mfc_put((void *)(str), argp, (uint32_t)envp, tag_id, 0, 0);

  // wait for 'get' command to complete. wait only on this tag_id.
```
waitag(tag_id);

  // release the tag from the tag manager
mfc_tag_release(tag_id);

  return (0);
}

Example 4-17  SPU initiated basic DMA between LS and main storage - PPU code
#include <libspe2.h>

  // macro for rounding input value to the next higher multiple of either
  // 16 or 128 (to fulfill MFC's DMA requirements)
#define spu_mfc_ceil128(value)  ((value + 127) & ~127)
#define spu_mfc_ceil16(value)   ((value +  15) &  ~15)

volatile char str[256]  __attribute__ ((aligned(16)));

int main(int argc, char *argv[])
{
  void *spe_argp, *spe_envp;
spe_context_ptr_t spe_ctx;
spe_program_handle_t *program;
uint32_t entry = SPE_DEFAULT_ENTRY;

  // Prepare SPE parameters
  strcpy( str, "I am here but I want to go there!"");
  printf("PPE: %s\n", str);

  spe_argp=(void*)str;
spe_envp=(void*)strlen(str);
spe_envp=(void*)spu_mfc.ceil16((uint32_t)spe_envp);//round up to 16B

  // Initialize and run the SPE thread using the four functions:
  // 1) spe_context_create 2) spe_image_open
  // 3) spe_program_load 4) spe_context_run

  // Wait for SPE thread to complete using spe_context_destroy
  // function (blocked untill SPE thread was complete).
printf("PPE: %s\n", str); // is he already there?
return (0);
}

**DMA list data transfer**

A DMA list is a sequence of transfer elements (or list elements) that, together with an initiating DMA list command, specify a sequence of DMA transfers between a single *continuous area* of the LS and *possibly discontinuous areas* in the main storage. Therefore, the DMA lists can be used to implement scatter-gather functions between the main storage and LS. All the data transfers that are issued in a single DMA list command share the same tag ID and are the same type of commands (*getl*, *putl*, or another command). The DMA list is stored in the LS of the same SPE.

In the following sections, we describe the steps that a programmer who wants to initiate a sequence of transfers by using a DMA-list should perform:

- In “Creating a DMA list” on page 126, we create and initialize the DMA list in the LS of an SPE. Either the local SPE, the PPE, or another SPE can do this step.
- In “Initiating the DMA list command” on page 127, we issue a DMA list command, such as *getl* or *putl*. Such DMA list commands can only be issued by programs that run on the local SPE.
- In “Waiting for completion of the data transfer” on page 128, we wait for the completion of the data transfers.
- Finally, in “DMA list transfer: Code example” on page 129, we provide a code example that illustrates the sequence of steps.

**Creating a DMA list**

Each transfer element in the DMA list contains three parameters:

- *notify* refers to the stall-and-notify flag that can be used to suspend list execution after transferring a list element whose stall-and-notify bit is set.
- *size* refers to the transfer size in bytes.
- *eal* refers to the lower 32-bits of an effective address in main storage.

---

**DMA-list parameter values**: The supported and recommended values of the DMA-list parameters are discussed in “Supported and recommended values for DMA-list parameters” on page 117.
The SPU software creates the list and stores it in the LS. The list basic element is an `mfc_list_element` structure that describes a single data transfer. This structure is defined in the `spu_mfcio.h` header file as shown in Example 4-18.

**Example 4-18   DMA list basic element - mfc_list_element struct**

typedef struct mfc_list_element {
    uint64_t notify : 1; // optional stall-and-notify flag
    uint64_t reserved : 16; // the name speaks for itself
    uint64_t size : 15; // transfer size in bytes
    uint64_t eal : 32; // lower 32-bits of an EA in main storage
} mfc_list_element_t;

Transfer elements are processed sequentially in the order in which they are stored. If the `notify` flag is set for a transfer element, the MFC stops processing the DMA list after performing the transfer for that element until the SPE program sends an acknowledgement. This procedure is described in “Waiting for completion of the data transfer” on page 128.

**Initiating the DMA list command**

After the list is stored in the LS, the execution of the list is initiated by a DMA list command, such as `getl` or `putl`, from the SPE whose LS contains the list. To initialize a DMA list transfer, the SPE programmer can call one of the corresponding functions of the `spu_mfcio.h` header file. Each function implements a single DMA list command such as in the following examples:

- The `mfc_getl` function implements the `getl` command.
- The `mfc_putl` function implements the `putl` command.

These functions are nonblocking in terms of issuing the DMA command. The software continues its execution after enqueueing the commands into the MFC SPU command queue but does not block until the DMA commands are issued on the EIB. However, these functions will block if the command queue is full and will wait until space is available in that queue. Refer to Table 4-3 on page 113 for the full list of supported commands.

Initializing a DMA list command requires similar steps and parameters as when initializing a basic DMA command. See “Initiating a DMA transfer” on page 122. However, a DMA list command requires two different types of parameters than those required by a single-transfer DMA command:

- EAL, which is written to the MFC_EAL channel, should be the starting LSA of the DMA list, rather than with the EAL that is specified in each transfer element separately.
- Transfer size, which is written to the MFC_Size channel, should be the size in bytes of the DMA list itself, rather than the transfer size that is specified in
each transfer element separately. The list size is equal to the number of
transfer elements, multiplied by the size of the mfc_list_element structure
(8 bytes).

The starting LSA and the EAH are specified only once in the DMA list command
that initiates the transfers. The LSA is internally incremented based on the
amount of data transferred by each transfer element. However, if the starting LSA
for each transfer element in a list does not begin on a 16-byte boundary, then the
hardware automatically increments the LSA to the next 16-byte boundary. The
EAL for each transfer element is in the 4 GB area defined by the EAH.

Waiting for completion of the data transfer
Two mechanisms enable the software to verify the completion of the DMA
transfers. The first mechanism is the same as basic (non-list) DMA commands
using MFC_WrTagMask and MFC_RdTagStat channels. It can be used to notify
the software about the completion of the entire transfer in the DMA list. This
procedure is explained in “Waiting for completion of a DMA transfer” on
page 122.

The second mechanism is to use the stall-and-notify flag that enables the
software to be notified about the completion of subset of the transfers in the list
by the MFC. The MFC halts the transfers in the list (but not only the operations)
until it is acknowledged by the software. This mechanism can be useful if the
software needs to update the characteristics of stalled subsequent transfers,
depending on the data that was just transferred to the LS on the previous
transfers. In any case, the number of elements in the queued DMA list cannot be
changed.

To use this mechanism, the SPE software and the local MFC perform the
following actions:
1. The software enables the stall-and-notify event of the DMA list command,
   which is illustrated in the notify_event_enable function of Example 4-20 on
   page 131.
2. The software sets the notify bit in a certain element in the DMA list for it to
   indicate when it is done.
3. The software issues a DMA list command on this list.
4. The MFC stops processing the DMA list after performing the transfer for that
   specific element, which activates the DMA list command stall-and-notify
   event.
5. The software handles the event, optionally modifies the subsequent transfer
   elements before they are processed by the MFC, and then acknowledges the
   MFC. This step is illustrated in the notify_event_handler function of
   Example 4-20 on page 131.
6. The MFC continues processing the subsequent transfer elements in the list, until perhaps another element sets the notify bit.

**DMA list transfer: Code example**

In this section, we provide a code example that shows how the SPU program can initiate a DMA list transfer. The example demonstrate the following techniques:

- The SPU code creates a DMA list on the LS.
- The SPU code activates the stall-and-notify bit in some of the elements in the list.
- The SPU code uses `spe_mfcio.h` definitions to check if the DMA transfer attributes are legal.
- The SPU code issues the `getl` command to transfer data from the main storage to the LS.
- The SPU code issues the `putl` command to transfer data from the LS to the main storage.
- The SPU code implements an *event handler* to the stall-and-notify events.
- The SPU code dynamically updates DMA list according to the data that was just transferred into the LS.
- The PPU code marks the SPU code to stop transferring data after some data elements by using the stall-and-notify mechanism.
- The PPU and SPU code synchronizes the completion of SPEs writing the output data. It is implemented by using a notification flag in the main storage and a barrier between writing the data to memory and updating the notification flag.

Example 4-19 shows the shared header file.

**Source code:** The code in Example 4-19 and Example 4-20 on page 131 is included in the additional material for this book. See “SPU initiated DMA list transfers between LS and main storage” on page 619 for more information.

Example 4-19  **SPU initiated DMA list transfers between LS and main storage - Shared header file**

```c
// common.h file

// DMA list parameters
#define DMA_LIST_LEN 512
#define ELEM_PER_DMA 16 // Guarantee alignment to 128 B
#define NOTIFY_INCR 16
```
#define TOTA_NUM_ELEM   ELEM_PER_DMA*DMA_LIST_LEN
#define BUFF_SIZE       TOTA_NUM_ELEM+128
#define MAX_LIST_SIZE   2048  // 2K

// commands and status definitions
#define CMD_EMPTY       0
#define CMD_GO          1
#define CMD_STOP        2
#define CMD_DONE        3
#define STATUS_DONE     1234567
#define STATUS_NO_DONE  ~(STATUS_DONE)

// data elements that SPE should work on
#define DATA_LEN 15

typedef struct {
    char cmd;
    char data[DATA_LEN];
} data_elem; // aligned to 16B

// the context that PPE forward to SPE
typedef struct{
    uint64_t ea_in;
    uint64_t ea_out;
    uint32_t elem_per_dma;
    uint32_t tot_num_elem;
    uint64_t status;
} parm_context; // aligned to 16B

#define MIN(a,b) (((a)>(b)) ? (b) : (a))
#define MAX(a,b) (((a)>(b)) ? (a) : (b))

// dummy function for calculating the output from the input
inline char calc_out_d( char in_d ){
    return in_d-1;
}
Example 4-20 shows the SPU code.

```c
#include <spu_intrinsics.h>
#include <spu_mfcio.h>
#include "common.h"

// Macro for waiting to completion of DMA group related to input tag
#define waitag(t) mfc_write_tag_mask(1<<t); mfc_read_tag_status_all();

static parm_context ctx __attribute__ ((aligned (128)));

// DMA data structures and and data buffer
volatile data_elem lsa_data[BUFF_SIZE] __attribute__ ((aligned (128)));
volatile mfc_list_element_t dma_list[MAX_LIST_SIZE] __attribute__ ((aligned (128)));
volatile uint32_t status __attribute__ ((aligned(128)));

// global variables
int elem_per_dma, tot_num_elem, byte_per_dma, byte_tota, dma_list_len;
int event_num=1, continue_dma=1;
int notify_incr=NOTIFY_INCR;

// enables stall-and-notify event
//==================================
static inline void notify_event_enable(  )
{
    uint32_t eve_mask;

    eve_mask = spu_read_event_mask();
    spu_write_event_mask(eve_mask | MFC_LIST_STALL_NOTIFY_EVENT);
}

// updates the remaining DMA list according to data that was already
// transferred to LS
//==================================
static inline void notify_event_update_list( )
{
    int i, j, start, end;

    start = (event_num-1)*notify_incr*elem_per_dma;
    end   = event_num*notify_incr*elem_per_dma-1;
```
// loop on only data elements that were transferred since last event
for (i=start; i<=end; i++){
  if ( lsa_data[i].cmd == CMD_STOP){

    // PPE wants us to stop DMAs - zero remaining DMAs
    dma_list[event_num*notify_incr+1].size=0;
    dma_list[dma_list_len-1].size=0;
    for (j=event_num*notify_incr; j<dma_list_len; j++){
      dma_list[j].size = 0;
      dma_list[j].notify = 0;
    }
    continue_dma = 0;
    break;
  }
}

// handle stall-and-notify event include acknowledging the MFC
//===================================================================
static inline void notify_event_handler( uint32_t tag_id )
{
  uint32_t eve_mask, tag_mask;

  // blocking function to wait for event
  eve_mask = spu_read_event_mask();

  spu_write_event_mask(eve_mask | MFC_LIST_STALL_NOTIFY_EVENT);

  // loop for checking that event is on the correct tag_id
  do{
    // loop for checking that stall-and-notify event occurred
    do{
      eve_mask = spu_read_event_status();

    }while ( !(eve_mask&(uint32_t)MFC_LIST_STALL_NOTIFY_EVENT) );

    // disable event stall-and-notify event
    eve_mask = spu_read_event_mask();
    spu_write_event_mask(eve_mask & (~MFC_LIST_STALL_NOTIFY_EVENT));

    // acknowledge stall-and-notify event
    spu_write_event_ack(MFC_LIST_STALL_NOTIFY_EVENT);

    // read the tag_id that caused the event. no information is
    // provided on which DMA list command in the tag group has
// stalled or which element in the DMA list command has stalled
tag_mask = mfc_read_list_stall_status();

}while (!tag_mask & (uint32_t)(1<<(tag_id)) );

// update DMA list according to data that was just transferred to LS
notify_event_update_list();

// acknowledge the MFC to continue
mfc_write_list_stall_ack(tag_id);

// re-enable the event
eve_mask = spu_read_event_mask();
spu_write_event_mask(eve_mask | MFC_LIST_STALL_NOTIFY_EVENT);
}

void exit_handler( uint32_t tag_id ){

// update the status so PPE knows that all data is in place
status = STATUS_DONE;

// barrier to ensure data is written to memory before writing status
mfc_putb((void*)&status, ctx.status, sizeof(uint32_t), tag_id,0,0);
waitag(tag_id);

mfc_tag_release(tag_id); // release tag ID before exiting
printf("<SPE: done\n");
}

int main(int speid , uint64_t argp){
    int i, j, num_notify_events;
    uint32_t addr, tag_id;

    // enable the stall-and-notify
    //============================================
    notify_event_enable( );

    // reserve DMA tag ID
    //============================================
    tag_id = mfc_tag_reserve();

    if(tag_id==MFC_TAG_INVALID){
        printf("SPE: ERROR - can't reserve a tag ID\n");
        return 1;
    }
get context information from system memory.

```
// get context information from system memory.
//=================================================================================
mfc_get((void*) &ctx, argp, sizeof(ctx), tag_id, 0, 0);
waitag(tag_id);
```

initialize DMA transfer attributes

```
// initialize DMA transfer attributes
//=================================================================================
tot_num_elem = ctx.tot_num_elem;
 elem_per_dma = ctx.elem_per_dma;
dma_list_len = MAX( 1, tot_num_elem/ elem_per_dma );
 byte_tota = tot_num_elem*sizeof(data_elem);
 byte_per_dma = elem_per_dma*sizeof(data_elem);
```

initialize data buffer

```
// initialize data buffer
//=================================================================================
for (i=0; i<tot_num_elem; ++i){
   lsa_data[i].cmd = CMD_EMPTY;
}
```

use spe_mfcio.h definitions to check DMA attributes' legitimate

```
// use spe_mfcio.h definitions to check DMA attributes' legitimate
//=================================================================================
if (byte_per_dma<MFC_MIN_DMA_SIZE || byte_per_dma>MFC_MAX_DMA_SIZE){
   printf("SPE: ERROR - illegal DMA transfer's size\n");
   exit_handler( tag_id ); return 1;
}
if (dma_list_len<MFC_MIN_DMA_LIST_SIZE ||
   dma_list_len>MFC_MAX_DMA_LIST_SIZE){
   printf("SPE: ERROR - illegal DMA list size.\n");
   exit_handler( tag_id ); return 1;
}
if (dma_list_len>=MAX_LIST_SIZE){
   printf("SPE: ERROR - DMA list size bigger then local list \n");
   exit_handler( tag_id ); return 1;
}
if(tot_num_elem>BUFF_SIZE){
   printf("SPE: ERROR - dma length bigger then local buffer\n");
   exit_handler( tag_id ); return 1;
}
```

create the DMA lists for the 'getl' command

```
// create the DMA lists for the 'getl' command
//=================================================================================
addr = mfc_ea2l(ctx.ea_in);
```
for (i=0; i<dma_list_len; i++) {
    dma_list[i].size = byte_per_dma;
    dma_list[i].eal = addr;
    dma_list[i].notify = 0;
    addr += byte_per_dma;
}

// update stall-and-notify bit EVERY ‘notify_incr’ DMA elements
num_notify_events = 0;
for (i=notify_incr-1; i<(dma_list_len-1); i+=notify_incr) {
    num_notify_events++;
    dma_list[i].notify = 1;
}

// issue the DMA list 'getl' command
mfc_getl((void*)lsa_data, ctx.ea_in, (void*)dma_list,
          sizeof(mfc_list_element_t)*dma_list_len, tag_id, 0, 0);

// handle stall-and-notify events
for (event_num=1; event_num<=num_notify_events; event_num++) {
    notify_event_handler(tag_id);
    if( !continue_dma ){ // stop dma since PPE mark us to do so
        break;
    }
}

// wait for completion of the 'getl' command
waitag(tag_id);

// calculate the output data
for (i=0; i<tot_num_elem; ++i){
    lsa_data[i].cmd = CMD_DONE;
    for (j=0; j<DATA_LEN; j++){
        lsa_data[i].data[j] = calc_out_d(lsa_data[i].data[j]);
    }
}
// + update the existing DMA lists for the 'putl' command
// + update only the address since the length is the same
//================================================================
addr = mfc_ea2l(ctx.ea_out);

for (i=0; i<dma_list_len; i++) {
    dma_list[i].eal = addr;
    dma_list[i].notify = 0;
    addr += byte_per_dma;
}

// + no notification is needed for the 'putl' command

// issue the DMA list 'getl' command
//================================================================
mfc_putl((void*)lsa_data,ctx.ea_out,(void*)dma_list,
        sizeof(mfc_list_element_t)*dma_list_len,tag_id,0,0);

// wait for completion of the 'putl' command
//================================================================
waitag(tag_id);
exit_handler(tag_id);
return 0;
}

Example 4-21 shows the corresponding PPU code.

Example 4-21  SPU initiated DMA list transfers between LS and main storage - PPU code
#include <libspe2.h>
#include <cbe_mfc.h>

#include "common.h"

// data structures to work with the SPE
volatile parm_context ctx __attribute__((aligned(16)));
volatile data_elem in_data[TOTA_NUM_ELEM] __attribute__((aligned(128)));
volatile data_elem out_data[TOTA_NUM_ELEM] __attribute__((aligned(128)));
volatile uint32_t status __attribute__((aligned(128)));

// take ‘spu_data_t’ structure and ‘spu_pthread’ function from
// Example 4-5 on page 90
int main(int argc, char *argv[]) {
    spe_program_handle_t *program;
    int i, j, error=0;

    printf("PPE: Start main \n");
    status = STATUS_NO_DONE;

    // initiate input and output data
    for (i=0; i<TOTA_NUM_ELEM; i++){
        in_data[i].cmd = CMD_GO;
        out_data[i].cmd = CMD_EMPTY;

        for (j=0; j<DATA_LEN; j++){
            in_data[i].data[j] = (char)j;
            out_data[i].data[j] = 0;
        }
    }

    // ================================================================
    // tell the SPE to stop in some random element (number 3) after 10
    // stall-and-notify events.
    // ================================================================
    in_data[3+10*ELEM_PER_DMA*NOTIFY_INCR].cmd = CMD_STOP;

    // initiate SPE parameters
    ctx.ea_in    = (uint64_t)in_data;
    ctx.ea_out   = (uint64_t)out_data;
    ctx.elem_per_dma = ELEM_PER_DMA;
    ctx.tot_num_elem = TOTA_NUM_ELEM;
    ctx.status   = (uint64_t)&status;

    data.argp = (void*)&ctx;

    // ... Omitted section:
    // creates SPE contexts, load the program to the local stores,
    // run the SPE threads, and waits for SPE threads to complete.

    // (the entire source code for this example is part of the book’s
    // additional material).

    // This subject of is also described in section 4.1.2, “Task
    parallelism and managing SPE threads” on page 83
// wait for SPE data to be written into memory
while (status != STATUS_DONE);

for (i=0, error=0; i<TOTA_NUM_ELEM; i++){
    if (out_data[i].cmd != CMD_DONE){
        printf("ERROR: command is not done at index %d\n",i);
        error=1;
    }
    for (j=0; j<DATA_LEN; j++){
        if (calc_out_d(in_data[i].data[j]) != out_data[i].data[j]){
            printf("ERROR: wrong output : entry %d char %d\n",i,j);
            error=1; break;
        }
    }
    if (error) break;
}
if(error) { printf(")PPE: program was completed with error\n");
} else { printf(")PPE: program was completed successfully\n");
}
return (0);
}

4.3.3 PPU initiated DMA transfer between LS and main storage

Software running on a PPU initiates DMA data transfers between the main storage and the LS of some SPEs by accessing the MFC facilities of the SPE through the MMIO interface. In this section, we describe how such PPU code can initiate basic data transfers between the main storage and LS of some SPEs.

For detailed information about the MMIO (or direct problem state) interface and the MFC commands, see the “SPE channel and related MMIO interface” chapter and the “DMA transfers and interprocessor communication” chapter respectively in the Cell Broadband Engine Programming Handbook.23

Tag IDs: The tag ID that is used for the PPE-initiated DMA transfer is not related to the tag ID that is used by the software that runs on this SPE. Each tag is related to a different queue of the MFC. Currently no mechanism is available for allocating tag IDs on the PPE side, such as the SPE tag manager. Therefore, the programmer should use a predefined tag ID. Since tag IDs 16 to 31 are reserved for the Linux kernel, the user must use only tag IDs 0 to 15.

23 See note 1 on page 78.
Another alternative for a PPU software to access the LS of some SPEs is to map the LS to the main storage and then use regular direct memory access. This issue is discussed in “Direct PPE access to LS of some SPE” on page 145.

**Basic DMA transfer between LS and main storage**

In this section, we explain how PPU software can transfer data between the LS of some SPEs and the main storage by using basics DMA commands. The term *basic commands* implies commands that should be explicitly issued for each DMA transaction separately, unlike DMA list commands. We explain how the PPU can initialize these commands and illustrate this by using a code example of get and put commands. The available DMA commands are described in “DMA commands” on page 112.

The naming of the commands is based on an SPE-centric view. For example, put means a transfer from the SPE LS to an effective address.

**DMA commands from the PPE:** The programmer should avoid initiating DMA commands from the PPE and initiate them by the local SPE. There are three reasons for doing this. First, accessing the MMIO by the PPE is executed on the interconnect bus, which has larger latency than the SPU that accesses the local channel interface. The latency is high because the SPE problem state is mapped as guarded, cache inhibited memory. Second, the addition of this traffic reduces the available bandwidth for other resources on the interconnect bus. Third, since the PPE is an expensive resource, it is better to have the SPEs do more work instead.

**Initiating a DMA transfer**

To initialize a DMA transfer, the PPE programmer can call one of the corresponding functions of the *libspe2.h* header file. Each function implements a single command, as shown in the following examples:

- The *spe_mfcio_get* function implements the get command.
- The *spe_mfcio_put* function implements the put command.

The functions are nonblocking so that the software continues its execution after issuing the commands. Table 4-3 on page 113 provides a complete list of the supported commands.

Another alternative is to use the inline function that is defined in the *cbe_mfc.h* file and supports all the DMA commands. This function is the _spe_mfc_dma function, which enqueues a DMA request by using the values that are provided. It supports all types of DMA commands according to the value of the cmd input parameter. For example, the cmd parameter is set to MFC_PUT_CMD for the put command, it is set to MFC_GETS_CMD for the gets command, and so on. This
function blocks until the MFC queue has space available (in the \texttt{cbe\_mfc.h} file) and is preferred from a performance point of view over the \texttt{libspe2.h} functions.

**DMA commands from the PPE:** When issuing DMA commands from the PPE, using the \texttt{cbe\_mfc.h} functions are preferred from performance point of view over the \texttt{libspe2.h} functions. While the \texttt{cbe\_mfc.h} functions are not well described in the SDK documentation, they are quite straightforward and easy to use. In our examples, we used the \texttt{libspe2.h} functions.

The programmer should be aware of the fact that the implementation of the functions involves a sequence of the following commands:

1. Write the LSA (local store address) parameter to the MFC\_LSA register.
2. Write the EAH and EAL parameters to the MFC\_EAH registers respectively. The software can implement this by two 32-bit stores or one 64-bit store.
3. Write the transfer size and tag ID parameters to the MFC\_Size and MFC\_TagID registers respectively. The software can implement this by one 32-bit store (MFC\_Size in upper 16 bits, MFC\_TagID in lower 16 bits) or along MFC\_ClassID\_CMD in one 64-bit store.
4. Write the class ID and \texttt{opcode} command to the MFC\_ClassID\_CMD register. The \texttt{opcode} command defines the transfer type, for example get or put).
5. Read the MFC\_CMDStatus register by using a single 32-bit store to determine the success or failure of the attempt to enqueue a DMA command, as indicated by the two least-significant bits of returned value:
   - 0 The enqueue was successful.
   - 1 A sequence error occurred while enqueuing the DMA. For example, an interrupt occurred, and then another DMA was started within interrupt handler). The software should restart the DMA sequence by going to step 1.
   - 2 The enqueue failed due to insufficient space in the command queue. The software can either wait for space to become available before attempting the DMA transfer again or simply continue attempting to enqueue the DMA until successful (go to step 1).
   - 3 Both errors occurred.

**DMA parameter values:** The supported and recommended value of the different DMA parameters are described in “Supported and recommended values for DMA parameters” on page 116.
Waiting for completion of a DMA transfer

After the DMA command is initiated, the software might wait for completion of the DMA transaction. The programmer calls one of the functions that are defined in libspe2.h header, for example the spe_mfcio_tag_status_read function. The function input parameters include a mask that defines the group ID and blocking behavior (continue waiting until completion or quit after one read).

The programmer must be aware of the fact that the implementation of this function includes a sequence of the following actions:

1. Set the Prxy_QueryMask register to the groups of interest. Each tag ID is represented by one bit. Tag 31 is assigned the most-significant bit, and tag 0 is assigned the least-significant bit.
2. Issue an eieio instruction before reading the Prxy_TagStatus register to ensure the effects of all previous stores are complete.
3. Read the Prxy_TagStatus register.
4. If the value is a nonzero number, at least one of the tag groups of interest has completed. If you are waiting for all the tag groups of interest to complete, you can perform a logical XOR between tag group status value and the tag group query mask. A result of 0 indicates that all groups of interest are complete.
5. Repeat steps 3 and 4 until the tag groups of interest are complete.

Another alternative is to use the inline functions that are defined in cbe_mfc.h file:

- The _spe_mfc_write_tag_mask function is a nonblocking function that writes the mask value to the Prxy_QueryMask register.
- The _spe_mfc_read_tag_status_immediate function is a nonblocking function that reads the Prxy_TagStatus register and returns the value read. Before calling this function, the _spe_mfc_write_tag_mask function must be called to set the tag mask.

Various other methods are possible to wait for the completion of the DMA transfer as described in the “PPE-initiated DMA transfers” chapter in the Cell Broadband Engine Programming Handbook. We chose to show the simplest one.

---

24 See note 1 on page 78.
Basic DMA get and put transfers: Code example

In this section, we provide a code example that shows how a PPU program can initiate basic DMA transfers between the LS and main storage. This example demonstrates the following techniques:

- The PPU code maps the LS to the share memory and retrieves a pointer to its EA base.
- The SPU code uses the mailbox to send PPU the offset to its data buffer in the LS.
- The PPU code initiates the DMA put command to transfer data from the LS to the main storage. Notice that the direction of this command might be confusing.
- The PPU code waits for completion of the put command before using the data.

Example 4-22 shows the PPU code. We use the MFC functions method to access the DMA mechanism from the PPU side. Each of these functions implements a few of the actions that were mentioned previously causing the code to be simpler. From a programmer’s point of view, you must be familiar with the number of commands that are involved in order to understand the impact on its application execution.

Source code: The code that is shown in Example 4-22 and Example 4-23 on page 144 is included in the additional material for this book. See “PPU initiated DMA transfers between LS and main storage” on page 619 for more information.

Example 4-22   PPU initiated DMA transfers between LS and main storage - PPU code

```c
#include <libspe2.h>
#include <cbe_mfc.h>
#define BUFF_SIZE 1024

int main(int argc, char *argv[]){
    int ret;

    // PPU’s data buffer
    volatile char my_data[BUFF_SIZE] __attribute__ ((aligned(128)));
    ...
uint32_t tag, status;

// MUST use only tag 0-15 since 16-31 are used by kernel
tag = 7; // choose my lucky number

spe_ctx = spe_context_create (....); // create SPE context
spe_program_load (....); // load SPE program to memory
pthread_create (....); // create SPE pthread

// collect from the SPE the offset in LS of the data buffer. NOT the
// most efficient using mailbox- but sufficient for initialization
while(spe_out_mbox_read( data.spe_ctx, &ls_offset, 1)<=0);

// initiate DMA 'put' command to transfer data from LS to main storage
do{
    ret=spe_mfcio_put( spe_ctx, ls_offset, (void*)my_data, BUFF_SIZE,
                        tag, 0,0);
}while( ret!=0);

// wait for completion of the put command
ret = spe_mfcio_tag_status_read(spe_ctx,0,SPE_TAG_ALL, &status);

if(ret!=0){
    perror ("Error status was returned");
    // ‘status’ variable may provide more information
    exit (1);
}

// MUST issue synchronization command before reading the ‘put’ data
__lwsync();

printf("SPE says: %s\n", my_data);

// continue saving the world or at least managing the 16 SPEs
return (0);
Example 4-23 shows the corresponding SPU code.

```
#include <spu_intrinsics.h>
#include <spu_mfcio.h>

#define BUFF_SIZE 1024

// SPU's data buffer
volatile char my_data[BUFF_SIZE] __attribute__ ((aligned(128)));

int main(int speid, uint64_t argp)
{
    strcpy((char*)my_data, "Racheli Paz lives in La-Paz.\n" );

    // send to PPE the offest the data buffer- stalls if mailbox is full
    spu_write_out_mbox((uint32_t)my_data);

    // continue helping PPU saving the world or at least do what he says
    return 0;
}
```

### DMA list data transfer

A DMA list is a sequence of transfer elements (or list elements). Together with an initiating DMA list command, the list specifies a sequence of DMA transfers between a single continuous area of the LS and possibly discontinuous areas in the main storage.

The PPE software can participate in the initiating of such DMA list commands by creating and initializing the DMA list in the LS of an SPE. The way in which PPU software can access the LS is described in “Direct PPE access to LS of some SPE” on page 145. After such a list is created, only code running on this SPU can proceed with the execution of the command itself. Refer to “DMA list data transfer” on page 126, in which we describe the entire process.

### 4.3.4 Direct problem state access and LS-to-LS transfer

In this section, we explain how applications can directly access the LS of an SPE. The intention is for applications that do not run on this SPE but that run on either the PPE or other SPEs.
PPE access to the LS is described in the following section, “Direct PPE access to LS of some SPE”. The programmer should avoid massive use of this technique because of performance considerations.

We explain the access of LS by other SPEs in “SPU initiated LS-to-LS DMA data transfer” on page 147. For memory bandwidth reasons, use this technique whenever it fits the application structure.

**Direct PPE access to LS of some SPE**

In this section, we explain how the PPE can directly access the LS of some SPEs. The programmer should avoid frequent PPE direct access to the LS and use DMA transfer instead. However, it might be useful to use direct PPE to LS access occasionally with a small amount of data in order to control the program flow, for example to write a notification.

A code running on the PPU can access the LS by performing the following actions:

1. Map the LS to the main storage and provide an effective address pointer to the LS base address. The `spe_ls_area_get` function of the `libspe2.h` header file implements this step as explained in the *SPE Runtime Management Library* document. Note that this type of memory access is not cache coherent.

2. (Optional) From the SPE, get the offset and compare it to the LS base of the data to be read or written. This step can be implemented by using the mailbox mechanism.

3. Access this region as any regular data access to the main storage by using direct load and store instructions.

**Important:** The LS stores the SPU program's instructions, program stack, and global data structure. Therefore, use care with the PPU code in accessing the LS to avoid overriding the SPU program's components. Let the SPU code manage its LS space. Using any other communication technique, the SPU code can send to the PPE the offset of the region in the LS that the PPE can access.

---

25 See note 7 on page 84.
Example 4-24 shows code that illustrates how a PPE can access the LS of an SPE:

- The SPU program forwards the offset of the corresponding buffer in the LS to the PPE by using the mailbox mechanism.
- The PPE program maps the base address of the LS to the main storage by using the libspe function.
- The PPE program adds the buffer offset to the LS base address to retrieve the buffer effective address.
- The PPE program uses the calculated buffer address to access the LS and copy some data to it.

**Source code:** The code in Example 4-24 is included in the additional material for this book. See “Direct PPE access to LS of some SPEs” on page 619 for more information.

**Example 4-24  Direct PPE access to LS of some SPEs**

```c
#include <ppu_intrinsics.h>

uint64_t ea_ls_base; // effective address of LS base
uint32_t ls_offset; // offset (LS address) of SPE's data buffer
uint64_t ea_ls_str; // effective address of SPE's data buffer

#define BUFF_SIZE 256

int main(int argc, char *argv[]) {
    uint32_t mbx;

    // create SPE thread as shown in Example 4-3 on page 86

    // map SPE's LS to main storage and retrieve its effective address
    if( (ea_ls_base = (uint64_t)(spe_ls_area_get(data.spe_ctx))) == NULL) {
        perror("Failed map LS to main storage"); exit(1);
    }

    // read from SPE the offset to the data buffer on the LS
    while(spe_out_mbox_read(data.spe_ctx, &ls_offset, 1) <= 0) {

        // calculate the effective address of the LS data buffer
```

The code in Example 4-24 is included in the additional material for this book. See “Direct PPE access to LS of some SPEs” on page 619 for more information.
SPU initiated LS-to-LS DMA data transfer

In this section, we provide a code example that shows how the SPU program can access the LS of another SPE in the chip. The LS is mapped to an effective address in the main storage that allows SPEs to use ordinary DMA operations to transfer data to and from this LS.

Use LS-to-LS data transfer whenever it fits the application structure. This type of data transfer is efficient because it goes directly from SPE to SPE on the internal EIB without involving the main memory interface. The internal bus has a much higher bandwidth than the memory interface (up to 10 times faster) and lower latency.

The following actions enable a group of SPEs to initiate the DMA transfer between each local storage:

1. The PPE maps the local storage to the main storage and provides an effective address pointer to the local storage base addresses. The spe_ls_area_get function of the libspe2.h header file implements this step as described in the SPE Runtime Management Library document.²⁶

2. The SPEs send, to the PPE, the offset compare to the LS base of the data to be read or written.

²⁶ See note 7 on page 84.
3. The PPE provides the SPEs the LS base addresses and the data offsets of the other SPEs. This step can be implemented by using the mailbox.

4. SPEs access this region as a regular DMA transfer between the LS and effective address on the main storage.

Source code: A code example that uses LS-to-LS data transfer to implement a multistage pipeline programming mode is available as part the additional material for this book. See “Multistage pipeline using LS-to-LS DMA transfer” on page 619 for more information.

4.3.5 Facilitating random data access by using the SPU software cache

In this section, we discuss the software cache library that is a part of the SDK package and is based on the following principles:

Software cache library: In this chapter, we use the term software cache library. However, its full name is SPU software managed cache. We use the shorted name for simplicity. For more information about the library specification, see the Example Library API Reference document.\(^a\)

\(^a\) Example Library API Reference is available on the Web at the following address:

- The library provides a set of SPU function calls to manage the data on the LS and to transfer the data between the LS and the main storage.
- The library maintains a cache memory that is statically allocated on the LS.
- From a programmer’s point of view, accessing the data by using the software cache is similar to using ordinary load and store instructions, unlike the typical DMA interface of the SPU for transferring data.
- For the data on the main storage that the program tries to access, the cache mechanism first checks whether it is already in the cache (that is, in the LS). If it is, then the data is taken from there, which saves the latency of bringing the data from the main storage. Otherwise, the cache automatically and transparently to the programmer performs the DMA transfer.
- The cache provides an asynchronous interface which, like double buffering, enables the programmer to hide the memory access latency by overlapping data transfer and computation.
The library has the following advantages over using a standard SDK functions call to activate the DMA transfer:

- Better performance in some applications can be achieved by taking advantage of locality of reference and saving redundant data transfers if the corresponding data is already in the LS.

  **More information:** Refer to “When and how to use the software cache” on page 156, which has examples for applications for which the software cache provides a good solution.

- Usage of familiar load and store instructions with an effective address are easier to program in most cases.

- Since the topology and behavior of the cache is configurable, it can be easily optimized to match data access patterns, unlike most hardware caches.

- The library decreases the development time that is needed to port some applications to SPE.

The software cache functions add computation overhead compared to ordinary DMA data transfers. Therefore, in case the data access pattern is sequential, from a performance point of view, use ordinary DMA data transfer instead.

**Note:** The software cache activity is local to a single SPU, managing the data access of such an SPU program to the main storage and the LS. The software cache does not coordinate between data accesses of different SPUs to main storage, nor does it handle coherency between them.

In this section, we discuss the following topics:

- In “Main features of the software cache” on page 150, we summarize the main features of the software cache and explain how the programmer can configure them.

- In “Software cache operation modes” on page 150, we discuss the two different modes that are supported by the cache to perform either synchronous or asynchronous data transfer.

- In “Programming by using the software cache” on page 152, we show how to practically program by using the software cache and include code examples.

- In “When and how to use the software cache” on page 156, we provide examples of an application where using the software cache is beneficial.
**Main features of the software cache**

The programmer can configure many of the features that are related to the software cache topology and behavior, which can create an advantage to the software cache over the hardware cache in some cases. By configuring these features, the programmer can iteratively adjust the cache attributes to best suit the specific application that currently runs in order to achieve optimal performance.

The software cache has the following main features:

- **Associativity:** Direct mapped, 2-way, or 4-way set associative
- **Access mode:** Read-only or read-write (The former has better performance.)
- **Cache line size:** 16 bytes to 4 KB (a power of two values)

Cache line: Unless explicitly mentioned, we use the term *cache line* to define the software cache line. While the hardware cache line is fixed to 128 bytes, the software line can be configured to any power of two values between 16 bytes and 4 KB.

- **Number of lines:** 1 to 4K lines (a power of two values)
- **Data type:** Any valid data type (However, all the cache has the same type.)

To set the software cache attributes, the programmer must statically add the corresponding definition to the program code. By using the method, the cache attributes are taken into account during compilation of the SPE program (not on run time) when many of the cache structures and functions are constructed.

In addition, the programmer must assign a specific name to the software cache so that the programmer can define several separate caches in the same program. This is useful in cases where several different data structures are accessed by the program and each structure has different attributes. For example, some structures are read-only, some are read-write, some are integers, and some are single precision.

By default, the software managed cache can use the entire range of the 32 tag IDs that are available for DMA transfers. It does not take into account other application uses of tag IDs. If a program also initiates DMA transfers, which require separate tag IDs, the programmer must limit the number of tag IDs that are used by the software cache by explicitly configuring a range of tag IDs that the software cache can use.

**Software cache operation modes**

The software cache supports two different modes in which the programmer can use the cache after it is created. The two supported modes are *safe mode* and *unsafe mode*, which are discussed in following sections.
Safe mode and synchronous interface
The safe interfaces provide the programmer with a set of functions to access data simply by using the data's effective address. The software cache library performs the data transfer between the LS and the main memory transparently to the programmer and manages the data that is already in the LS.

One advantage of using this method is that the programmer does not need to worry about the LS addresses and can use effective addressees as any other PPE program does. From a programmer's point of view, this method is quite simple.

Data access function calls with this mode are done synchronously and are performed according to the following guidelines:

- If data is already in the LS (in the cache), a simple load from the LS is performed.
- If data is not currently in the LS, the software cache function performs DMA between the LS and the main storage. The program is blocked until the DMA is completed.

Such a synchronous interface has the disadvantage of having a long latency when the software cache needs to transfer data between the LS and main storage.

Unsafe mode and asynchronous interface
The unsafe mode provides a more efficient means of accessing the LS compared to the safe mode. The software cache provides functions to map effective addresses to the LS addresses. The programmer should use those LS addresses later to access the data, unlike in safe mode where the effective addresses are used.

In this mode, as in safe mode, the software cache keeps tracking the data that is already in the LS and performs a data transfer between the LS and the main storage only if the data is not currently in LS.

One advantage of using this method is that the programmer can have the software cache asynchronously prefetch the data by “touching” it. The programmer can implement a double-buffering-like data transfer. In doing so, the software cache starts transferring the next data to be processed, while the program continues performing computation on the current data.

The disadvantage of using this mode is that programming is slightly more complex. The programmer must access the data by using the LS address and use software cache functions to lock the data in case the data is updated. For
optimal performance, software cache functions for prefetching the data can be called.

**Programming by using the software cache**

In this section, we demonstrate how the programmer can use the software cache in an SPU program. The following programming techniques are shown in the next sections:

- “Constructing a software cache” shows how to construct a software cache and define its topology and behavior attributes.
- “Synchronous data access using safe mode” on page 153 shows how to perform synchronous data access by using the library’s safe mode.
- “Asynchronous data access by using unsafe mode” on page 154 shows how to perform asynchronous data access by using the library’s unsafe mode.

**Source code:** The code in Example 4-25 on page 153 through Example 4-27 on page 155 are included in the additional material for this book. See “SPU software managed cache” on page 620 for more information.

**Constructing a software cache**

To construct a software cache on an SPU program, the programmer must define the required attributes as well as other optional attributes for the cache topology and behavior. If the programmer does not define these attributes, the library sets default values for the attributes. In either case, the programmer must be familiar with all of the attributes since their values can effect the performance.

Next in the code following the definitions is the cache header file, which is in the /opt/cell/sdk/usr/spu/include/cache-api.h directory. Multiple caches can be defined in the same program by redefining the attributes and re-includeing the cache-api.h header file. The only restriction is that the CACHE_NAME must be different for each cache.

Example 4-25 on page 153 shows code for constructing the software cache. It shows the following actions:

- Constructing a software cache named MY_CACHE and defining both its mandatory and optional attributes
- Reserving the tag ID to be used by the software cache
Example 4-25  Constructing the software cache

```c
unsigned int tag_base; // should be defined before the cache

// Mandatory attributes
#define CACHE_NAME         MY_CACHE // name of the cache
#define CACHED_TYPE        int    // type of basic element in cache

// Optional attributes
#define CACHE_TYPE CACHE_TYPE_RW // rw type of cache
#define CACHELINE_LOG2SIZE 7    // 2^7 = 128 bytes cache line
#define CACHE_LOG2NWAY      2    // 2^2 = 4-way cache
#define CACHE_LOG2NSETS     4    // 2^4 = 16 sets
#define CACHE_SET_TAGID(set) (tag_base + (set & 7)) // use 8 tag IDs
#define CACHE_STATS//collect statistics
#include <cache-api.h>

int main(unsigned long long spu_id, unsigned long long parm)
{
    // reserve 8 tags for the software cache
    if((tag_base=mfc_multi_tag_reserve(8))==MFC_TAG_INVALID){
        printf( "ERROR: can't reserve a tags\n" ); return 1;
    }

    // can use the cache here
    ...

Synchronous data access using safe mode
After the caches is defined, the programmer can use its function calls to access
data. In this section, we show how to perform synchronous data access by using
the safe mode.

When using this mode, only the effective addresses are used to access the main
memory data. There is no need for the programmer to be aware of the LS
address to which the data was transferred (that is, by the software cache).

Example 4-26 on page 154 shows the code for the following actions:

- Using the safe mode to perform synchronous data access
- Flushing the cache so that the modified data is written into main memory
Reading variables from the main memory by using their effective address, modifying them, and writing them back to memory by using their effective address

Example 4-26  Synchronous data access by using safe mode

Use Example 4-25 on page 153 code to construct the cache and initialize the tag IDs

```c
int a, b;
unsigned eaddr_a, eaddr_b;

// initialize effective addresses from PPU parameter
eaddr_a = parm;
eaddr_b = parm + sizeof(int);

// read a and b from effective address
a = cache_rd(MY_CACHE, eaddr_a);
b = cache_rd(MY_CACHE, eaddr_b);

// write values into cache (no write-through to main memory)
cache_wr(MY_CACHE, eaddr_b, a);
cache_wr(MY_CACHE, eaddr_a, b);

// at this point only the variables in LS are modified

// writes all modified (dirty) cache lines back to main memory
cache_flush(MY_CACHE);
```

Asynchronous data access by using unsafe mode

In this section, we show how to perform asynchronous data access by using the unsafe mode. In addition, we show how the programmer can print cache statistics, which provide information about the cache activity. The programmer can use these statistics later to tune the cache topology and behavior.

When using this mode, the software cache maps the effective address of the data in the main memory into the local storage. The programmer should use the mapped local addresses later to use the data.
Example 4-27 shows the code for performing the following actions:

- Using unsafe mode to perform synchronous data access
- Touching a variable so that the cache starts asynchronous prefetching of the variable from the main memory to the local storage
- Waiting until the prefetched data is present in the LS before modifying it
- Flushing the cache so that the modified data is written into main memory
- Printing the software cache statistics

Example 4-27  Asynchronous data access by using the unsafe mode

```
// Use the beginning of the program from Example 4-26 on page 154
...

int *a_ptr, *b_ptr;

// asynchronously touch data 'b' so cache will start to prefetch it
b_ptr = cache_touch(MY_CACHE, eaddr_b);

// synchronously read data 'a' - blocked till data is present in LS
a_ptr = cache_rw(MY_CACHE, eaddr_a);

// MUST lock variables in LS since it will be modified.
// ensures that it will not cast out while the reference is held.
cache_lock(MY_CACHE, a_ptr);

// 'a' is locked in cache - can now safely be modified through ptr
*a_ptr = *a_ptr+10;

// blocking function that waits till 'b' is present in LS
cache_wait(MY_CACHE, b_ptr);

// need to lock 'b' since it will be updated
cache_lock(MY_CACHE, b_ptr);

// now 'b' is in cache - can now safely be modified through ptr
*b_ptr = *b_ptr+20;

// at this point only the variables in LS are modified

// writes all modified (dirty) cache lines back to main memory
cache_flush(MY_CACHE);

//print software cache statistics
```
When and how to use the software cache

In this section, we explain two cases in which you should use the software cache. In each of the following sections, we define one such case and explain why you should use the software cache in this case, mainly in regard to whether you should select safe mode or unsafe mode.

Case 1: First pass SPU implementation

The first case refers to situations in which there is a need to develop a first pass implementation of an application on the SPU in a relatively short amount of time. If the programmer uses the software cache in safe mode, the program is not significantly different nor requires more programming compared to other single processor programs, such as a program that runs on the PPE.

Case 2: Random data access with high cache hit rate

Some applications have a random or unpredicted data access pattern, which can be difficult for implementing a simple and efficient double or multi-buffering mechanism. This second case, refers to streaming applications that contain many iterations, and in each iteration, a few blocks of data are read and used as input for computing output blocks. Consider the following examples:

- Data blocks that are accessed by the program are scattered in memory and are relatively small.

Scattered in memory: The intention is not that the data itself is necessarily scattered in memory, but that each iteration of the algorithm uses several different blocks that are not continuous in memory.
An indirect mechanism: A program first reads index vectors from the main storage and those vectors contain the location of the next blocks of data that need to be processed.

Only after computing the results of current the iteration, the program knows which blocks to read next.

If the application also has a high cache hit rate, then the software cache can provide better performance compared to other techniques. Such a high rate can occur if blocks that are read in one iteration are likely to be used in the sequential iterations. Another similarity is regarding blocks that are read in some iteration and are close enough to the blocks of the previous iteration, that is in the same software cache line.

A high cache hit rate ensures that, in most cases, when a structure is accessed by the program, the corresponding data is already in the LS. Therefore, the software cache can take the data from the LS instead of transferring it again from main memory.

If the hit rate is significantly high, performing synchronous data access by using safe mode provides good performance since waiting for data transfer completion does not occur often. However, in most cases, the programmer might use asynchronous data access in the unsafe mode and measure whether performance improvement is achieved.

### 4.3.6 Automatic software caching on SPE

A simple way for an SPE to reference data on the main storage is to use an extension to the C language syntax that enables the sharing of data between an SPE and the PPE or between the SPEs. This extension makes it easier to pass pointers, so that the programmer can use the PPE to perform certain functions on behalf of the SPE. Similarly this mechanism can be used to share data between all SPEs through variables in the PPE address space. This mechanism is based on the software cache safe mode mechanism that was discussed in 4.3.5, “Facilitating random data access by using the SPU software cache” on page 148, but provides a more user friendly interface to activate it.

To use this mechanism, the programmer must use the `__ea` address space identifier when declaring a variable to indicate to the SPU compiler that a memory reference is in the remote (or effective) address space, rather than in the local storage. The compiler automatically generates code to access these data objects by DMA into the local storage and caches references to these data objects.
This identifier can be used as an extra type qualifier such as `const` or `volatile` in type and variable declarations. The programmer can qualify variable declarations in this way, but not variable definitions.

Accessing an `__ea` variable from an SPU program creates a copy of this value in the local storage of the SPU. Subsequent modifications to the value in main storage are not automatically reflected in the copy of the value in local storage. The programmer is responsible for ensuring data coherence for `__ea` variables that are accessed by both SPE and PPE programs.

Example 4-28 shows how to use this variable.

```
Example 4-28   Using the __ea variable

// Variable declared on the PPU side.
extern __ea int ppe_variable;

// Can also be used in typedefs.
typedef __ea int ppe_int;

// SPU pointer variable point to memory in main storage address space
__ea int *ppe_pointer;
```

The SPU program initiates this pointer to a valid effective address:
```
// Init the SPU pointer according to 'ea_val' which is a valid effective
// address that PPU forward to the SPU (e.g. using mailbox)
ppe_pointer = ea_val;
```

After the pointer is initiated, it can be used as any SPU pointer while the software cache maps it into DMA memory access, for example:
```
for (i = 0; i < size; i++) {
    *ppe_pointer++ = i;  // memory accesses use software cache
}
```

Another case is pointers in the LS space of the SPE that can be cast to pointers in the main storage address space. This action transforms an LS address into an equivalent address in the main storage because the LS is also mapped to the main storage domain. Consider the following example:
```
int x;
__ea int *ppe_pointer_to_x = &x;
```

The pointer variable `ppe_pointer_to_x` can be passed to the PPE process by a mailbox and be used by the PPE code to access the variable `x` in the LS. The programmer must be aware of the ordering issues in case both the PPE access
this variable (from the main storage) and the SPE access it (from the LS domain). Similarly this pointer can be used to transfer data between the LS and another LS by the SPEs.

GCC for the SPU provides the command line options shown in Table 4-6 to control the runtime behavior of programs that use the __ea extension. Many of these options specify parameters for the software-managed cache. In combination, these options cause GCC to link the program to a single software-managed cache library that satisfies those options. Table 4-6 describes these options.

Table 4-6  GCC options for supporting main storage access from the SPE

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-mea32</td>
<td>Generates code to access variables in 32-bit PPU objects. The compiler defines a preprocessor macro <strong>EA32</strong> so that applications can detect the use of this option. This is the default.</td>
</tr>
<tr>
<td>-mea64</td>
<td>Generates code to access variables in 64-bit PPU objects. The compiler defines a preprocessor macro <strong>EA64</strong> so that applications can detect the use of this option.</td>
</tr>
<tr>
<td>-mcache-size=X</td>
<td>Specifies an X KB cache size (X=8, 16, 32, 64 or 128)</td>
</tr>
<tr>
<td>-matomic-updates</td>
<td>Uses DMA atomic updates when flushing a cache line back to PPU memory. This is the default.</td>
</tr>
<tr>
<td>-mno-atomic-updates</td>
<td>Negates the -matomic-updates option.</td>
</tr>
</tbody>
</table>

You can find a complete example of using the __ea qualifiers to implement a quick sort algorithm on the SPU accessing PPE memory in the SDK /opt/cell/sdk/src/examples/ppe_address_space directory.

4.3.7 Efficient data transfers by overlapping DMA and computation

One of the unique features of the CBEA is that the DMA engines in each of the SPEs enables asynchronous data transfer. In this section, we discuss the fundamental techniques to achieve overlapping between data transfers and computation by using the DMA engines. These techniques are important because they enable a dramatic increase in the performance of many applications.
Motivation
Consider a simple SPU program that repeats the following steps:
1. Access incoming data using DMA from the main storage to the LS buffer B.
2. Wait for the transfer to complete.
3. Compute on data in buffer B.

This sequence is not efficient because it wastes a lot of time waiting for the completion of the DMA transfer and has no overlap between the data transfer and the computation. Figure 4-2 illustrates the time graph for this scenario.

![Figure 4-2 Serial computation and data transfer](image)

Double buffering
We can significantly accelerate the previous process by allocating two buffers, \( B_0 \) and \( B_1 \), and overlapping computation on one buffer with data transfer in the other. This technique is called *double buffering*, which is illustrated by the flow diagram scheme in Figure 4-3.

![Figure 4-3 Double buffering scheme](image)
Double buffering is a private class of *multibuffering*, which extends this idea by using multiple buffers in a circular queue instead of only the two buffers of double buffering. In most cases, usage of the two buffers in the double buffering case is enough to guarantee overlapping between the computation and data transfer.

However, if the software must wait for completion of the data transfer, the programmer might consider extending the number of buffers and move to a multibuffering scheme. Obviously this requires more memory on the LS, which might be a problem in some cases. Refer to “Multibuffering” on page 166 for more information about the multibuffering technique.

In the code that follows, we show an example of the double buffering mechanism. Example 4-29 is the header file, which is common to the SPE and PPE side.

**Source code:** The code in Example 4-29 through Example 4-31 on page 164 is included in the additional material for this book. See “Double buffering” on page 620 for more information.

The code demonstrates the use of the barrier on the SPE side to ensure that all the output data that SPE updates in memory is written into memory before the PPE tries to read it.

**Example 4-29  Double buffering code - Common header file**

```c
// common.h file -----------------------------------------------
#define ELEM_PER_BLOCK 1024 // # of elements to process by the SPE
#define NUM_OF_ELEM 2048*ELEM_PER_BLOCK // total # of elements

#define STATUS_DONE      1234567
#define STATUS_NO_DONE ~(STATUS_DONE)

typedef struct {
    uint32_t *in_data;
    uint32_t *out_data;
    uint32_t *status;
    int size;
} parm_context;
```

---
Example 4-30 shows the SPU code that contains the double buffering mechanism.

Example 4-30  Double buffering mechanism - SPU code

```c
#include <spu_intrinsics.h>
#include <spu_mfcio.h>
#include "common.h"

// Macro for waiting to completion of DMA group related to input tag
#define waitag(t) mfc_write_tag_mask(1<<t); mfc_read_tag_status_all();

// Local store structures and buffers.
volatile parm_context ctx __attribute__ ((aligned(16)));
volatile uint32_t ls_in_data[2][ELEM_PER_BLOCK] __attribute__ ((aligned(128)));
volatile uint32_t ls_out_data[2][ELEM_PER_BLOCK] __attribute__ ((aligned(128)));
volatile uint32_t status __attribute__ ((aligned(128)));
uint32_t tag_id[2];

int main(unsigned long long spu_id, unsigned long long argv)
{
    int buf, nxt_buf, cnt, nxt_cnt, left, i;
    volatile uint32_t *in_data, *nxt_in_data, *out_data, *nxt_out_data;

    tag_id[0] = mfc_tag_reserve();
    tag_id[1] = mfc_tag_reserve();

    // Fetch the parameter context, waiting for it to complete.
    mfc_get((void*)&ctx, (uint32_t)argv, sizeof(parm_context),
             tag_id[0], 0, 0);
    waitag(tag_id[0]);

    // Init parameters
    in_data = ctx.in_data;
    out_data = ctx.out_data;
    left = ctx.size;
    cnt = (left<ELEM_PER_BLOCK) ? left : ELEM_PER_BLOCK;

    // Prefetch first buffer of input data.
    buf = 0;
    mfc_getb((void *)(ls_in_data), (uint32_t)(in_data),
              cnt*sizeof(uint32_t), tag_id[0], 0, 0);
```

If you have any questions or need further assistance, please let me know.
while (cnt < left) {
    left -= SPU_Mbox_Statnt;

    nxt_in_data  = in_data + cnt;
    nxt_out_data = out_data + cnt;
    nxt_cnt = (left<ELEM_PER_BLOCK) ? left : ELEM_PER_BLOCK;

    // Prefetch next buffer so it is available for next iteration.
    // IMPORTANT: Put barrier so that we don't GET data before
    // the previous iteration's data is PUT.
    nxt_buf = buf^1;

    mfc_getb((void*)((char*)ls_in_data[nxt_buf][0]),
             (uint32_t)(nxt_in_data), nxt_cnt*sizeof(uint32_t),
             tag_id[nxt_buf], 0, 0);

    // Wait for previously prefetched buffer
    waitag(tag_id[buf]);

    for (i=0; i<ELEM_PER_BLOCK; i++) {
        ls_out_data[buf][i] = ~(ls_in_data[buf][i]);
    }

    // Put the output buffer back into main storage
    mfc_put((void*)((char*)ls_out_data[buf][0]),
             (uint32_t)(out_data),
             cnt*sizeof(uint32_t),tag_id[buf],0,0);

    // Advance parameters for next iteration
    in_data  = nxt_in_data;
    out_data = nxt_out_data;

    buf = nxt_buf;
    cnt = nxt_cnt;
}

// Wait for previously prefetched buffer
waitag(tag_id[buf]);

// process buffer
for (i=0; i<ELEM_PER_BLOCK; i++) {
    ls_out_data[buf][i] = ~(ls_in_data[buf][i]);
}

// Put the output buffer back into main storage
// Barrier to ensure all data is written to memory before status
mfc_putb((void*)(&ls_out_data[buf][0]), (uint32_t)(out_data),
cnt*sizeof(uint32_t), tag_id[buf],0,0);

// Wait for DMAs to complete
waitag(tag_id[buf]);

// Update status in memory so PPE knows that all data is in place
status = STATUS_DONE;

mfc_put((void*)&status, (uint32_t)(ctx.status), sizeof(uint32_t),
tag_id[buf],0,0);
waitag(tag_id[buf]);

mfc_tag_release(tag_id[0]);
mfc_tag_release(tag_id[1]);

return (0);
}

Example 4-31 shows the corresponding PPU code.

**Example 4-31  Double buffering mechanism - PPU code**

```c
#include <libspe2.h>
#include <cbe_mfc.h>
#include <pthread.h>
#include "common.h"

volatile parm_context ctx __attribute__ ((aligned(16)));
volatile uint32_t in_data[NUM_OF_ELEM] __attribute__ ((aligned(128)));
volatile uint32_t out_data[NUM_OF_ELEM] __attribute__ ((aligned(128)));
volatile uint32_t status __attribute__ ((aligned(128)));

// Take 'spu_data_t' structure and 'spu_thread' function from
// Example 4-5 on page 90

int main(int argc, char *argv[])
{
    spe_program_handle_t *program;
    int i, error;

    status = STATUS_NO_DONE;
```
// Init input buffer and zero output buffer
for (i=0; i<NUM_OF_ELEM; i++){
    in_data[i] = i;
    out_data[i] = 0;
}

ctx.in_data  = in_data;
ctx.out_data = out_data;
ctx.size     = NUM_OF_ELEM;
ctx.status   = &status;

data.argp = &ctx;

// ... Omitted section:
// creates SPE contexts, load the program to the local stores,
// run the SPE threads, and waits for SPE threads to complete.

// (the entire source code for this example is part of the book’s
// additional material).

// This subject of is also described in 4.1.2, “Task parallelism and
// managing SPE threads” on page 83

// Wait for SPE data to be written into memory
while (status != STATUS_DONE);

for (i=0, error=0; i<NUM_OF_ELEM; i++){
    if (in_data[i] != (¬out_data[i])){
        printf("ERROR: wrong output at index %d\n", i);
        error=1; break;
    }
}
if(error){ printf(")PPE: program was completed with error\n"); }else{ printf(")PPE: program was completed successfully\n"); }
return 0;
}
Multibuffering

Data buffering can be extended to use more than two buffers if there is no complete overlapping between the computation and the data transfer, causing the software to significantly wait for the completion of the DMA transfers. Extending the number of buffers extends the amount of memory that is needed to store the buffers. Therefore, the programmer should guarantee that enough space is available in the LS for doing so.

Building on similar concepts to double buffering, multibuffering uses multiple buffers in a circular queue. The following example shows the pseudo code of a multibuffering scheme:

1. Allocate multiple LS buffers, B0..Bn.
2. Initiate transfers for buffers B0..Bn. For each buffer Bi, apply the tag group identifier i to transfers involving that buffer.
3. Beginning with B0 and moving through each of the buffers in round-robin fashion:
   a. Set the tag group mask to include only the i tag and request a conditional tag status update.
   b. Compute on Bi.
   c. Initiate the next transfer on Bi.

This algorithm waits for and processes each Bi in round-robin order, regardless of when the transfers complete with respect to one another. In this regard, the algorithm uses a strongly ordered transfer model. Strongly ordered transfers are useful when the data must be processed in a known order, which happens in many streaming model applications.

4.3.8 Improving the page hit ratio by using huge pages

In this section, we explain how the programmer can use huge pages to enhance the data access and performance of a given application. We show the commands that are required for configuring huge pages on a system and a short code example on how to use the huge pages within a program. In addition, refer to Example 4-34 on page 174, which shows how to use huge pages with the NUMA API.

The huge page support on the SDK aims to address the issue of reducing the latency of the address translation mechanism on the SPEs. This mechanism is implemented by using 256 entry translation look-aside buffers (TLBs) that reside on the SPEs and store the information regarding address translation. The operating system on the PPE is responsible for managing the buffers.
The following process runs whenever the SPE tries to access data on the main storage:

1. The SPU code initiates the MFC DMA command for accessing data on the main storage and provides the effective address of the data in the main storage.

2. The SPE synergistic memory management (SMM) checks whether the effective address falls within one of the TLB entries:\(^{27}\)
   - If it exists (page hit), use this entry to translate to the real address and exit the translation process.
   - If it does not exist (a page miss), continue to step 3.

3. The SPU halts program execution and generates an external interrupt to the PPE.

4. The operating systems on the PPE allocate the page and writes the required information to the TLB of this particular SPE by using memory access to the problem state of this SPE.

5. The PPE signals to the SPE that translation is complete.

6. The MFC starts transferring the data, and the SPU code continues running.

This mechanism causes the SPU program to halt until the translation process is complete, which can take a significant amount of time. This may be not efficient if the process repeats itself many times during the program execution. However, the process occurs only the first time a page is accessed, unless the translation information in the TLB is replaced with information from other pages that are later accessed.

Therefore, using huge pages can significantly improve the performance in cases where the application operates on large data sets. In such cases, using huge pages can significantly reduce the amount of time in which this process occurs (only once for each page).

The SDK supports the huge TLB file system, which allows the programmer to reserve 16 MB huge pages of pinned, contiguous memory. For example, if 50 pages are configured, it provides 600 MB of pinned contiguous memory. In the worst case where each SPE accesses the entire memory range, a TLB miss occurs only once for each of the 50 pages since the TLB has enough room to store all of those pages. For comparison, the size of ordinary pages on the operating system that runs on the Cell/B.E. system is either 4 KB or 64 KB.

---

\(^{27}\) The SMM unit is responsible for address translation in the SPE.
The following issues are related to the huge pages mechanism:

- Configuring the huge pages is not related to a specific program but to the operating system.

- Any program that runs on the system can use the huge pages by explicitly mapping data buffers on the corresponding huge files.

- The area that is used by the huge pages is pinned in the system memory. Therefore, it equivalently reduces the amount of system memory bytes that are available for other purposes, that is any memory allocation that does not explicitly use huge pages.

To configure huge pages, a root user must execute a set of commands. The commands can be executed at any time and create memory mapped files in the /huge/ path that stores the huge pages content.

The first part of Example 4-32 on page 169 shows the commands that are required to set 20 huge pages that provide 320 MB of memory. The last four commands in this part, the groupadd, usermod, chgrp, and chmod commands, provide permission to the user of the huge pages files. Without executing the commands, only the root user can access the files later and use the huge pages. The second part of this example demonstrates how to verify whether the huge pages were successfully allocated.

In many cases, the programmer might have difficulties in configuring adequate huge pages usually because the memory is fragmented. Rebooting the system is required in those cases. The alternative and recommended way is to add the first part of the command sequence shown Example 4-32 on page 169 to the startup initialization script, such as /etc/rc.d/rc.sysinit. By doing so, the huge TLB file system is configured during the system boot.

Some programmers might use huge pages while also using NUMA to restrict memory allocation to a specific node as described in 4.3.9, “Improving memory access using NUMA” on page 171. The number of available huge pages for the specific node in this case is half of what is reported in /proc/meminfo. This is because, on Cell/B.E.-based blade systems, the huge pages are equally distributed across both memory nodes.
Example 4-32  Configuring huge pages

> Part 1: Configuring huge pages:

mkdir -p /huge
echo 20 > /proc/sys/vm/nr_hugepages
mount -t hugetlbfs nodev /huge
groupadd hugetlb
usermod -a -G hugetlb <user>
chgrp -R hugetlb /huge
chmod -R g+w /huge

> Part 2: Verify that huge pages are successfully configured:

cat /proc/meminfo

The following output should be printed:
MemTotal:   1010168 kB
MemFree:     155276 kB
...          
HugePages_Total: 20
HugePages_Free:  20
Hugepagesize: 16384 kB

After the huge pages are configured, any application can allocate data on the corresponding memory mapped file. The programmer explicitly invokes mmap of a /huge file of the specified size.

Example 4-33 on page 170 shows a code example which opens a huge page file using the open function and allocates 32 MB of private huge paged memory using mmap function (32 MB indicated by the Ox2000000 parameter of the mmap function).

Source code: The code in Example 4-33 on page 170 is included in the additional material for this book. See “Huge pages” on page 620 for more information.

The mmap function: The mmap function succeeds even if there are insufficient huge pages to satisfy the request. Upon first access to a page that cannot be backed by a huge TLB file system, the application process is terminated and the message “killed” is shown. Therefore, the programmer must ensure that the number of huge pages that are requested does not exceed the number of huge pages that are available.
Another useful standard Linux library that handles the access to huge pages from the program code is *libhugetlbs*.\(^28\) This library provides an API for dynamically managing the huge pages in a way that is similar to working with ordinary pages.

Example 4-33  PPU code for using huge pages

```c
#include <stdio.h>
#include <stdlib.h>
#include <sys/mman.h>
#include <sys/stat.h>
#include <fcntl.h>

int main(int argc, char *argv[])
{
    void *ptr;
    int fmem;
    char *mem_file = "/huge/myfile.bin";

    // open a huge pages file
    if ( (fmem = open(mem_file, O_CREAT|O_RDWR, 0755))== -1){
        perror("ERROR: Can't open huge pages file"); exit(1);
    }
    remove(mem_file);

    // map 32MB (0x2000000) huge pages file to main storage
    // get pointer to effective address
    ptr = mmap(0, 0x2000000, PROT_READ|PROT_WRITE, MAP_PRIVATE,fmem,0);
    if(ptr==NULL){
        perror("ERROR: Can't map huge pages"); exit(1);
    }
    printf("Map huge pages to 0x%llx\n",(unsigned long long int)ptr);

    // now we can use ‘ptr’ effective addr. pointer to store our data
    // for example forward to the SPEs to use it

    return (0);
}
```

\(^28\) See [http://sourceforge.net/projects/libhugetlbs](http://sourceforge.net/projects/libhugetlbs)
4.3.9 Improving memory access using NUMA

The first two Cell/B.E.-based blade systems, the BladeCenter QS20 and BladeCenter QS21 are both NUMA systems, which consist of two Cell/B.E. processors, each with its own system memory. The two processors are interconnected through a FlexIO interface by using the fully coherent Broadband Interface (BIF) protocol.

Since coherent access is guaranteed, from a software point of view, a program that runs on either of the two processors can coherently access either of the two attached memories. Therefore, the programmer can choose to ignore the NUMA architecture by having the data stored in two different memories and program as though the program runs on an SMP system. However, in many cases, doing so results in performance that is far from optimal.

The bandwidth between processor elements or processor elements and memory is greater if the access is local and does not have to communicate across the FlexIO. In addition, the access latency is slightly higher on node 1 (Cell/B.E. 1) compared to node 0 (Cell/B.E. 0), regardless of whether they are local or non-local.

To maximize the performance of a single application, the programmer can specify processor and memory binding to either reduce FlexIO traffic or exploit the aggregated bandwidth of the memory available on both nodes.

**Memory bandwidth-limited applications:** For applications that are memory bandwidth limited, consider allocating memory on both nodes and exploit the aggregated memory bandwidth. The optimal case is where the data and tasks execution can be perfectly divided between nodes. That is, the processor on node 0 primarily accesses memory on this node, and the same for node 1.

Linux provides a NUMA API to address this issue and to enable allocating memory on a specific node. When doing so, the programmer can use the NUMA API in either of the following ways:

- Allocating memory on the same processor on the current thread runs
- Guaranteeing that this thread keeps running on a specific processor (node affinity)

---

29 Refer to the white paper *A NUMA API for LINUX* on the Web at the following address:  
In the following sections, we discuss the two separate interfaces that Linux provides to control and monitor the NUMA policy and program considerations regarding NUMA:

- In “NUMA program level API (libnuma library)” on page 172, we discuss the libnuma shared library that provides an application level API.
- In “NUMA command utility (numactl)” on page 176, we discuss the numactl command utility.
- In “NUMA policy considerations” on page 177, we present the main consideration the programmer should make when deciding whether and how to use NUMA.

After NUMA is configured and the application completes its execution, the programmer can use the NUMA numastat command to retrieve statistics regarding the status of the NUMA allocation and data access on each of the nodes. This information can be used to estimate the effectiveness of the current NUMA configuration.

**NUMA program level API (libnuma library)**

Linux provides a shared library name, libnuma, which implements a set of APIs for controlling and tracing the NUMA policy. The library function calls can be called from any application-level program that allows programming flexibility and have the advantage of creating a self-contained program that manages the NUMA policy that is unique to them.

To use the NUMA API, the programmer must perform the following tasks:

- Include the numa.h header file in the source code.
- Add the -lnuma flag to the compilation command in order to link the library to the application.

Additional information is available in the man pages of this library and can be retrieved by using the man numa command.

Example 4-34 on page 174 shows a suggested method for using NUMA and a corresponding PPU code. The example is inspired by the matrix multiply demonstration of the SDK in the /opt/cell/sdk/src//demos/matrix_mul directory.

**Node:** NUMA terminology uses the term node in the following example to refer to one Cell/B.E. processor. Two are present on a Cell/B.E.-based blade.
The following principles are behind the NUMA example that we present:

1. Use the NUMA API to allocate two continuous memory regions, one on each of the node’s memories.

2. The allocation is done by using huge pages to minimize the page miss of the SPE. Notice that the huge pages are equally distributed across both memory nodes on a Cell/B.E.-based blade system. Refer to 4.3.8, “Improving the page hit ratio by using huge pages” on page 166, which further discusses huge pages.

3. Duplicate the input data structures (matrix and vector, in this case) by initiating two different copies, one on each of the regions that were allocated in step 1.

4. Use NUMA to split the SPE threads, so that each half of the thread is initiated and runs on a separate node.

5. The threads that run on node number 0 are assigned to work on the memory region that was allocated on this node, and node 1’s threads are assigned to work on node 1’s memory region.

In this example, we needed to duplicate the input data since the entire input matrix is needed for the threads. While this is not the optimal solution, in many other applications, this duplication is not necessary. The input data can be divided between the two nodes. For example, when adding two matrixes, one half of those matrixes can be located on one node’s memory and the second half can be loaded on the other node’s memory.

Consider the following additional comments regarding the combining of the NUMA API with the other SDK’s functions:

- The spe_cpu_info_get SKD function can be used to retrieve the number of physical Cell/B.E. processors and the specific number of physical SPEs that are currently available. Example 4-34 on page 174 demonstrates the usage of this function.

- The SDK affinity mechanism of the SPEs can be used in conjunction with the NUMA API to add affinity between SPEs to the SPEs to near memory binding that is provided by the NUMA. SPE affinity is relevant mainly when there is significant SPE-to-SPE communication and is discussed in 4.1.3, “Creating SPEs affinity by using a gang” on page 94.
Example 4-34  Code for using NUMA

#include <numa.h>

char *mem_file = "/huge/matrix_mul.bin";
char *mem_addr0=NULL, *mem_addr1=NULL;

#define MAX_SPUS16
#define HUGE_PAGE_SIZE(size_t)(16*1024*1024)

// main===============================================================
int main(int argc, char *argv[])
{
    int i, nodes, phys_spus, spus;
    unsigned int offset0, offset1;
    nodemask_t mask0, mask1;

    // calculate the number of SPU for the program
    spus = <number of required SPUs>;

    phys_spus = spe_cpu_info_get(SPE_COUNT_PHYSICAL_SPES, -1);

    if (spus > phys_spus) spus = phys_spus;

    // check NUMA availability and intiate NUMA data structures
    if (numa_available() >= 0) {
        nodes = numa_max_node() + 1;

        if (nodes > 1){
            // Set NUMA masks; mask0 for node # 0, mask1 for node # 1
            nodemask_zero(&mask0);
            nodemask_set(&mask0, 0);
            nodemask_zero(&mask1);
            nodemask_set(&mask1, 1);
        } else{
            printf("WARNING: Can't use NUMA - insufficient # of nodes\n");
        }
    } else{
        printf("WARNING: Can't use NUMA - numa is not available.\n");
    }

    // calculate offset on the huge pages for input buffers
    offset0 = <offset for node 0's buffer>
    offset1 = <offset for node 1's buffer>
// allocate inout buffers - mem_addr0 on node 0, mem_addr1 on node 1
mem_addr0 = allocate_buffer(offset0, &mask0);
mem_addr1 = allocate_buffer(offset1, &mask1);

// Initialize the data in mem_addr0 and mem_addr1

// Create each of the SPU threads
for (i=0; i<spus; i++) {
    if (i < spus/2) {
        // lower half of the SPE threads uses input buffer of ndoe 0
        threads[i].input_buffer = mem_addr0;

        // binds the current thread and its children to node 0
        // they will only run on the CPUs of node 0 and only be able
        // to allocate memory from this node
        numa_bind(&mask0);
    } else{
        // similarly - second half use buffer of node 1
        threads[i].input_buffer = mem_addr1;
        numa_bind(&mask1);
    }

    // create SPE thread - will be binded to run only on
    // NUMA's specified node
    spe_context_create(...);
    spe_program_load(...);
    pthread_create(...);
}

for (i=0; i<spus; i++) {
    pthread_join(...); spe_context_destroy(...);
}

// allocate_buffer=======================================================
// allocate a cacheline aligned memory buffer from huge pages or the
char * allocate_buffer(size_t size, nodemask_t *mask)
{
    char *addr;
    int fmem = -1;
    size_t huge_size;
// sets memory allocation mask. The thread will only allocate memory
// from the nodes set in 'mask'.
if (mask) {
    numa_set_membind(mask);
}

// map huge pages to memory
if ((fmem=open (mem_file, O_CREAT|O_RDWR, 0755))==-1) {
    printf("WARNING: unable to open file (errno=%d).\n", errno);
    exit(1);
}
remove(mem_file);
huge_size = (size + HUGE_PAGE_SIZE-1) & ~(HUGE_PAGE_SIZE-1);

addr=(char*)mmap(0, huge_size, PROT_READ|PROT_WRITE,
    MAP_PRIVATE,fmem,0);
if (addr==MAP_FAILED) {
    printf("ERROR: unable to mmap file (errno=%d).\n", errno);
    close (fmem); exit(1);
}

// Perform a memset to ensure the memory binding.
if (mask) {
    (void*)memset(addr, 0, size);
}
return addr;

NUMA command utility (numactl)

Linux provides the numactl command utility to enable control and trace on the
NUMA policy. The programmer can combine the numactl commands in a script
that executes the appropriate commands and later runs the application.

For example, the following command invokes a program that allocates all
processors on node 0 with a preferred memory allocation on node 0:
numactl --cpunodebind=0 --preferred=0 ./matrix_mul

The following command is a shorter version that performs the same action:
numactl -c 0 -m 0 ./matrix_mul

To read the man pages of this command, run the man numactl command.
One advantage of using this method is that there is no need to recompile the program to run with different settings of the NUMA configuration. Alternatively, using the command utility enables less flexibility to the programmer compared to calling the API of libnuma library from the program itself.

Controlling the NUMA policy by using the command utility is usually sufficient in cases where all SPU threads can run on a single Cell/B.E. processor. If more than one processor is required (usually because more than eight threads are necessary), and the application requires dynamic allocation of data, it is usually difficult to use only the command utility. Using the libnuma library API from the program is more appropriate and allows greater flexibility in this case.

**NUMA policy considerations**

Choosing an optimal NUMA policy depends upon the application’s data access patterns and communication methods. We suggest the following guidelines when the programmer must decide whether to use the NUMA commands or API and which NUMA policy to implement:

- For applications that are memory bandwidth limited, consider allocating memory on both nodes and exploiting the aggregated memory bandwidth. If possible, partition application data so that processors on node 0 primarily access memory on node 0 only. Likewise, processors on node 1 primarily access memory on node 1 only.

- Choose a NUMA policy that is compatible with typical system usage patterns. For example, if multiple applications are expected to run simultaneously, the programmer should not bind all processors to a single node forcing an overcommit scenario that leaves one of the nodes idle. In this case, we recommend not constraining the Linux scheduler with any specific bindings.

- If the access patterns are not predictable and the SPEs are allocated on both nodes, then use an interleaved memory policy, which can improve the overall memory throughput.

- In a Cell/B.E. system, use node 0 over node 1 because node 0 usually has better memory access performance.

- Consider the operating system services when choosing the NUMA policy. For example, if the application incorporates extensive GbE networking communications, the TCP stack will consume some PPU resources on node 0 for eth0. In such cases, we recommend binding the application to node 1.

- Avoid over committing processor resources. Context switching of SPE threads is not instantaneous, and the scheduler quanta for the threads of the SPE is relatively large. Scheduling overhead is minimized by not over-committing resources.
4.4 Inter-processor communication

The Cell/B.E. contains several mechanisms that enable communication between the PPE and SPEs and between the SPEs. These mechanisms are mainly implemented by the MFCs (one instance of MFC exists in each of the eight SPEs). The code that runs on an SPU can interact with the MFC of the associated SPE by using the channel interface. The PPU code or code that runs on the other SPUs can interact with this MFC by using the MMIO interface.

In the following sections, we discuss the primary communication mechanisms between the PPE and SPEs:

- In 4.4.1, “Mailboxes” on page 179, we discuss the mailbox mechanism that can send 32-bit messages to and from the SPE. This mechanism is used mainly to control communication between an SPE and the PPE or between SPEs.

- In 4.4.2, “Signal notification” on page 190, we discuss the signal notifications (signaling) mechanism that signals 32-bit messages to an SPE. This mechanism is used to control communication from the PPE or other SPEs to another SPE. It can be configured for one-sender-to-one-receiver signalling or many-senders-to-one-receiver signalling.

- In 4.4.3, “SPE events” on page 203, we explain how to use an event to create asynchronous communication between the processors.

- In 4.4.4, “Atomic unit and atomic cache” on page 211, we explain how to implement a fast-shared data structure for inter-processor communication by using the Atomic Unit and the Atomic Cache hardware mechanism.

Refer to 4.2, “Storage domains, channels, and MMIO interfaces” on page 97, in which we describe the MFC interfaces and the different programming methods in which programs can interact with the MFC. In this section, we use only the MFC functions method in order interact with the MFC.

Another mechanism to apply inter-processor communication is DMA data transfers. For example, an SPE can compute output data and use DMA to transfer this data to the main memory. Later the SPE can notify the PPE that the data is ready by using additional DMA to a notification variable in the memory that the PPE polls. See 4.3, “Data transfer” on page 110, in which we describe the available data transfer mechanisms and how the programmer can initiate them.

Both mailboxes and signals are mechanism that can be used for program control and sending short messages between the different processors. While those mechanisms have a lot in common, there are differences between the two mechanisms. In general, a mailbox implements a queue for sending separate
32-bit messages, while signaling is similar to interrupts, which can be accumulated when being written and reset when being read. Table 4-7 compares the two mechanisms.

Table 4-7 Comparison between mailboxes and signals

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Mailboxes</th>
<th>Signals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direction</td>
<td>One inbound and two outbound.</td>
<td>Two inbound (toward the SPE).</td>
</tr>
<tr>
<td>Interrupts</td>
<td>One mailbox can interrupt a PPE. Two mailbox-available event interrupts.</td>
<td>Two signal-notification event interrupts.</td>
</tr>
<tr>
<td>Message accumulation</td>
<td>No.</td>
<td>Yes, using logical OR mode (many-to-one). The alternative is overwrite mode (one-to-one).</td>
</tr>
<tr>
<td>Unique SPU commands</td>
<td>No, programs use channel reads and writes.</td>
<td>Yes, sndsig, sndsigf, and sndsigb enable an SPU to send signals to another SPE.</td>
</tr>
<tr>
<td>Destructive read</td>
<td>Reading a mailbox consumes an entry.</td>
<td>Reading a channel resets all 32 bits to 0.</td>
</tr>
<tr>
<td>Channel count</td>
<td>Indicates the number of available entries.</td>
<td>Indicates a waiting signal.</td>
</tr>
<tr>
<td>Number</td>
<td>Three mailboxes: four deep incoming, one deep outgoing, one deep outgoing with interrupt.</td>
<td>Two signal registers.</td>
</tr>
</tbody>
</table>

### 4.4.1 Mailboxes

In this section, we discuss the mailbox mechanism, which is easily enables the sending of 32-bit messages between the different processors on the chip (PPE and SPEs). In this section, we discuss the following topics:

- In “Mailbox overview” on page 180, we provide an overview about the mechanism and its hardware implementation.
- In “Programming interface for accessing mailboxes” on page 182, we describe the main software interfaces for an SPU or PPU program to use the mailbox mechanism.
- In “Blocking versus nonblocking access to the mailboxes” on page 183, we explain how the programmer can implement either blocking or nonblocking access to the mailbox on either an SPU or PPU program.
In “Practical scenarios and code examples for using mailboxes” on page 184, we provide practical scenarios and techniques for using the mailboxes and emphasize some code examples.

Monitoring the mailbox status can be done asynchronously by using events that are generated whenever a new mailbox is written or read by external source, for example, a PPE or other SPE. Refer to 4.4.3, “SPE events” on page 203, in which we discuss the events mechanism in general.

**Mailbox overview**

The mailbox mechanism is easy to use and enables the software to exchange 32-bit messages between the local SPU and the PPE or local SPU and other SPEs. The term *local SPU* refers to the SPU of the same SPE where the mailbox is located. The mailboxes are accessed from the local SPU by using the channel interface and from the PPE or other SPEs by using the MMIO interface.

**SPE communication:** Mailboxes can also be used as a communications mechanism between SPEs. This is accomplished by an SPE doing DMA of data into the mailbox of another SPE by using the effective addressed problem state mapping.

The mailbox mechanism is similar to the signaling mechanism. Table 4-7 on page 179 compares the two mechanisms.

**Mailbox access:** Local SPU access to the mailbox is internal to an SPE and has small latency (less than or equal to six cycles for nonblocking access). Alternatively, PPE or other SPE access to the mailbox is done through the local memory EIB. The result is larger latency and overloading of the bus bandwidth, especially when polling to wait for the mailbox to become available.

The MFC of each SPE contains three mailboxes divided into two categories:

- **Outbound mailboxes**
  
  Two mailboxes are used to send messages from the local SPE to the PPE or other SPEs:
  
  - SPU Write Outbound mailbox (SPU_WrOutMbox)
  - SPU Write Outbound Interrupt mailbox (SPU_WrOutIntrMbox)

- **Inbound mailbox**
  
  One mailbox, SPU Read Inbound mailbox (SPU_RdInMbox), is used to send messages to the local SPE from the PPE or other SPEs.
Table 4-8 summarizes the main attributes of the mailboxes and the differences between outbound and inbound mailboxes. It also describes the differences between accessing the mailboxes from the SPU programs and accessing them from the PPU and other SPE programs.

Table 4-8  Attributes of inbound and outbound mailboxes

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Inbound mailboxes</th>
<th>Outbound mailboxes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direction</td>
<td>Messages from the PPE or another SPEs to the local SPE.</td>
<td>Messages from the local SPE to the PPE or another SPEs.</td>
</tr>
<tr>
<td>Read/Write</td>
<td>▶ Local SPE reads. ▶ PPE writes.&lt;sup&gt;b&lt;/sup&gt;</td>
<td>▶ Local SPE write. ▶ PPE reads.&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td># mailboxes</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td># entries</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Counter&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Counts the number of valid entries: ▶ Decrement when the SPU program reads from the mailbox. ▶ Incremented when the PPU program writes to the mailbox.&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Counts the number of empty entries: ▶ Decrement when the SPU program writes to the mailbox. ▶ Incremented when the PPU program reads from the mailbox.&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Buffer</td>
<td>A first-in-first-out (FIFO) queue. The SPU program reads the oldest data first.</td>
<td>A FIFO queue. The SPU program reads the oldest data first.</td>
</tr>
<tr>
<td>Overrun</td>
<td>A PPU program that writes new data when the buffer is full overruns the last entry in the FIFO.&lt;sup&gt;b&lt;/sup&gt;</td>
<td>The SPU program that writes new data when the buffer is full blocks until space is available in the buffer. For example, PPE reads from the mailbox.&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Blocking</td>
<td>▶ The SPU program blocks when trying to read an empty buffer and continues only when there is a valid entry. For example, the PPE writes to the mailbox.&lt;sup&gt;b&lt;/sup&gt; ▶ The PPU program never blocks.&lt;sup&gt;b&lt;/sup&gt; Writing to the mailbox when full overrides the last entry, and the PPU immediately continues.</td>
<td>▶ The SPU program blocks when trying to write to the buffer when it is full and continues only when there is an empty entry. For example, the PPE reads from the mailbox.&lt;sup&gt;b&lt;/sup&gt; ▶ The PPU program never blocks.&lt;sup&gt;b&lt;/sup&gt; Reading from the mailbox when it is empty returns invalid data, and the PPU program immediately continues.</td>
</tr>
</tbody>
</table>

<sup>a</sup> This per-mailbox counter can be read by a local SPU program by using a separate channel or by the PPU or other SPUs program using a separate MMIO register.

<sup>b</sup> Or other SPE that accesses the mailbox of the local SPE.
Programming interface for accessing mailboxes

The simplest way to access the mailboxes is through the MFC functions that are part of SDK package library:

- A local SPU program can access the mailboxes by using the spu_*_mbox functions in the spu_mfcio.h header file.

- A PPU program can access the mailboxes by using the spe_*_mbox* functions in the libraries2.h header file.

- Other SPUs program can access the mailboxes by using the DMA functions of the spu_mfcio.h header file, which enables reading or writing of the mailboxes that are mapped to main storage as part of the problem state of the local SPU.

For more information about the spu_mfcio.h functions, see the “SPU mailboxes” chapter in the C/C++ Language Extensions for Cell Broadband Engine Architecture document.30 For information about the libraries2.h functions, see the “SPE mailbox functions” chapter in the SPE Runtime Management Library document.31

Table 4-9 summarizes the simple functions in the files for accessing the mailboxes from a local SPU program or from a PPU program. In addition to the value of the mailboxes messages, the counter that is mentioned in Table 4-9 can also be read by the software by using the SPU *_stat_* functions or the PPU *_status functions.

Table 4-9  MFC functions for accessing the mailboxes

<table>
<thead>
<tr>
<th>Name</th>
<th>SPU code functions (channel interface)</th>
<th>Blocking</th>
<th>PPU code functions (MMIO interface)</th>
<th>Blocking</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPU write outbound mailbox</td>
<td>spu_write_out_mbox</td>
<td>Yes</td>
<td>spe_out_mbox_read</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>spu_stat_out_mbox</td>
<td>No</td>
<td>spe_out_mbox_status</td>
<td>No</td>
</tr>
<tr>
<td>SPU write outbound int. mailbox</td>
<td>spu_write_out_intr_mbox</td>
<td>Yes</td>
<td>spe_out_intr_mbox_read</td>
<td>User3</td>
</tr>
<tr>
<td></td>
<td>spu_stat_out_intr_mbox</td>
<td>No</td>
<td>spe_out_intr_mbox_status</td>
<td>No</td>
</tr>
<tr>
<td>SPU read inbound mailbox</td>
<td>spu_read_in_mbox</td>
<td>Yes</td>
<td>spe_in_mbox_write</td>
<td>User</td>
</tr>
<tr>
<td></td>
<td>spu_stat_in_mbox</td>
<td>No</td>
<td>spe_in_mbox_status</td>
<td>No</td>
</tr>
</tbody>
</table>

30 See note 2 on page 80.
31 See note 7 on page 84.
To access a mailbox of the local SPU from other SPU, the following actions are required:

1. The PPU code maps the controls area of the SPU to main storage by using the `libspe2.h` file’s `spe_ps_area_get` function with the `SPE_CONTROL_AREA` flag set.

2. The PPE forwards the control area base address of the SPU to another SPU.

3. The other SPU uses ordinary DMA transfers to access the mailbox. The effective address should be control area base, plus offset to a specific mailbox register.

**Blocking versus nonblocking access to the mailboxes**

By using the SDK library functions for accessing the mailboxes (see “Programming interface for accessing mailboxes” on page 182), the programmer can implement either blocking or nonblocking mechanisms.

For the SPU, the instructions to access the mailbox are blocking by nature and are stalled when the mailbox is not available (empty for read or full for write). The SDK simply implements the instructions.

For the PPU, the instructions to access the mailbox are nonblocking by nature. The SDK functions provide a software abstraction of blocking behavior functions for some of the mailboxes, which is implemented by polling the mailbox counter until an entry is available.

The programmer can explicitly read the mailbox status (the counter mentioned in Table 4-8 on page 181) by calling the `_stat_*` functions for the SPU program and the `_status` functions for the PPU program.

**Nonblocking access:** The nonblocking approach is slightly more complicated to program, but enables the program to perform other tasks in case the FIFO is empty instead being stalled to wait for a valid entry.

Table 4-10 on page 184 summarizes the programming approaches for performing either blocking or nonblocking access to the mailbox on a PPU or SPU program.
Table 4-10  Blocking versus nonblocking access to mailboxes - Programming approaches

<table>
<thead>
<tr>
<th>Processor</th>
<th>Mailbox</th>
<th>Blocking</th>
<th>Nonblocking</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPU</td>
<td>In</td>
<td>Reads the mailbox by using the spu_read_in_mbox function.</td>
<td>Before reading the mailbox, it polls the counter by using the spu_stat_in_mbox function until FIFO is not empty.</td>
</tr>
<tr>
<td></td>
<td>Out</td>
<td>Writes to the mailbox by using the spu_write_out_mbox function.</td>
<td>Before writing to the mailbox, it polls the counter by using the spu_stat_out_mbox function until FIFO is not full.</td>
</tr>
<tr>
<td></td>
<td>OutIntr</td>
<td>Writes to the mailbox by using the spu_write_out_intr_mbox function.</td>
<td>Before writing to the mailbox, it polls the counter by using the spu_stat_out_intr_mbox function until FIFO is not full.</td>
</tr>
<tr>
<td>PPU</td>
<td>In</td>
<td>Calls the spe_in_mbox_write function and sets the “behavior” parameter to blocking.</td>
<td>Calls the spe_in_mbox_write function and sets the “behavior” parameter to nonblocking.</td>
</tr>
<tr>
<td></td>
<td>Out</td>
<td>Not implemented. a</td>
<td>Calls the spe_out_mbox_read function.</td>
</tr>
<tr>
<td></td>
<td>OutIntr</td>
<td>Calls the spe_out_intr_mbox_read function and sets the “behavior” parameter to blocking.</td>
<td>Calls the spe_out_intr_mbox_read function and sets the “behavior” parameter to nonblocking.</td>
</tr>
</tbody>
</table>

a. The programmer should check the function return value to see that the data that was read it valid.

Practical scenarios and code examples for using mailboxes

When using the mailbox, it is important to be aware of the following attributes of mailbox access:

- Local SPU access is internal to the SPE and has small latency (less than or equal to six cycles for nonblocking access).
- Local SPU access to a mailbox that is not available (empty for read or full for write) is blocking. To avoid blocking, the program can first read the counter as explained later.
- PPE or other SPE access is done through the local memory EIB, so that they have larger latency and overload the bus bandwidth, especially when polling the mailbox counter waiting for the mailbox to become available.
- PPU or local SPU access to a mailbox can be either blocking or nonblocking when using SDK library functions, as discussed in “Blocking versus nonblocking access to the mailboxes” on page 183.

In the following sections, we describe different scenarios for using the mailbox mechanism and provide a mailbox code example.
**Notifying the PPE about data transfer completion by using a mailbox**

A mailbox can be useful when an SPE must notify the PPE about completion of transferring data that was previously computed by the SPE to the main memory.

The implementation of this type of mechanism requires the following actions:

1. The SPU code places the computational results in main storage by using DMA.
2. The SPU code waits for the DMA transfer to complete.
3. The SPU code writes to an outbound mailbox to notify the PPE that its computation is complete. This ensures that only the LS buffers of the SPE are available for reuse but does not guarantee that data has been coherently written to main storage.
4. The PPU code reads the outbound mailbox of the SPE and is notified that computation is complete.
5. The PPU code issues an `lwsync` instruction to be sure that the results are coherently written to memory.
6. The PPU code reads the results from memory.

To implement step 4, the PPU might need to poll the mailbox status to see if there is valid data in this mailbox. Such polling is not efficient because it causes overhead on the bus bandwidth, which can affect other data transfer on the bus, such as SPEs reading from main memory.

**Alternative:** An SPU can notify the PPU that it has completed computation by using a fenced DMA to write notification to an address in the main storage. The PPU may poll this area on the memory, which may be local to the PPE in case the data is in the L2 cache, so that it minimizes the overhead on the EIB and memory subsystem. Example 4-19 on page 129, Example 4-20 on page 131, and Example 4-21 on page 136 provide code for such a mechanism.

**Exchanging parameters between the PPE and SPE by using a mailbox**

A mailbox can be used for any short-data transfer purpose, such as sending storage effective addresses from the PPE to SPE.

Because the operating system runs on the PPE, only the PPE originally is aware of the effective addresses of different variables in the program, for example, when the PPE dynamically allocates data buffers or when it maps the local storage or problem state of the SPE to an effective address on the main storage. The inbound mailboxes can be used to transfer those addresses to the SPE.
Example 4-35 and Example 4-36 on page 188 provide code for such a mechanism.

Similarly, any type of function or command parameters can be forwarded from the PPE to the SPE by using this mechanism.

In the other direction, an SPE can use the outbound mailbox to notify the PPE about a local storage offset of a buffer that is located on the local storage and can be accessed later by either the PPE or another SPE. Refer to the following section, “Code example for using mailboxes”, for a code example for such a mechanism.

**Code example for using mailboxes**
The code examples in this section cover the following techniques:

- Example 4-35 shows the PPU code for accessing the SPE mailboxes by using either nonblocking methods or blocking methods. Most of the methods described in the previous sections are illustrated. This example also shows how to map the control area of the SPEs to the main storage to enable the SPEs to access each other's mailbox.

- Example 4-36 on page 188 shows the SPU code for accessing the local mailboxes by using either nonblocking methods and blocking methods. The code also sends a mailbox to the mailbox of another SPE.

- Example 4-39 on page 199 shows the functions that implement the writing to a mailbox of another SPE by using DMA transactions. The code also contains functions for reading the status of the mailbox of another SPE.

**Source code:** The code in Example 4-35, Example 4-36 on page 188, and Example 4-39 on page 199 is included in the additional material for this book. See “Simple mailbox” on page 620 for more information.

**Example 4-35  PPU code for accessing the SPEs’ mailboxes**

```c
// add the ordinary SDK and C libraries header files...
// take 'spu_data_t' structure and 'spu_pthread' function from
// Example 4-5 on page 90

extern spe_program_handle_t spu;
volatile parm_context ctx[2] __attribute__ ((aligned(16)));
volatile spe_spu_control_area_t* mfc_ctl[2];

// main====================================================================
int main()
{
```
int num, ack;
uint64_t ea;
char str[2][8] = {"0 is up","1 is down"};

for( num=0; num<2; num++){
    // SPE_MAP_PS flag should be set when creating SPE context
    data[num].spe_ctx = spe_context_create(SPE_MAP_PS,NULL);
}

// ... Omitted section:
// load the program to the local stores, and run the SPE threads.

// (the entire source code for this example is part of the book's
// additional material)

// This is also described in 4.1.2, “Task parallelism and managing
SPE threads” on page 83

// STEP 0: map SPEs’ MFC problem state to main storage (get EA)
for( num=0; num<2; num++){
    if (!mfc_ctl[num] = (spe_spu_control_area_t*)spe_ps_area_get(
        data[num].spe_ctx, SPE_CONTROL_AREA))
    {
        perror("Failed mapping MFC control area");exit (1);
    }
}

// STEP 1: send each SPE its number using BLOCKING mailbox write
for( num=0; num<2; num++){

    // write 1 entry to in_mailbox
    // we don't know if we have available space so use blocking
    spe_in_mbox_write(data[num].spe_ctx,(uint32_t*)&num,1,
    SPE_MBOX_ALL_BLOCKING);
}

// STEP 2: send each SPE the EA of other SPE's MFC area and a string
// Use NON-BLOCKING mailbox write after first verifying
// availability of space.
for( num=0; num<2; num++){

    ea = (uint64_t)mfc_ctl[(num==0)?1:0];

    // loop till we have 4 entries available
    while(spe_in_mbox_status(data[num].spe_ctx)<4){
        // PPE can do other things meanwhile before check status again
    }
Example 4-36  SPU code for accessing local mailboxes and another SPE’s mailbox

// add the ordinary SDK and C libraries header files...
#include "spu_mfcio_ext.h"  // the file described in Example 4-39 on page 199

uint32_t my_num;

// Macro for waiting to completion of DMA group related to input tag:
#define waitag(t) mfc_write_tag_mask(1<<t); mfc_read_tag_status_all();

int main( )
{
    uint32_t data[2],ret, mbx, ea_mfc_h, ea_mfc_l, tag_id;
    uint64_t ea_mfc;

if ((tag_id = mfc_tag_reserve()) == MFC_TAG_INVALID){
    printf("SPE: ERROR can't allocate tag ID\n"); return -1;
}

// STEP 1: read from PPE my number using BLOCKING mailbox read
while( spu_stat_in_mbox() <= 0 );
my_num   = spu_read_in_mbox();

// STEP 2: receive from PPE the EA of other SPE's MFC and string
// use BLOCKING mailbox, but to avoid bloking we first read
// status to check that we have 4 valid entries
while( spu_stat_in_mbox() < 4 ){
    // SPE can do other things meanwhile before check status again
}

    ea_mfc_h = spu_read_in_mbox(); // read EA lower bits
    ea_mfc_l = spu_read_in_mbox(); // read EA higher bits

data[0] = spu_read_in_mbox(); // read 4 bytes of string
    data[1] = spu_read_in_mbox(); // read 4 more bytes of string

    ea_mfc = mfc_hl2ea( ea_mfc_h, ea_mfc_l);

// STEP 3: send my ID as acknowledge to PPE using BLOCKING mbx write
spu_write_out_mbox( my_num + 1313000 ); //add dummy constant to pad MSb

// STEP 4: write message to other SPE's mailbox using BLOCKING write
mbx = my_num + 1212000; //add dummy constant to pad MSb

    ret = write_in_mbox( mbx, ea_mfc, tag_id);
    if (ret!=1){ printf("SPE: fail sending to other SPE\n");return -1;}

// STEP 5: read mailbox written by other SPE
    data[0] = spu_read_in_mbox();

        return 0;
}
4.4.2 Signal notification

In this section, we discuss the signal notification mechanism, which is an easy-to-use mechanism that enables a PPU program or SPU program to signal a program that is running on another SPU.

We discuss the following topics in this section:

- In “Signal notification overview” on page 190, we provide an overview of this mechanism and its hardware implementation.
- In “Programming interface for accessing signaling” on page 192, we describe the main software interfaces for an SPU or PPU program to use the signal notification mechanism.
- In “Practical scenarios and code examples for using signaling” on page 193, we provide practical scenarios and techniques for using the signal notification and emphasize some code examples. We also provide printing macros for tracing inter-processor communication, such as sending mailboxes and signaling between PPE and SPE and SPE and SPE. These macros can be useful when tracing a flow of a given parallel program.

Monitoring the signal status can be done asynchronously by using events that are generated whenever a new signal is sent by an external source, for example a PPE or other SPE. Refer to 4.4.3, “SPE events” on page 203, in which we discuss the events mechanism in general.

**Signal notification overview**

Signal notification is an easy-to-use mechanism that enables a PPU program to signal an SPE by using 32-bit registers. It also enables an SPU program to signal a program that is running on another SPU by using the signal mechanism of the other SPU. The term *local SPU* is used in this section to define the SPU of the same SPE where the signal register is located.

Each SPE contains two identical signal notification registers: Signal Notification 1 (SPU_RdSigNotify1) and Signal Notification 2 (SPU_RdSigNotify2).

Unlike the mailboxes, the signal notification has only one direction and can send information toward the SPU that resides in the same SPE as the signal registers (and not vice versa). Programs can access the signals by using the following interfaces:

- A local SPU program reads the signal notification by using the channel interface.
- A PPU program signals an SPE by writing the MMIO interface to it.
An SPU program signals another SPE by using special signaling commands, which include \texttt{sndsig}, \texttt{sndsigf}, and \texttt{sndsigb}. (The “f” and “b” suffix in the commands indicate “fence” and “barrier” respectively.) The commands are implemented by using the DMA \texttt{put} commands and optionally contain ordering information.

When the local SPU program reads a signal notification, the value of the signal’s register is reset to 0. Reading the signal’s MMIO (or problem state) register by the PPU or other SPUs does not reset their value.

Regarding writing the PPU or other SPUs to the signal registers, two different modes can be configured:

- **OR mode (many-to-one)**
  
  The MFC accumulates several writes to the signal-notification register by combining all the values that are written to this register by using a logical OR operation. The register is reset when the SPU reads it.

- **Overwrite mode (one-to-one)**
  
  Writing a value to a signal-notification register overwrites the value in this register. This mode is similar to using an inbound mailbox and has similar performance.

Configuring the signaling mode can be done by the PPU when it creates the corresponding SPE context.

**OR mode:** By using the OR mode, signal producers can send their signals at any time and independently of other signal producers. (No synchronization is needed.) When the SPU program reads the signal notification register, it becomes aware of all the signals that have been sent since the most recent read of the register.

Similar to the mailboxes, the signal notification register maintains a counter, which has a different behavior in the signaling case:

- The counter indicates only whether there are pending signals (at least one bit set) and not the number of writes to the register that have taken place.

- A value of 1 indicates that there is at least one event pending, and a value of 0 indicates that no signals are pending.

- The counter can be read by a program that is running on either the local SPU, PPU, or other SPUs.
In regard to the blocking behavior, accessing the signal notification has the following characteristics:

- PPU code writing to the signal register is nonblocking. Whether it overrides its previous value depends on the configured mode (OR or overwrite mode as explained previously).
- SPU code writing to the signal register of another SPU behaves similarly to the DMA put command and blocks only if the MFC FIFO is full.
- A local SPU reading the signal register is blocking when no events are pending. Reading is completed immediately in case there is at least one pending event.

Table 4-7 on page 179 summarizes the similarities and differences between the signal notification and mailbox mechanism.

**Programming interface for accessing signaling**

The simplest way to access the signal notification mechanism is through the MFC functions that are part of SDK package library:

- The local SPU program can read the local SPE’s signals by using the `spu_read_signal*` and `spu_stat_signal*` functions in the `spu_mfcio.h` header file for reading the signal register and the status (counter) respectively.
- The other SPU program can signal other SPUs by using the `mfc_sndsig*` functions (where the `*` is “b,” “f,” or blank) in the `spu_mfcio.h` header file, which enables signaling of the other SPU by doing a write operation on its memory mapped problem state.
- The PPU program can access the signals by using two main functions in the `libspe2.h` header file:
  - The `spe_signal_write` function to send a signal to an SPU
  - Optionally setting the `SPE_CFG_SIGNOTIFY1_OR` flag when creating the SPE context (by using the `spe_context_create` function) to enable OR mode.

The `spu_mfcio.h` functions are described in the “SPU signal notification” chapter in the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document. The `libspe2.h` functions are described in the “SPE SPU signal notification functions” chapter in the *SPE Runtime Management Library* document.

---

32 See note 2 on page 80.
33 See note 7 on page 84.
To signal a local SPU from another SPU, the following actions are required:

1. The PPU code maps the signaling area of the SPU to the main storage by using the `spe_ps_area_get` function of the `libspe2.h` file with the `SPE_SIG_NOTIFY_x_AREA` flag set.

2. The PPE forwards the signaling area base address of the SPU to another SPU.

3. Another SPU uses the `mfc_sndsig` function of the `spu_mfcio.h` file to access the signals. The effective address should signal the area base plus offset to a specific signal register.

The programmer can take either the blocking or nonblocking approach when reading the signals from the local SPU. The programming methods for either approach are similar to the approach for mailboxes as explained in “Blocking versus nonblocking access to the mailboxes” on page 183. However, setting signals from the PPU program or other SPUs is always nonblocking.

**Practical scenarios and code examples for using signaling**

Similar to the mailboxes mechanism, local SPU access to the signals notification is internal to an SPE and have small latency (less than 6 cycles for nonblocking access). PPE or MMIO access of other SPUs to the mailbox are done through the local memory EIB, which has larger latency. However, because it is uncommon (and usually not useful) to poll the signal notification from the MMIO side, overloading the bus bandwidth is not a significant issue.

In regard to blocking behavior, a local SPU that reads the signals register when no bits are set is blocking. To avoid blocking, the program can first read the counter. The PPE or other SPUs that signal another SPU is always nonblocking.

When using the OR mode, the PPE or other SPUs usually do not need to poll the signals counter because the events are accumulated. Otherwise (overwrite mode) the signals have a similar behavior to inbound mailboxes.

In the following sections, we describe two different scenarios for using the signals notification mechanism. We also provide a signals code example.

Because the signals behave similar to mailboxes in overwrite mode, the scenarios for using this mode are similar to those described in “Practical scenarios and code examples for using mailboxes” on page 184.

**A signal value as a processor ID**

In this section, we describe one useful scenario for using the OR mode. This mode can be useful when one processor needs to asynchronously send a notification, for example, about reaching a certain step in the program, to an SPE.
and uses the signal value to identify which processor has sent the signal. In this scenario, the SPE can receive notification from different sources.

The following sequence is suggested for implementing such a mechanism:

1. Each processor (PPE and SPE) is assigned with one bit in the signaling register.
2. A processor that wants to signal an SPE writes to the signal register of the SPE with the processor’s corresponding bit set to 1 and other bits set to 0.
3. An SPE that reads its signal register checks which bits are set. For each bit that is set, the SPE knows that the corresponding processor has sent a signal to this SPE.
4. The SPE that receives the signal can then obtain more information from the sending processor, for example, by reading its mailbox or memory.

**A signal value as an event ID**

In this section, we describe one useful scenario for using the OR mode. The OR mode can be useful when a single source processor, usually the PPE, must asynchronously send notification about an event (for example, about the need to execute some command) to an SPE (or a few SPEs). In this scenario, there are several different events in the program, and the signal value is used to identify which event has occurred at this time.

The following sequence is suggested to implement such a mechanism:

1. Each event in the program is assigned with one bit in the signaling register.
2. A PPE that wants to signal an SPE about an event writes to the signal register of the SPE with the event’s corresponding bit set to 1 and other bits are 0.
3. An SPE that reads its signal register checks which bits are set. For each bit that is set, the SPE knows that the corresponding event occurred and handles it.

**Code example for using signals notification**

The code samples that follow show the following techniques:

- Example 4-37 on page 195 shows the PPU code that signals an SPE. Because the SPE is configured to the OR mode, we use nonblocking access. The example also shows how to map the signaling area of the SPEs to the main storage to enable the SPEs to signal each other.

- Example 4-38 on page 197 shows the SPU code that reads the local signals by using either nonblocking methods or blocking methods. The SPUs signal each other in a loop until they receive an asynchronous signal from the PPU to stop.
Example 4-39 on page 199 shows the SPU code that contains functions that access the mailbox and signals of another SPE. The code also contains functions for writing to the other SPE’s mailbox and reading the mailbox status by using DMA transactions.

Example 4-40 on page 202 shows the PPU and SPU printing macros for tracing interprocessor communication, such as sending mailboxes and signaling between the PPE and SPE and the SPE and SPE.

Source code: The code in Example 4-37 through Example 4-40 on page 202 is included in the additional material for this book. See “Simple signals” on page 621 for more information.

Example 4-37  PPU code for signaling the SPEs

```c
#include <cbea_map.h>
#include <com_print.h> // the code from Example 4-40 on page 202
extern spe_program_handle_t spu;

volatile spe_sig_notify_1_area_t *ea_sig1[2];
volatile spe_sig_notify_2_area_t *ea_sig2[2];

int main()
{
    int num, ret[2], mbx[2];
    uint32_t sig=0x80000000; // bit 31 indicates signal from PPE
    uint64_t ea;

    for( num=0; num<2; num++){
        // SPE_MAP_PS flag should be set when creating SPE context
        data[num].spe_ctx = spe_context_create(SPE_MAP_PS, NULL);
    }

    // ... Omitted section:
    // load the program to the local stores. and run the SPE threads

    // (the entire source code for this example is part of the book’s
    // additional material).

    // This subject of is also described in TBD_REF: Chapter 4.1.2 Task
    parallelism and managing SPE threads
```
// STEP 0: map SPE's signals area to main storage (get EA)
for( num=0; num<2; num++){
    if ((ea_sig1[num] = (spe_sig_notify_1_area_t*)spe_ps_area_get(data[num].spe_ctx, SPE_SIG_NOTIFY_1_AREA))==NULL){
        perror("Failed mapping Signal1 area");exit (1);
    }
    if ((ea_sig2[num] = (spe_sig_notify_2_area_t*)spe_ps_area_get(data[num].spe_ctx, SPE_SIG_NOTIFY_2_AREA))==NULL){
        perror("Failed mapping Signal2 area");exit (1);
    }
}

// STEP 1: send each SPE the EA of the other SPE's signals area
// first time writing to SPE so we know mailbox has 4 entries empty
for( num=0; num<2; num++){
    spe_in_mbox_write(data[num].spe_ctx, (uint32_t*)&num,1,
        SPE_MBOX_ANY_NONBLOCKING);
    ea = (uint64_t)ea_sig1[(num==0)?1:0];
    spe_in_mbox_write(data[num].spe_ctx, (uint32_t*)&ea,2,
        SPE_MBOX_ANY_NONBLOCKING);

    // wait we have 2 entries free and then send the last 2 entries
    while(spe_in_mbox_status(data[num].spe_ctx)<2);
    ea = (uint64_t)ea_sig2[(num==0)?1:0];
    spe_in_mbox_write(data[num].spe_ctx, (uint32_t*)&ea,2,
        SPE_MBOX_ANY_NONBLOCKING);
}

// STEP 2: wait for SPEs to start signaling loop
for( num=0; num<2; num++){
    while(!spe_out_mbox_status(data[num].spe_ctx));
    spe_out_mbox_read(data[num].spe_ctx, (uint32_t*)&mbx[num], 1);
    prn_p_mbx_s2m(4,1,sig);
};

// STEP 3: wait for while - let SPEs signal one to another
for( num=0; num<20000000; num++){
    mbx[0] = mbx[0] *2;
}

// STEP 4: send the SPEs a signal to stop
prn_p_sig_m2s(4,0,sig);
prn_p_sig_m2s(4,1,sig);
```c
ret[0] = spe_signal_write(data[0].spe_ctx, SPE_SIG_NOTIFY_REG_1, sig);
ret[1] = spe_signal_write(data[1].spe_ctx, SPE_SIG_NOTIFY_REG_2, sig);

if (ret[0]==-1 || ret[1]==-1){
    perror("Failed writing signal to SPEs"); exit (1);
}

// STEP 5: wait till SPEs tell me that they're done
for( num=0; num<2; num++){
    while(!spe_out_mbox_status(data[num].spe_ctx));
    spe_out_mbox_read(data[num].spe_ctx, (uint32_t*)&mbx[num], 1);
    prn_p_mbx_s2m(5,num,mbx[num]);
}

// STEP 6: tell SPEs that they can complete execution
for( num=0; num<2; num++){
    mbx[num] = ~mbx[num];
    spe_in_mbox_write(data[num].spe_ctx, (uint32_t*)&mbx[num], 2,
        SPE_MBOX_ANY_NONBLOCKING);
    prn_p_mbx_m2s(6,num,mbx[num]);
}

// ... Omitted section:
// waits for SPE threads to complete.

// (the entire source code for this example is part of the book’s
// additional material).

return (0);
```

**Example 4-38  SPU code for reading local signals and signaling other SPE**

// add the ordinary SDK and C libraries header files...
#include "spu_mfcio_ext.h" // the code from Example 4-39 on page 199
#include <com_print.h> // the code from Example 4-40 on page 202

#define NUM_ITER 1000

uint32_t num;

// Macro for waiting to completion of DMA group related to input tag:
# 1. Write tag mask. 2. Read status untill all tag’s DMA are completed

```c
#define waitag(t) mfc_write_tag_mask(1<<t); mfc_read_tag_status_all();
```

```c
int main()
{
  uint32_t in_sig,out_sig,mbx,idx,i,ea_h,ea_l,tag_id;
  uint64_t ea_sig[2];

  if ((tag_id= mfc_tag_reserve())==MFC_TAG_INVALID){
    printf("SPE: ERROR can't allocate tag ID\n"); return -1;
  }

  // STEP 1: read from PPE my number using BLOCKING mailbox read
  num     = spu_read_in_mbox();
  idx     = (num==0)?1:0;
  out_sig = (1<<num);

  // STEP 2: receive from PPE EA of other SPE's signal area and string
  while( spu_stat_in_mbox()<4 ); //wait till we have 4 entries
  for (i=0;i<2;i++){
    ea_h = spu_read_in_mbox(); // read EA lower bits
    ea_l = spu_read_in_mbox(); // read EA higher bits
    ea_sig[i] = mfc_hl2ea( ea_h, ea_l);
  }

  // STEP 3: Tell the PPE that we are going to start loopoing
  mbx = 0x44332211; prn_s_mbx_m2p(3,num,mbx);
  spu_write_out_mbox( mbx );

  // STEP 4: Start looping- signal other SPE and read my signal
  if (num==0){
    write_signal2( out_sig, ea_sig[idx], tag_id);
    while(1){
      in_sig = spu_read_signal1();
      if (in_sig&0x80000000){ break; } // PPE signals us to stop
    }
  }else{ //num==1
    write_signal2( out_sig, ea_sig[idx], tag_id);
  }
}
```
while(1){
    in_sig = spu_read_signal2();
    if (in_sig&0x80000000){ break; } // PPE signals us to stop

    if (in_sig&0x00000001){ // receive signal from other SPE
        prn_s_sig_m2s(4,num,out_sig);
        write_signal1( out_sig, ea_sig[idx], tag_id);
    }else{
        printf("}}SPE%d<<NA: <%08x>
",num,in_sig); return -1;
    } }  

prn_s_sig_p2m(4,num,in_sig);

// STEP 5: tell the PPE that we're done
mbx = 0x11223344*(num+1); prn_s_mbx_m2p(5,num,mbx);
spu_write_out_mbox( mbx );

// STEP 6: block mailbox from PPE to not finish before other SPE
mbx = spu_read_in_mbox(); prn_s_mbx_p2m(5,num,mbx);
mfc_tag_release(tag_id);
return 0;
}

Example 4-39   SPU code for accessing the mailbox and signals of another SPE

spu_mfcio_ext.h

#include <spu_intrinsics.h>
#include <spu_mfcio.h>

static uint32_t msg[4]__attribute__ ((aligned (16)));

// mailbox status register definitions
#define SPU_IN_MBOX_OFFSET 0x0C // offset from control area base
#define SPU_IN_MBOX_OFFSET_SLOT 0x3 // 16B alignment= (OFFSET&0xF)>>2

// mailbox status register definitions
#define SPU_MBOX_STAT_OFFSET 0x14 // offset from control area base
#define SPU_MBOX_STAT_OFFSET_SLOT 0x1 // 16B alignment= (OFFSET&0xF)>>2
// signal notify 1 and 2 registers definitions
#define SPU_SIG_NOTIFY_OFFSET 0x0C // offset from signal areas base
#define SPU_SIG_NOTIFY_OFFSET_SLOT 0x3 // 16B alignment (OFFSET&0xF)>>2

// returns the value of mailbox status register of remote SPE
inline int status_mbox(uint64_t ea_mfc, uint32_t tag_id)
{
    uint32_t status[4], idx;
    uint64_t ea_stat_mbox = ea_mfc + SPU_MBOX_STAT_OFFSET;

    idx = SPU_MBOX_STAT_OFFSET_SLOT;

    mfc_get((void *)&status[idx], ea_stat_mbox, sizeof(uint32_t),
            tag_id, 0, 0);
    mfc_write_tag_mask(1<<tag_id);
    mfc_read_tag_status_any();

    return status[idx];
}

// returns the status (counter) of inbound_mailbox of remote SPE
inline int status_in_mbox(uint64_t ea_mfc, uint32_t tag_id)
{
    int status = status_mbox( ea_mfc, tag_id);
    status = (status&0x0000ff00)>>8;
    return status;
}

// returns the status (counter) of outbound_mailbox of remote SPE
inline int status_out_mbox(uint64_t ea_mfc, uint32_t tag_id)
{
    int status = status_mbox( ea_mfc, tag_id);
    status = (status&0x000000ff);
    return status;
}

//returns status (counter) of inbound_interrupt_mailbox of remote SPE
inline int status_outintr_mbox(uint64_t ea_mfc, uint32_t tag_id)
{
    int status = status_mbox( ea_mfc, tag_id);
    status = (status&0xffff0000)>>16;
    return status;
}
// writing to a remote SPE's inbound mailbox
inline int write_in_mbox(uint32_t data, uint64_t ea_mfc,
                       uint32_t tag_id)
{
    int status;
    uint64_t ea_in_mbox = ea_mfc + SPU_IN_MBOX_OFFSET;
    uint32_t mbx[4], idx;

    while( (status = status_in_mbox(ea_mfc, tag_id))<1);

    idx = SPU_IN_MBOX_OFFSET_SLOT;
    mbx[idx] = data;

    mfc_put((void *)&mbx[idx], ea_in_mbox, sizeof(uint32_t), tag_id, 0, 0);
    mfc_write_tag_mask(1<<tag_id);
    mfc_read_tag_status_any();

    return 1; // number of mailbox being written
}

// signal a remote SPE's signal1 register
inline int write_signal1(uint32_t data, uint64_t ea_sig1,
                         uint32_t tag_id)
{
    uint64_t ea_sig1_notify = ea_sig1 + SPU_SIG_NOTIFY_OFFSET;
    uint32_t idx;

    idx = SPU_SIG_NOTIFY_OFFSET_SLOT;
    msg[idx] = data;

    mfc_sndsig( &msg[idx], ea_sig1_notify, tag_id, 0, 0);
    mfc_write_tag_mask(1<<tag_id);
    mfc_read_tag_status_any();

    return 1; // number of mailbox being written
}

// signal a remote SPE's signal1 register
inline int write_signal2(uint32_t data, uint64_t ea_sig2, uint32_t tag_id)
{
    uint64_t ea_sig2_notify = ea_sig2 + SPU_SIG_NOTIFY_OFFSET;
    uint32_t idx;

    idx = SPU_SIG_NOTIFY_OFFSET_SLOT;

msg[idx] = data;

mfc_sndsig( &msg[idx], ea_sig2_notify, tag_id, 0,0);
mfc_write_tag_mask(1<<tag_id);
mfc_read_tag_status_any();

return 1; // number of mailbox being written

Example 4-40   PPU and SPU macros for tracing inter-processor communication

com_print.h ======================================================
// add the ordinary SDK and C libraries header files...

// Printing macros for tracing PPE-SPE and SPE-SPE communication
// Syntax: prn_X_Y_Z2W:
// X: 'p' when printing from the PPE, 's' printing from SPE
// Y: 'mbx' for mailbox, 'sig' for signaling
// Z: 'm' source is me, 's' source SPE, 'p' source PPE
// W: 'm' destination is me, 's' destination SPE, 'p' destination PPE
// Parameters (i,s,m) stands for:
// i: some user-defined index for example step # in program execution
// s: For PPE - # of SPE with-which we communicate,
//     For SPE - # of local SPE
// m: message value (mailbox 32b value, signal value)

#define prn_p_mbx_m2s(i,s,m) printf("%d)PPE>>SPE%02u: <%08x>\n",i,s,m);
#define prn_p_mbx_s2m(i,s,m) printf("%d)PPE<<SPE%02u: <%08x>\n",i,s,m);
#define prn_p_sig_m2s(i,s,m) printf("%d)PPE->SPE%02u: <%08x>\n",i,s,m);
#define prn_p_sig_s2m(i,s,m) printf("%d)PPE<-SPE%02u: <%08x>\n",i,s,m);
#define prn_s_mbx_m2p(i,s,m) printf("%d{SPE%02u>>PPE: <%08x>\n",i,s,m);
#define prn_s_mbx_p2m(i,s,m) printf("%d{SPE%02u<<PPE: <%08x>\n",i,s,m);
#define prn_s_mbx_m2s(i,s,m) printf("%d{SPE%02u<-SPE: <%08x>\n",i,s,m);
#define prn_s_mbx_s2m(i,s,m) printf("%d{SPE%02u->SPE: <%08x>\n",i,s,m);
#define prn_s_sig_m2p(i,s,m) printf("%d{SPE%02u->PPE: <%08x>\n",i,s,m);
#define prn_s_sig_p2m(i,s,m) printf("%d{SPE%02u<-PPE: <%08x>\n",i,s,m);
#define prn_s_sig_m2s(i,s,m) printf("%d{SPE%02u->SPE: <%08x>\n",i,s,m);
#define prn_s_sig_s2m(i,s,m) printf("%d{SPE%02u<-SPE: <%08x>\n",i,s,m);
4.4.3 SPE events

In this section, we discuss the SPE events mechanism that enables code that runs on the SPU to trace events that are external to the program execution. The SDK package provides a software interface that also enables a PPE program to trace events that occurred on the SPEs.

We include the following topics in this section:

- In “SPE events overview” on page 203, we provide an overview about the SPE events mechanism and its hardware implementation.
- In “Programming interface for accessing events” on page 205, we describe the main software interfaces for an SPU or PPU program to use the SPE events mechanism.
- In “Practical scenarios and code example for using events” on page 206, we provide practical scenarios and techniques for using the mailboxes and emphasize the code examples.

SPE events overview

With the SPE events mechanism, code that runs on the SPU can trace events that are external to the program execution. The events can either be set internally by the hardware of this specific SPE or, due to such external events as sending mailbox messages of signal notification, set by the PPE or the SPEs.

In addition, the SDK package provides a software interface so that a PPE program can trace events that occurred on the SPEs and create an event handler to service those events. Only a subset of four events is supported by this mechanism. Refer to “Programming interface for accessing events” on page 205 for a discussion about this mechanism.

The main events that can be monitored fall into the following categories:

- MFC DMA
  These events are related to the DMA commands of the MFC. Specifically, refer to Example 4-20 on page 131 which shows an example for handling the MFC DMA list command stall-and-notify.
- Mailbox or signals
  This category refers to the external write or read to a mailbox or signal notification registers.
Synchronization events
These events are related to multisource synchronization or a lock line reservation (atomic) operation.

Decrementer
These events are set whenever the decrementer’s elapsed time has expired.

The events are generated asynchronously to the program execution, but the programmer can choose to monitor and correspond to those events either synchronously or asynchronously:

Synchronous monitoring
The program explicitly checks the events status in one of the following ways:
– Nonblocking
  It polls for pending events by testing the event counts in a loop.
– Blocking
  It reads the event status, which stalls when no events are pending.

Asynchronous monitoring
The program implements an event interrupt handler.

Intermediate approach
The program sprinkles bisled instructions, either manually or automatically throughout the application code by using code-generation tools, so that they are executed frequently enough to approximate asynchronous event detection.

Four different 32-bit channels can enable SPU software to manage the events mechanism. The channels have an identical bit definition, while each event is represented by a single bit. The SPE software should use the following sequence of actions to deal with SPE events:

1. Initialize event handling by write to the “SPU Write Event Mask” channel and set the bits that correspond to the events that the program wants to monitor.
2. Monitor some events that are pending by using either a synchronous, asynchronous, or intermediate approach as described previously.
3. Recognize which events are pending by reading from the “SPU Read Event Status” channel and see which bits were set.
4. Clear events by writing a value to “SPU Write Event Acknowledge” and set the bit that corresponds to the pending events in the written value.
5. Service the events by executing application-specific code to handle the specific events that are pending.
Similarly to the mailbox or signal notification mechanism, each of the registers maintains a counter that can be read by the SPU software. The only counter that is usually relevant to the software is the one related to the “SPU Read Status” channel. The software can read this channel to detect the number of events that are pending. If the counter returns a value of 0, then no enabled events are pending. If the counter returns a value of 1, then enabled events have been raised since the last read of the status.

Table 4-11 summarizes the four available SPE event channels.

<table>
<thead>
<tr>
<th>Name</th>
<th>RW</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPU Write Event Mask (SPU_WrEventMask)</td>
<td>W</td>
<td>To enable only the events that are relevant to its operation, the SPU program can initialize a mask value with the event bits set to 1 only for the relevant events.</td>
</tr>
<tr>
<td>SPU Read Event Status (SPU_RdEventStat)</td>
<td>R</td>
<td>Reading this channel reports events that are both pending at the time of the channel read and are enabled. The corresponding bit is set in “SPU Write Event Mask.”</td>
</tr>
<tr>
<td>SPU Write Event Acknowledgment (SPU_WrEventAck)</td>
<td>W</td>
<td>Before an SPE program services the events reported in “SPU Read Event Status,” it should write a value to the “SPU Write Event Acknowledge” to acknowledge (clear) the events that will be processed. Each bit in the written value acknowledges the corresponding event.</td>
</tr>
<tr>
<td>SPU Read Event Mask (SPU_RdEventMask)</td>
<td>R</td>
<td>This channel enables the software to read the value that was recently written to “SPU Write Event Mask.”</td>
</tr>
</tbody>
</table>

**Programming interface for accessing events**

The simplest way to access the events is through the MFC functions that are part of the SDK package library:

- The SPU programmer can manage events with the following functions in the spu_mfcio.h header file:
  - Enable events by using the spu_read_event_mask and spu_write_event_mask functions, which access “Event Mask” channel.
  - Monitor events by using the spu_read_event_status and spu_stat_event_status functions, which read the value and counter of the “Event Status” channel.
  - Acknowledge events by using the spu_write_event_ack function, which writes into the “Event Acknowledgment” channel.
  - Retrieve the events that are pending by using the MFC_*_EVENT that is defined, for example MFC_SIGNAL_NOTIFY_1_EVENT and MFC_OUT_MBOXAVAILABLE_EVENT)
The PPU program can trace the events that are set on the SPE and implement an event handler by using several functions in the libspe library (SPE Runtime Management, defined in the libspe2.h header file):

- Use the spe_event_handler_* functions to create, register, unregister, and destroy the event handler.
- Use the spe_event_wait function to synchronously wait for events. This is a semi-blocking function. It is stalled until the input timeout parameter expires.
- Use the other functions to service the detected event. Use the appropriate function depending on the event. For example, use the functions for reading the mailbox when a mailbox-related event occurred.

The spu_mfcio.h functions are described in the “SPU event” chapter in the C/C++ Language Extensions for Cell Broadband Engine Architecture document. The libspe2.h functions are described in “SPE event handling” chapter in the SPE Runtime Management Library document.

The programmer can use either a blocking or nonblocking approach when reading events from the local SPU. The programming methods for these approaches are similar to those discussed for the mailboxes in “Blocking versus nonblocking access to the mailboxes” on page 183. However, reading events from the PPE side is a semi-blocking function, which is stalled until the input timeout parameter expires.

There is no specific mechanism to allow an SPE to trace the events of another SPE, but it might be possible to implement such a mechanism in the software. In most cases, such a mechanism is not practical.

**Practical scenarios and code example for using events**

Similar to the mailbox mechanism, local SPU access to the event channels is internal to the SPE and has small latency (less than or equal to six cycles for nonblocking access). PPE or other SPE access to the event registers has higher latency.

Regarding blocking behavior, a local SPU that reads the events register when no bits are set is blocking. To avoid blocking, the program can first read the counter.

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34 See note 2 on page 80.
35 See note 7 on page 84.
Based on the event that is monitored, the events can be used for the following scenarios:

- **DMA list dynamic updates**
  
  Monitor stall-notify-event to update the DMA list according to the data that was transferred to the local storage from the main storage. See Example 4-20 on page 131 for this type of scenario.

- **Profiling or implementing a watchdog on an SPU program**
  
  Use the decrementer to periodically profile the program or implement a watchdog about the program execution.

Example 4-41 on page 209 shows another scenario for using the SPE events, in which code is included for implementing an event handler on the PPU.

**SPU as computation server**

An SPE event can be used to implement a mechanism in which the SPU acts as a computation server that executes commands that are generated and forwarded to it by the PPU code.

One option is to implement it as an *asynchronous* computation server. The SPU program implements the asynchronous events handler mechanism for handling incoming mailboxes from the PPE:

1. The SPU code asynchronously waits for an inbound mailbox event.
2. The PPU code forwards to the SPU the commands that should be executed (and possibly other information) by writing commands to the inbound mailbox.
3. The SPU code monitors the pending mailbox event and determines the command that should be executed.
4. Additional information can be forward from the PPU to the SPU by using more mailbox messages or DMA transfer.
5. The SPU processes the command.

The SPU side for such a mechanism can be implemented as an interrupt (events) handler as explained in the “Developing a basic interrupt handler” chapter in the *Cell Broadband Engine Programming Handbook.*

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36 See note 1 on page 78.
Another option is to implement a synchronous computation server on the SPU side and implement the event handler on the PPU side:

- The SPU code synchronously polls and executes the commands that are defined in its inbound mailbox.
- The PPU code implements an event handler for the SPU events. Whenever the PPU monitors the SPU that has read the mailbox, it writes the next command to the SPU mailbox.

Refer to the following section, “PPU code example for implementing the SPE events handler”, in which we suggest how to implement such an event handler on the PPU.

The second synchronous computation server can have advantages when compared to the asynchronous version. It allows overlapping between different commands because the PPU can write to the SPU the next command at the same time that the SPU is working on the current command.

There is a large latency between the generation of the SPE events and the execution of corresponding PPU event handler (which involves running kernel functions). For this reason, only if the delay between one command and another is large, then using the second synchronous computation server make sense and provides good performance results.

**PPU code example for implementing the SPE events handler**

In this section, we show how PPU code can implement a handler for SPE events. The code contains a simplified version of the PPU program for implementing the synchronous computation server that is described in “SPU as computation server” on page 207. There is a large latency between the generation of the SPE event and the execution of the corresponding PPU event handler (roughly 100K cycles) because it involves running kernel functions.

Example 4-41 on page 209 contains the corresponding PPU code that creates and registers an event handler for monitoring whenever the inbound mailbox is no longer full. Any time the mailbox is not full, which indicates that the SPU has read a command from it, the PPU puts new commands in the mailbox.

The goal of this example is only to demonstrate how to implement a PPE handler for SPE events and use the event of an SPE read from an inbound mailbox. While supporting only this type of event is not always practical, it can be easily extended to support a few different types of other events. For example, it can support an event in which an SPE has stopped execution, a PPE-initiated DMA operation has completed, or an SPE has written to the outbound mailbox. For more information, see the method for making a callback to the PPE side of the SPE thread (stop and signal mechanism) as described in the “PPE-assisted library facilities” chapter in the *SPE Runtime Management Library* document.37
The SPU code is not shown, but generally it should include a simple loop that reads an incoming message from the mailbox and processes it.

Source code: The code in Example 4-41 is included in the additional material for this book. See “PPE event handler” on page 621 for more information.

Example 4-41 Event handler on the PPU

```c
#include files....
#include <com_print.h> // the code from Example 4-40 on page 202

#define NUM_EVENTS 1
#define NUM_MBX 30

// take ‘spu_data_t’ structure and ‘spu_pthread’ function from
// Example 4-5 on page 90

int main()
{
    int i, ret, num_events, cnt;
    spe_event_handler_ptr_t event_hand;
    spe_event_unit_t event_uni, pend_events[NUM_EVENTS];
    uint32_t mbx=1;
    data.argp = NULL;

    // SPE_EVENTS_ENABLE flag should be set when creating SPE thread
    // to enable events tracing
    if ((data.spe_ctx = spe_context_create(SPE_EVENTS_ENABLE, NULL))
        == NULL)
    {
        perror("Failed creating context"); exit(1);
    }

    // create and register handle event handler
    event_hand = spe_event_handler_create();
    event_uni.events = SPE_EVENT_IN_MBOX;
    event_uni.spe = data.spe_ctx;
    ret = spe_event_handler_register(event_hand, &event_uni);

    // more types of events may be registered here

    // load the program to the local stores, and run the SPE threads.
    if (!program = spe_image_open("spu/spu")) {
```

37 See note 7 on page 84.
perror("Fail opening image"); exit(1);
}

if (spe_program_load(data.spe_ctx, program)) {
    perror("Failed loading program"); exit(1);
}

if (pthread_create(&data.pthread, NULL, &spu_pthread, &data)) {
    perror("Failed creating thread"); exit(1);
}

// write 4 first messages to make the mailbox queue full
for (mbx=1; mbx<5; mbx++){
    prn_p_mbx_m2s(mbx,0,mbx);
    spe_in_mbox_write(data.spe_ctx, &mbx,1,SPE_MBOX_ANY_BLOCKING);
}

// loop on all pending events
for ( ; mbx<NUM_MBX; ) {
    // wait for events to be set
    num_events =spe_event_wait(event_hand,pend_events,NUM_EVENTS,-1);

    // few events were set - handle them
    for (i = 0; i < num_events; i++) {
        if (pend_events[i].events & SPE_EVENT_IN_MBOX){

            // SPE read from mailbox- write to mailbox till it is full
            for (cnt=spe_in_mbox_status(pend_events[i].spe);cnt>0;
                    cnt--){
                mbx++;
                prn_p_mbx_m2s(mbx,0,mbx);
                ret = spe_in_mbox_write(pend_events[i].spe, &mbx,1,
                                         SPE_MBOX_ANY_BLOCKING);
            }
        }
    }

    //if we register more types of events- we can handle them here
}

// wait for all the SPE pthread to complete
if (pthread_join(data.pthread, NULL)) {
    perror("Failed joining thread"); exit (1);
}
4.4.4 Atomic unit and atomic cache

In this section, we explain how to implement a fast shared data structure for interprocessor communication by using the atomic unit and atomic cache hardware mechanism.

We include the following topics:

- In “Overview of the atomic unit and atomic cache” on page 211, we provide an overview of the atomic unit and atomic cache mechanism.

- In “Programming interface for accessing atomic unit and cache” on page 212, we describe the main software interfaces for an SPU or PPU program to use the atomic unit and atomic cache mechanism.

- In “Code example for using atomic unit and atomic cache” on page 214, we provide a code example for using the atomic unit and atomic cache.

Overview of the atomic unit and atomic cache

All of the atomic operations supported by the SPE are implemented by a specific atomic unit inside each MFC, which contains a dedicated local cache for cache-line reservations. This cache is called the atomic cache. The atomic cache has a total capacity of six 128-byte cache lines, of which four are dedicated to atomic operations.

When all the SPEs and the PPE perform atomic operations on a cache line with an identical effective address, a reservation for that cache line is present in at least one of the MFC units. When this occurs, the cache snooping and update processes are performed by transferring the cache line contents to the requesting SPE or PPE over the EIB, without requiring a read or write to the main system memory. Hardware support is essential for efficient atomic operations on shared data structures that consist of up to 512 bytes, divided into four 128-byte blocks mapped on a 128-byte aligned data structure in the local storage of the
SPEs. Such a system can be used as a fast broadcast interprocessor communication system.

The approach of exploiting this facility is to extend the principles behind the handling of a mutex lock or an atomic addition and ensure that the operations that are involved always affect the same four cache lines.

**Programming interface for accessing atomic unit and cache**

Two programming methods are available to exploit this functionality:

- The simplest method involves using the procedures `atomic_read` and `atomic_set` on both the SPU and PPU. These procedures provide access to individually shared 32-bit variables. They can be atomically set to specific values or atomically modified by simple arithmetic operations by using `atomic_add`, `atomic_inc`, and `atomic_dec`.

  The atomic procedures are part of a “sync” library that is delivered with SDK 3.0 and is implemented by using more basic reservation-related instructions. Refer to 4.5.3, “Using the sync library facilities” on page 238, in which we discuss this library further.

- The more powerful method allows multiple simultaneous atomic updates to a shared structure, using more complex update logic. The size of the shared variable can be up to four lines of a 128-byte atomic cache (assuming that no other mechanism uses this cache).

  To use this facility, the same sequence of operations that is required to handle a shared lock is performed by using atomic instructions as described as follows.

The following sequence of operations must be performed in the SPU program in order to set a lock on a shared variable:

1. Perform the reservation for the cache line designated to contain the part of the shared data structure that is to be updated by using `mfc_getllar`. This operation triggers the data transfer from the atomic unit that contains the most recent reservation or from the PPU cache to the requesting atomic unit of the SPE over the EIB.

2. The data structure mapped in the SPU local storage now contains the most up-to-date values. Therefore, the code can copy the values to a temporary buffer and update the structure with modified values according with the program logic.

3. Attempt the conditional update for the updated cache line by using `mfc_putllc`, and if unsuccessful, repeat the process from step 1.
4. Upon the successful update of the cache line, the program can continue to have both the previous structure values in the temporary buffer and the modified values in the local storage mapped structure.

The following sequence of operations must be performed in the PPU program in order to set a lock on a shared variable:

1. Perform the reservation for the cache line designated to contain the part of the shared data structure that is to be updated by using __lwarx or __ldarx. This operation triggers the data transfer from the atomic unit that contains the most recent reservation to the PPU cache over the EIB.

2. The data structure that was contained in the specified effective address, which resides in the PPU cache, now contains the most current values. Therefore, the code can copy the values to a temporary buffer and update the structure with modified values according with the program logic.

3. Attempt the conditional update for the updated cache line by using __stwcx or __stdcx, and if unsuccessful, repeat the process from step 1.

4. Upon the successful update of the cache line, the program can continue to have both the previous structure values contained in the temporary buffer and the modified values in the structure at the specified effective address.

A fundamental difference between the behavior of the PPE and SPE in managing atomic operations is worth noting. While both use the cache line size (128 bytes) as the reservation granularity, the PPU instructions operate on a maximum of 4 bytes (__lwarx and __stwcx) or 8 bytes (__ldarx and __stdcx) at once. The SPE atomic functions update the entire cache line contents.

For more details about how to use the atomic instructions on the SPE (mfc_getllar and mfc_putllc) and on the PPE (__lwarx, __ldarx, __stwcx, and __stdcx), see 4.5.2, “Atomic synchronization” on page 233.

The atomic cache in one of the MFC units or the PPE cache should always hold the desired cache lines before another SPE or the PPE requests a reservation on those lines. Provided that this occurs, the data refresh relies entirely on the internal data bus, which offers high performance.

The libsync synchronization primitives also use the cache line reservation facility in the SPE MFC. Therefore, special care is necessary to avoid conflicts that can occur when simultaneously exploiting manual usage of the atomic unit and other atomic operations provided by libsync.
Code example for using atomic unit and atomic cache

In this section, we provide a code example that shows how to use the atomic unit and atomic cache to communicate between the SPEs. The code example shows how to use the atomic instructions on the SPEs (mfc_getllar and mfc_putllc) to synchronize the access some shared structure.

Example 4-42 shows PPU code that initiates the shared structure, runs the SPE threads, and when the threads complete, reads the shared variable. No atomic access to this structure is done by the PPE.

Example 4-43 on page 215 show SPU code that uses the atomic instructions to synchronize the access to the shared variables between the SPEs.

Example 4-42   PPU code for using the atomic unit and atomic cache

```c
// add the ordinary SDK and C libraries header files...
// take ‘spu_data_t’ structure and ‘spu_pthread’ function from
// Example 4-5 on page 90

#define SPU_NUM 8

spu_data_t data[SPU_NUM];

typedef struct {
    int processingStep; // contains the overall workload processing step
    int exitSignal; // variable to signal end of processing step
    uint64_t accumulatedTime[8]; // contains workload dynamic execution
        // statistics (max. 8 SPE)
    int accumulatedSteps[8];
    char _dummyAlignment[24]; // dummy variables to set the structure
        // size equal to cache line (128 bytes)
} SharedData_s;

// Main memory version of the shared structure
// size of this structure is a single cache line
static volatile SharedData_s SharedData __attribute__ ((aligned(128)));

int main(int argc, char *argv[])
{
    int i;
    spe_program_handle_t *program;

    // Initialize the shared data structure
    SharedData.exitSignal = 0;
```
SharedData.processingStep = 0;

for (i = 0; i < SPU_NUM; ++i) {
    SharedData.accumulatedTime[i] = 0;
    SharedData.accumulatedSteps[i] = 0;
    data[i].argp = (void*)&SharedData;
    data[i].spu_id = (void*)i;
}

// ... Omitted section:
// creates SPE contexts, load the program to the local stores,
// run the SPE threads, and waits for SPE threads to complete.

// (the entire source code for this example is part of the book’s
// additional material).

// This subject of is also described in TBD_REF: Chapter 4.1.2 Task
parallelism and managing SPE threads

// Output the statistics
for (i = 0; i < SPU_NUM; ++i) {
    printf("SPE %d - Avg. processing time (decrementer steps):
           %lld\n", i, SharedData.accumulatedTime[i] / 
        SharedData.accumulatedSteps[i]);
}

return (0);

Example 4-43  SPU code for using the atomic unit and atomic cache

// add the ordinary SDK and C libraries header files...

Same ‘SharedData_s’ structure definition as in Example 4-42 on page 214

// local version of the shared structure
// size of this structure is a single cache line
static volatile SharedData_s SharedData __attribute__ ((aligned(128)));

// effective address of the shared structure
uint64_t SharedData_ea;
int main( uint64_t spuid, uint64_t argp, uint64_t envp )
{
    unsigned int status, t_start, t_spu;
    int exitFlag = 0, spuNum = envp, i;
    SharedData_ea = argp;

    // Initialize random number generator for fake workload example
    srand( spu_read_decrementer() );

    do{
        exitFlag = 0;

        // Start performance profile information collection
        spu_write_decrementer(0x7fffffff);
        t_start = spu_read_decrementer();

        // Data processing here
        // ...
        // Fake example workload:
        // 1) The first random number < 100 ends first step of the
        //    process
        // 2) The first number < 10 ends the second step of the process
        // Different SPEs process a different amount of data to generate
        // different execution time statistics.
        // The processingStep variable is shared, so all the SPEs will
        // process the same step until one encounters the desired result
        // Multiple SPEs can reach the desired result, but the first one
        // to reach it will trigger the advancement of processing step

        switch( SharedData.processingStep ){
        case 0:
            for( i = 0 ; i < (spuNum * 10) + 10 ; ++i ){
                if( rand() <= 100 ){ //found the first result
                    exitFlag = 1;
                    break;
                }
            }
            break;

        case 1:
            for( i = 0 ; i < (spuNum * 10) + 10 ; ++i ){
                if( rand() <= 10 ){ // found the second result
                    exitFlag = 1;
                    break;
                }
            }
            break;
        }
    }while( exitFlag != 1 );
}
break;
}
}
break;
// End performance profile information collection
t_spu = t_start - spu_read_decrementer();

// ... 
// Because we have statistics on all the SPEs average workload 
// time we can have some inter-SPE dynamic load balancing, 
// especially for workloads that operate in pipelined fashion 
// using multiple SPEs 

do{
    // get and lock the cache line of the shared structure
    mfc_getllar((void*)&SharedData, SharedData_ea, 0, 0);
    (void)mfc_read_atomic_status();

    // Update shared structure
    SharedData.accumulatedTime[spuNum] += (uint64_t) t_spu;
    SharedData.accumulatedSteps[spuNum]++;

    if( exitFlag ){
        SharedData.processingStep++;
        if(SharedData.processingStep > 1)
            SharedData.exitSignal = 1;
    }

    mfc_putllc((void*)&SharedData, SharedData_ea, 0, 0);
    status = mfc_read_atomic_status() & MFC_PUTLLC_STATUS;
} while (status);

}while (SharedData.exitSignal == 0);

return 0;
}
4.5 Shared storage synchronizing and data ordering

While the Cell/B.E. processor executes instructions in program order, it loads and stores data by using a “weakly” consistent storage model. With this storage model, storage accesses can be reordered dynamically. In doing so, there is an opportunity for improved overall performance and reduced effect on memory latency on instruction throughput.

By using this model, the programmer must explicitly order access to storage by using a special synchronization instruction, whenever storage occurs in the program order. If this is not done correctly, real-time bugs can result that are difficult to debug. For example, the program can run correctly on one system and fail on another, or run correctly on one execution and fail on another on the same system. Alternatively, over usage of the synchronization instructions can significantly reduce the performance because they take a lot of time to complete.

In this section, we discuss the Cell/B.E. storage model and the software utilities to control the data transfer ordering. For the reasons mentioned previously, it is important to understand this topic in order to obtain efficient and correct results. To learn more about this topic, see the “Shared-storage synchronization” chapter in the Cell Broadband Engine Programming Handbook.\(^{38}\)

We discuss the following topics:

- In 4.5.1, “Shared storage model” on page 221, we discuss the Cell/B.E. shared storage model and how the different components on the system can force ordering between the data transfers by using special ordering instructions. This section contains the following subtopics:
  - “PPE ordering instructions” on page 221 explains how code that runs the PPU can order the PPE data transfers on the main storage with respect to all other elements, such as other SPEs, in the system.
  - “SPU ordering instructions” on page 223 explains how the code that runs the SPU can order SPU data access to the LS with respect to all other elements that can access it, such as the MFC and other elements (for example PPE and other SPEs) in the system that access the LS through the MFC. It also explains how to synchronize access to the MFC channels.
  - “MFC ordering mechanisms” on page 227 explains the MFC ordering mechanism. The instructions are similar to the PPU ordering instructions, but from the SPU side. They enable the SPU code to order SPE data transfers on the main storage (done by the MFC) with respect to all other elements (such as PPE and other SPEs) in the system.

\(^{38}\) See note 1 on page 78.
In 4.5.2, “Atomic synchronization” on page 233, we explain the instructions that enable the different components on the Cell/B.E. chip to synchronize atomic access to some shared data structures.

In 4.5.3, “Using the sync library facilities” on page 238, we describe the sync library, which provides more high-level synchronization functions (based on the instructions mentioned previously). The supported C functions closely match those found in current traditional operating systems such as mutex, atomic increment, and decrements of variables and conditional variables.

In 4.5.4, “Practical examples of using ordering and synchronization mechanisms” on page 240, we describe specific real-life scenarios for using the ordering and synchronization instructions in the previous sections.

Table 4-12 on page 220 summarizes the effects of the different ordering and synchronization instructions on three storage domains: main storage, local storage, and channel interface. It shows the effects of instructions that are issued by different components: the PPU code, the SPU code, and the MFC. Regarding the MFC, data transfers are executed by the MFC following commands that were issued to the MFC by either the SPU code (using the channel interface) or PPU code (using the MMIO interface).

**Shaded table cells**: The gray shading in a table cell indicates that the instruction, command, or facility has no effect on the referenced domain.
| Issuer | Instruction, command, or facility | Main-storage domain | LS domain<br>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Accesses by PPE</td>
<td>Accesses by all other processor elements and devices</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Issuing thread</td>
<td>Both threads</td>
</tr>
<tr>
<td>PPU</td>
<td>sync$^c$</td>
<td>All accesses</td>
<td></td>
</tr>
<tr>
<td></td>
<td>lwsync$^e$</td>
<td>Accesses to memory-coherence-required locations</td>
<td></td>
</tr>
<tr>
<td></td>
<td>eieio</td>
<td>Accesses to caching-inhibited and guarded locations</td>
<td>Accesses to caching-inhibited and guarded locations</td>
</tr>
<tr>
<td></td>
<td>isync</td>
<td>Instruction fetches</td>
<td></td>
</tr>
<tr>
<td>SPU</td>
<td>sync</td>
<td>All accesses</td>
<td>All accesses</td>
</tr>
<tr>
<td></td>
<td>dsync</td>
<td></td>
<td>Load and store accesses</td>
</tr>
<tr>
<td></td>
<td>syncc</td>
<td></td>
<td>All accesses</td>
</tr>
<tr>
<td>MFC</td>
<td>mfcsync</td>
<td>All accesses</td>
<td>All accesses</td>
</tr>
<tr>
<td></td>
<td>mfceieio</td>
<td></td>
<td>Accesses to caching-inhibited and guarded locations</td>
</tr>
<tr>
<td></td>
<td>barrier</td>
<td></td>
<td>All accesses</td>
</tr>
<tr>
<td></td>
<td>&lt;f&gt;, &lt;b&gt;</td>
<td></td>
<td>All accesses for the tag group</td>
</tr>
<tr>
<td></td>
<td>MFC Multisource Synchronization Facility</td>
<td>All accesses</td>
<td>All accesses</td>
</tr>
</tbody>
</table>

a. The LS of the issuing SPE.
b. The channels of the issuing SPE.
c. The PowerPC sync instruction with L = 0.
4.5.1 Shared storage model

Unlike the in-order execution of instructions in the Cell/B.E. system, the processor loads and stores data by using a weakly consistent storage model. This means that the sequence in which any of the following orders are executed might be different from each other:

- The order of any processor element (PPE or SPE) performing storage access
- The order in which the accesses are performed with respect to another processor element
- The order in which the accesses are performed in main storage

To ensure that access to the shared storage is performed in program order, the software must place memory-barrier instructions between storage accesses.

The term storage access refers to access to main storage that is caused by a load, a store, a DMA read, or a DMA write. There are two orders to consider:

- Order of instructions execution
  The Cell/B.E. processor is an in-order machine, which means that, from a programmer's point of view, the instructions are executed in the order specified by the program.
- Order shared-storage access
  The order in which shared-storage access is performed might be different from both the program order and the order in which the instructions that caused the access are executed.

PPE ordering instructions

PPU ordering instructions enable the code that runs the PPU to order the PPE data transfers on the main storage with respect to all other elements, such as other SPEs, in the system. The ordering of storage access and instruction execution can be explicitly controlled by the PPE program by using barrier instructions. These instructions can be used between storage-access instructions to define a memory barrier that divides the instructions into those that precede the barrier instruction and those that follow it.
PPE supported barrier instructions are defined as intrinsics in the `ppu_intrinsics.h` header file, so that the programmer can easily use them in any C code application. Table 4-13 describes the two types of such instruction, storage barriers and instruction barriers.

**Table 4-13  PPE barrier intrinsics**

<table>
<thead>
<tr>
<th>Intrinsic</th>
<th>Description</th>
<th>Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Storage barriers</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>__sync()</td>
<td>Known as the <em>heavyweight sync</em>, ensures that all instructions that precede the sync have completed before the sync instruction completes and that no subsequent instructions are initiated until after the sync instruction completes. This does not mean that the previous storage accesses have completed before the sync instruction completes.</td>
<td>Ensure that the results of all storage into a data structure, caused by storage instructions executed in a critical section of a program, are seen by other processor elements before the data structure is seen as unlocked.</td>
</tr>
<tr>
<td>__lwsync()</td>
<td>Also known as <em>lightweight sync</em>, creates the same barrier as the sync instruction for storage access that is memory coherence. Therefore, unlike the sync instruction, it orders only the PPE main storage access and has no effect on the main storage access of other processor elements.</td>
<td>Use when ordering is required only for coherent memory, because it executes faster than sync.</td>
</tr>
<tr>
<td>__eieio()</td>
<td>Means “enforce in-order execution of I/O.” All main storage access caused by instructions preceding <code>eieio</code> have completed, with respect to main storage, before any main storage access caused by instructions following the <code>eieio</code>. The <code>eieio</code> instruction does not order access with differing storage attributes, for example, if an <code>eieio</code> is placed between a caching-enabled store and a caching-inhibited.</td>
<td>Use when managing shared data structures, accessing memory-mapped I/O (such as the SPE MMIO interface), and preventing load or store combining.</td>
</tr>
<tr>
<td><strong>Instruction barriers</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>__isync()</td>
<td>Ensures that all PPE instructions preceding the isync are completed before <code>isync</code> is completed. Causes an issue stall and blocks all other instructions from both PPE threads until the <code>isync</code> instruction completes.</td>
<td>In conjunction with self-modifying PPU code, execute after an instruction is modified and before it is run. Can also be used during context switching when the MMU translation rules are being changed.</td>
</tr>
</tbody>
</table>
Table 4-14 summarizes the use of the storage barrier instructions for two common types of main-storage memory:

- **System memory**
  The coherence main memory of the system. The XDR main memory falls into this category, as does the local storage when it is accessed from the EIB (when from the other PPE or SPEs).

- **Device memory**
  Memory that is caching-inhibited and guarded. In a Cell/B.E. system, it is typical of memory-mapped I/O devices, such as the double data rate (DDR) that is attached to the south bridge. Mapping of local storage of the SPEs to the main storage is caching-inhibited but not guarded.

In Table 4-14, “Yes” (and “No”) mean that the instruction performs (or does not perform) a barrier function on the related storage sequence. “Rec.” (for “recommended”) means that the instruction is the preferred one. “Not rec.” means that the instruction will work but is not the preferred one. “Not req.” (for “not required”) and “No effect” mean the instruction has no effect.

<table>
<thead>
<tr>
<th>Storage-access instruction sequence</th>
<th>System memory</th>
<th>Device memory</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>sync</td>
<td>lwsync</td>
</tr>
<tr>
<td>load-barrier-load</td>
<td>Yes</td>
<td>Rec.</td>
</tr>
<tr>
<td>load-barrier-store</td>
<td>Yes</td>
<td>Rec.</td>
</tr>
<tr>
<td>store-barrier-load</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>store-barrier-store</td>
<td>Yes</td>
<td>Rec.</td>
</tr>
</tbody>
</table>

a. Two stores to caching-inhibited storage are performed in the order specified by the program, regardless of whether they are separated by a barrier instruction.

**SPU ordering instructions**

SPU ordering instructions enable the code that runs the SPU to order SPU data access to the LS with respect to all other elements that can access it, such as the MFC and other elements in the system that access the LS through the MFC (for example PPE, other SPEs). They also synchronize the access to the MFC channels.
An LS can experience asynchronous interaction from the following streams that access it:

- Instruction fetches by the local SPU
- Data loads and stores by the local SPU
- DMA transfers by the local MFC or the MFC of another SPE
- Loads and stores in the main-storage space by other processor elements

With regard to an SPU, the Cell/B.E. in-order execution model guarantees only that SPU instructions that access the LS of that SPU appear to be performed in program order with respect to that SPU. However, it is not necessarily with respect to external accesses to that LS or with respect to the instruction fetch of the SPU.

Therefore, from an architecture point of view, an SPE might write data to the LS and immediately generate an MFC put command that reads this data (and transfers it to the main storage). In this case, without synchronization instructions, it is not guaranteed that the MFC will read the latest data, since it is not guaranteed that the MFC reading the data is performed after the SPU that writes the data. From practical point of view, there is no need to add the synchronization command to guarantee this ordering. Executing the six commands for issuing the DMA always takes longer than executing the former write to the LS.

From a programmer’s point of view, in the absence of external LS writes, an SPU load from an address in its LS returns the data written by that most-recent store of the SPU to that address. This statement is not necessarily true for an instruction fetch from that address, which might not guarantee the return of that recent data. The statement that follows regarding instruction fetch effect occurs only in cases of self-modifying code.

Consider a case where the LS and MFC resources of some SPEs that are mapped to the system-storage address space are accessed by software running on the PPE or other SPEs. In this case, there is no guarantee that two accesses to two different resources are ordered, unless a synchronization command, such as eieio or sync, is explicitly executed by the PPE or other SPEs, as explained in “PPE ordering instructions” on page 221.

In the following descriptions, we use the terms SPU load and SPU store to describe the access by the same SPU that executes the synchronization instruction. Several practical examples for using the SPU ordering instructions are discussed in the “Synchronization and ordering” chapter of the Synergistic Processor Unit Instruction Set Architecture Version 1.2 document.39 Specifically,
refer to the subtopic “External local storage access” in that chapter. It shows how
the instructions can be used when the processor, which is external to the SPE
(for example, the PPE), accesses the LS, for example, to write data to the LS and
later notifies the code that runs on the associated SPU that the writing of the data
is completed by writing to another address in this same LS.

The SPU instruction set provides three synchronization instructions. The easiest
way to use the instructions is through intrinsics, in which the programmer must
include the spu_internals.h header file. Table 4-15 briefly describes the
intrinsics and their main usage.

Table 4-15  

<table>
<thead>
<tr>
<th>Intrinsic</th>
<th>Description</th>
<th>Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>spu_sync</td>
<td>The sync (synchronize) instruction causes the SPU to wait until all pending</td>
<td>This instruction is most often used in conjunction with self-modifying SPU code. It must be used before attempting to execute new code that either arrives through the DMA transfers or is written with store instructions.</td>
</tr>
<tr>
<td></td>
<td>instructions of loads and stores to the LS and channel accesses have been</td>
<td></td>
</tr>
<tr>
<td></td>
<td>completed before fetching the next instruction.</td>
<td></td>
</tr>
<tr>
<td>spu_dsync</td>
<td>The dsync (synchronize data) instruction ensures that data has been stored</td>
<td>Architecturally, Cell/B.E. DMA transfers may interfere with store instructions and the store buffer. Therefore, the dsync instruction is meant to ensure that all DMA store buffers are flushed to the LS. That is, all previous stores to the LS will be seen by subsequent LS accesses. However, current cell implementation does not require the dsync instruction for doing so because it is handled by the hardware.</td>
</tr>
<tr>
<td></td>
<td>in the LS before the data becomes visible to the local MFC or other external</td>
<td></td>
</tr>
<tr>
<td></td>
<td>devices.</td>
<td></td>
</tr>
<tr>
<td>spu_sync_c</td>
<td>The syncc (synchronize channel) instruction ensures that channel synchronization</td>
<td>Ensures that the effects on SPU state that are caused by a previous write to a nonblocking channel are propagated and influence the execution of the instructions that follow.</td>
</tr>
<tr>
<td></td>
<td>is followed by the same synchronization provided by the sync instruction.</td>
<td></td>
</tr>
</tbody>
</table>
The instructions have both coherency and instruction-serializing effects, which are summarized in Table 4-16.

### Table 4-16 Effects of the SPU ordering instructions

<table>
<thead>
<tr>
<th>Intrinsic</th>
<th>Ensures these coherency effects</th>
<th>Forces these instruction serialization effects</th>
</tr>
</thead>
</table>
| spu_dsync   | ▶ Subsequent external reads access data written by prior SPU stores.  
▶ Subsequent SPU loads access data written by external writes. | ▶ Forces SPU load and SPU store access of LS due to instructions before the dsync to be completed before completion of dsync.  
▶ Forces read channel operations due to instructions before the dsync to be completed before completion of the dsync.  
▶ Forces SPU load and SPU store access of LS due to instructions after the dsync to occur after completion of the dsync.  
▶ Forces read-channel and write-channel operations due to instructions after the dsync to occur after completion of the dsync. |
| spu_sync    | ▶ The two effects of spu_dsync.  
▶ Subsequent instruction fetches access data written by prior SPU stores and external writes. | ▶ All access of LS and channels due to instructions before the sync to be completed before completion of sync.  
▶ All access of LS and channels due to instructions after the sync to occur after completion of the sync. |
| spu_sync_c  | ▶ The two effects of spu_dsync.  
▶ The second effect of spu_sync.  
▶ Subsequent instruction processing is influenced by all internal execution states modified by previous write instructions to a channel. | ▶ All access of LS and channels due to instructions before the sync to be completed before completion of sync.  
▶ All access of LS and channels due to instructions after the sync to occur after completion of the sync. |

Table 4-17 shows which SPU synchronization instructions are required between LS writes and LS reads to ensure that reads access data written by prior writes.

### Table 4-17 Synchronization instructions for access to an LS

<table>
<thead>
<tr>
<th>Writer</th>
<th>Reader</th>
<th>SPU instruction fetch</th>
<th>SPU load</th>
<th>External read&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>SPU store</td>
<td>sync</td>
<td>Nothing required</td>
<td>dsync</td>
<td></td>
</tr>
<tr>
<td>External write&lt;sup&gt;a&lt;/sup&gt;</td>
<td>sync</td>
<td>dsync</td>
<td>N/A</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup> By any DMA transfer (from the local MFC or a non-local MFC), the PPE, or other device, other than the SPU that executes the synchronization instruction.
MFC ordering mechanisms
The SPU can use the MFC channel interface to issue commands to the associated MFC. The PPU or other SPUs outside of this SPE similarly can use the MMIO interface of the MFC to send commands to a particular MFC. For each interface, the MFC independently accepts only queuable commands that are entered into one of the MFC SPU command queues (one queue for the channel interfaces and another for the MMIO). The MFC then processes these commands, possibly out of order to improve efficiency.

However, the MFC supports an ordering mechanism that can be activated through each of the following main interfaces:

- The Channel interface allows an SPU code to control the order in which the MFC executes the commands that were previously issued by this SPU by using the channel interface.

- Similarly but independently, the PPU or other SPUs can use the MMIO interface to control the order in which the MFC issues command that were previously queued on its MMIO interface.

  The effect of the commands, regardless of whether they are issued through either of the two interfaces, controls the order of the MFC data transfers on the main storage with respect to all other elements in the system, for example the PPE and other SPEs.

There are two types of ordering commands:

- Fence or barrier command options
  A tag specific mechanism is activated by appending a fence or barrier options to either data transfer or signal commands.

- Barrier commands
  A separate barrier command can be issued to order the command against all preceding and all succeeding commands in the queue, regardless of the tag group.

In the following sections, we further describe these two types of commands.

Fence or barrier command options
The fence or barrier command options ensure local ordering of storage accesses made through the MFC with respect to other devices in the system. The local ordering ensures the ordering of the MFC commands with respect to the particular MFC tag group (commands that have similar tag ID) and the command queue, that is the MFC proxy command queue and MFC SPU command queue. Both ordinary DMA and DMA list commands are supported as well as signaling commands.
Programmers can enforce ordering among DMA commands in a tag group with a fence or barrier option by appending an “f” for fence, or a “b” for barrier to the signaling commands, such as sndsig, or data transfer commands, such as getb and putf. The simplest way to do this is to use the supported MFC functions:

- The SPU code can use the functions call defined by the spu_mfcio.h header file. For example, use the mfc_getf and mfc_putb functions to issue the fenced get command and barrier put command respectively.

- The PPU code can use the functions call defined by the libspe2.h header file. For example, use the spe_mfcio_getf and spe_mfcio_putb functions to issue the fenced get command and the barrier put command respectively.

Table 4-18 lists the supported tag-specific ordering commands.

<table>
<thead>
<tr>
<th>Option</th>
<th>Commands</th>
</tr>
</thead>
<tbody>
<tr>
<td>Barrier</td>
<td>getb, getbs, getlb, putb, putbs, putrb, putrlb, sndsigb</td>
</tr>
<tr>
<td>Fence</td>
<td>getf, getfs, getlf, putf, putfs, putrf, putrlf, sndsigf</td>
</tr>
</tbody>
</table>

The fence and barrier options have different effects:

- The fence command is not executed until all previously issued commands within the same tag group have been performed. Commands that are issued after the fence command might be executed before the fence command.

- The barrier command and all the commands that are issued after the barrier command are not executed until all previously issued commands in the same tag group have been performed.

When the data transfers are issued, the storage system can complete the requests in an order that is different than the order in which they are issued, depending on the storage attributes.

Figure 4-4 on page 229 illustrates the different effects of the fence and barrier command. The row of white boxes represents command-execution slots, in real-time, in which the DMA commands (the solid red (darker) and green (lighter) boxes) might execute. Each DMA command is assumed to transfer the same amount of data. Therefore, all boxes are the same size. The arrows show how the DMA hardware, using out-of-order execution, might execute the DMA commands over time.
The commands are useful and efficient in synchronizing SPU code data access to the shared storage with access to other elements in the system. A common use of the command is in a double-buffering mechanism, as explained in “Double buffering” on page 160 and shown in the code in Example 4-30 on page 162. For more examples, see 4.5.4, “Practical examples of using ordering and synchronization mechanisms” on page 240.

**Barrier commands**

The barrier commands order storage access that is made through the MFC with respect to all other MFCs, processor elements, and other devices in the system. While the Cell Broadband Engine Architecture (CBEA) specifies those commands as having tag-specific effects (controls only the order in which transfers related to one tag-ID group are executed compare to each other), the current Cell/B.E. implementation has no tag-specific effects.

The commands can be activated only by the SPU that is associated with the MFC that uses the channel interface. There is no support from the MMIO interface. However, the PPU can achieve similar effects by using the non-MFC-specific ordering instructions that are described in “PPE ordering instructions” on page 221.
The easiest way to use the instructions from the SPU side is through intrinsics. In doing so, the programmer must include the `spu_mfcio.h` header file. Table 4-19 briefly describes these intrinsics and their main usage.

<table>
<thead>
<tr>
<th>Intrinsic</th>
<th>Description</th>
<th>Usage</th>
</tr>
</thead>
<tbody>
<tr>
<td>mfc_sync</td>
<td>The <code>mfcsync</code> command is similar to the PPE <code>sync</code> instruction and controls the order in which the MFC commands are executed with respect to storage access by all other elements and in the system.</td>
<td>This command is designed to use inter-processor or device synchronization. Since it creates a large load on the memory system, use it only between commands that involve storage with different storage attributes. Otherwise, use other synchronization commands.</td>
</tr>
<tr>
<td>mfc_eieio</td>
<td>The <code>mfceieio</code> command controls the order in which the DMA commands are executed with respect to the storage access by all other system elements, only when the storage that is accessed has the attributes of caching-inhibited and guarded (typical for I/O devices). The command is similar to the PPE <code>eieio</code> instruction. For details regarding the effects of accessing different types of memories, see Table 4-14 on page 223.</td>
<td>This command is used for managing shared data structures, performing memory-mapped I/O, and preventing load and store from combining in main storage. The fence and barrier options of other commands are preferred from performance point of view and should be used if they are sufficient.</td>
</tr>
<tr>
<td>mfc_barrier</td>
<td>The <code>barrier</code> command orders all subsequent MFC commands with respect to all MFC commands that precede the barrier command in the DMA command queue, independent of the tag groups. The barrier command will not complete until all preceding commands in the queue have completed. After the command completes, subsequent commands in the queue can be started.</td>
<td>This command is used for managing data structures that are in the main storage and are shared by other elements in the system.</td>
</tr>
</tbody>
</table>

**MFC multisource synchronization facility**

The Cell/B.E. processor contains multiple address and communication domains:- the main-storage domain, eight local LS-address domains, and eight local channel domains. The MFC multisource synchronization facility ensures the cumulative ordering of storage access that is performed by multiple sources, such as the PPE and SPEs, across all address domains. This is unlike the PPE `sync` instruction and other similar instructions that provide such cumulative ordering only with respect to the main-storage domain.
The MFC multisource synchronization facility addresses this cumulative-ordering need by providing two independent multisource synchronization-request methods:

- The **MMIO interface** allows the PPE or other processor elements or devices to control synchronization from the main-storage domain.

- The **Channel interface** allows an SPE to control synchronization from its LS-address and channel domain.

Each of these two synchronization-request methods ensures that all write transfers to the associated MFC are sent and received by the MFC before the MFC synchronization-request is completed. This facility does not ensure that read data is visible at the destination when the associated MFC is the source.

The two methods operate independently, so that synchronization requests through the MMIO register have no effect on synchronization requests through the channel, and vice versa.

**MMIO interface of the MFC multisource synchronization facility**

The MFC multisource synchronization facility can be accessed from the main storage domain by the PPE, other processor elements, or devices by using the MMIO MFC_MSSync (MFC multisource synchronization) register. A programmer accesses this facility through two functions that are defined in the libspe2.h header file and are further described in SPE Runtime Management Library document.40

Example 4-44 shows how the PPU programmer can achieve cumulative ordering by using the two correspondinglibspe2.h functions.

*Example 4-44  MMIO interface of MFC multisource synchronization facility*

```c
#include “libspe2.h”

// Do dome MFC DMA operation between memory and LS of some SPE
// PPE/other-SPEs use our MFC to transfer data between memory and LS

int status;

spe_context_ptr_t spe_ctx;

// init one or more SPE threads (also init ‘spe_ctx’ variable)

// Send a request to the MFC of some SPE to start tracking outstanding
// transfers which are sent to this MFC by either te associated SPU or
```

40 See note 7 on page 84.
// PPE or other-SPEs.

status = spe_mssync_start();

if (status==-1){
    // do whatever need to do on ERROR but do not continue to next step
}

// Check if all the transfers that are being tracked are completed.
// Repeat this step till the function returns 0 indicating the
// completions of those transfers

while(1){
    status = spe_mssync_status(spe_ctx); // nonblocking function

    if (status==0){
        break; // synchronization was completed
    }else{
        if (status==-1){
            // do whatever need to do on ERROR
            break; //unless we already exit program because of the error
        }
    }
}

Channel interface of the MFC multisource synchronization facility

MFC multisource synchronization facility can be accessed by the local SPU code from the LS domain by using the MFC_WrMSSyncReq (MFC write multisource synchronization request) channel. A programmer accesses this facility by using the two functions that are defined in the spu_mfcio.h header file and are further described in C/C++ Language Extensions for Cell Broadband Engine Architecture document.41 Example 4-45 on page 233 shows how the SPU programmer achieves cumulative ordering by using the two corresponding spu_mfcio.h functions.

---

41 See note 2 on page 80.
Example 4-45  Channel interface of MFC multisource synchronization facility

```c
#include "spu_mfcio.h"

uint32_t status;

// Do dome MFC DMA operation between memory and LS
// PPE/other-SPEs use our MFC to transfer data between memory and LS

// Send a request to the associated MFC to start tracking outstanding
// transfers which are sent to this MFC by either this SPU or PPE or
// other-SPEs
mfc_write_multi_src_sync_request();

// Check if all the transfers that are being tracked are completed.
// Repeat this step till the function returns 1 indicating the
// completions of those transfers

do{
    status = mfc_stat_multi_src_sync_request(); // nonblocking function
} while(!status);
```

An alternative is to use an asynchronous event that can be generated by the MFC to indicate the completion of the requested data transfer. Refer to the “MFC multisource synchronization facility” chapter in the *Cell Broadband Engine Programming Handbook*, which describes this alternative and other issues related to MFC multisource synchronization facility.\(^{42}\)

### 4.5.2 Atomic synchronization

The atomic operations that are supported by Cell/B.E. processor are implemented on both the SPE and the PPE. They enable the programmer to create synchronization primitives, such as semaphores and mutex locks, in order to synchronize storage access or other functions. Use these feature with special care to avoid livelocks and deadlocks.

The atomic operations that are implemented in the Cell/B.E. processor are not blocking, which enables the programmer to implement algorithms that are lock-free and wait-free.

\(^{42}\) See note 1 on page 78.
Atomic operations are described in detail in the “PPE atomic synchronization” chapter in the *Cell Broadband Engine Programming Handbook*. This same chapter contains usage examples and shows how the atomic operations can be used to implement synchronization primitives, such as mutex, atomic addition (part of semaphore implementation), and condition variables. We recommend that the advanced programmer reads this chapter to further understand this mechanism.

**Atomic synchronization instructions**

With the atomic mechanism, the programmer can set a lock, called a *reservation*, on an aligned unit of real storage, called a *reservation granule*, that contains the address that we want to lock. The Cell/B.E. reservation granule is 128 bytes and corresponds to the size of a PPE cache line, which means the programmer can set a lock on a block of 128 bytes that is also aligned to a 128-byte address.

The atomic synchronization mechanism includes the following instructions:

- **Load-and-reserve instructions** load the address value from memory and then set a reservation on the reservation granule that contains the address.

- **Store-conditional instructions** verify that the reservation is still set on the granule. Only if it is set, the store operation is carried out. Otherwise if the reservation does not exist, the instruction completes without altering the storage. By using the hardware set indication bit in the CRT register, the programmer can determine whether the store was successful.

The reservation is cleared by setting another reservation or by executing a conditional store to any address. Another processor element can also clear the reservation by accessing the same reservation granule.

A pair of load-and-reserve and store-conditional instructions permits an atomic update of a variable in main storage. This update enables the programmer to implement various synchronization primitives such as semaphore, mutex lock, test-and-set, and fetch-and-increment, and any atomic update of a single aligned word or double word in memory. The following sequence shows how this mechanism can implement a semaphore:

1. Reads a semaphore by using load-and-reserve.
2. Computes a result based on the value of the semaphore.
3. Uses store-conditional to write the new value back to the semaphore location only if that location has not been modified (for example, by another processor) since it was read in step 1.

---

43 ibid.
4. Determines if the store was successful:
   - If successful, the sequence of instructions from steps 1 to step 3 gives the appearance of having been executed atomically.
   - Otherwise, the other processor accesses the semaphore, so that the software can repeat this process (back to step 1).

These actions also control the order in which memory operations are completed, with respect to asynchronous events, and the order in which memory operations are seen by other processor elements or memory access mechanisms.

**PPE and SPE atomic implementation**

The atomic synchronization is implemented on both the PPE and the SPE:

- On the PPE side, atomic synchronization is implemented through a set of assembly instructions. A set of specific intrinsics makes the PPU instructions easily accessible from the C programming language because each of these intrinsics has a one-to-one assembly language mapping. The programmer must include the `ppu_intrinsics.h` header file to use them.
- On the SPE side, atomic synchronization is implemented on the SPE with a set of MFC synchronization commands that are accessible through a set of functions provided by the `spu_mfcio.h` file.

Table 4-20 summarizes both the PPE and SPE atomic instructions. For each PPE instruction that is attached, the MFC commands of an SPE implements a similar mechanism. For all PPE instructions, a reservation (lock) is set for the entire cache line in which this word resides.

---

### Table 4-20   Atomic primitives of PPE and SPE

<table>
<thead>
<tr>
<th>PPE</th>
<th>Description</th>
<th>SPE (MFC)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Load and reserve instructions</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>__ldarx</code></td>
<td>Loads a double word (cache line) and sets a reservation.</td>
<td><code>mfc_getllar</code></td>
<td>Transfers a cache line from the LS to the main storage and creates a reservation (lock). Is not tagged and is executed immediately (not queued behind other DMA commands).</td>
</tr>
<tr>
<td><code>__lwarx</code></td>
<td>Loads a word and sets reservation.(^a)</td>
<td><code>mfc_putllc</code></td>
<td>-</td>
</tr>
<tr>
<td><strong>Store conditional instructions</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>__stdcx</code></td>
<td>Stores a double word (cache line) only if a reservation (lock) exists.</td>
<td><code>mfc_putllc</code></td>
<td>Transfers a cache line from the LS to the main storage only if a reservation (lock) exists.</td>
</tr>
</tbody>
</table>
Two pairs of atomic instructions are implemented on both the PPE and SPE. The first pair is \texttt{__ldarx/_lwarx} and \texttt{mfc\_getllar}, and the second pair is \texttt{__stdcx/_stwcx} and \texttt{mfc\_putllc}, for the PPE and SPE respectively. These functions provide atomic read-modify-write operations that can be used to derive other synchronization primitives between a program that runs on the PPU and a program that runs on the SPU or SPUs. Example 4-46 on page 237 and Example 4-47 on page 237 show the PPU code and SPU code for implementing a mutex-lock mechanism using the two pairs of atomic instructions for PPE and SPE.

Refer to 4.5.3, “Using the sync library facilities” on page 238, which explains how the \textit{sync library} implements many of the standard synchronization mechanisms, such as mutex and semaphore, by using the atomic instructions. Example 4-46 is based on the sync library code from the \texttt{mutex\_lock.h} header file.

\textbf{Synchronization mechanisms:} The programmer should consider using the various synchronization mechanisms that are implemented in the sync library instead of explicitly using the atomic instructions that are described in this chapter. For more information, see 4.5.3, “Using the sync library facilities” on page 238.

Refer to the “PPE Atomic Synchronization” chapter in the \textit{Cell Broadband Engine Programming Handbook}, which provides more code examples that show how synchronization mechanisms can be implemented on both a PPE program and an SPE program to achieve synchronization between the two programs.\footnote{See note 1 on page 78.} Example 4-46 is based on one of those examples.
Example 4-46  PPE implementation of mutex_lock function in sync library

```c
#include "ppu_intrinsics.h"

// assumes 64 bit compilation of the code
void mutex_lock(uint64_t mutex_ptr) {

  uint32_t done = 0;

  do {
    if (__lwarx((void*)mutex_ptr) == 0)
      done = __stwcx((void*)mutex_ptr, (uint32_t)1);
  } while (done == 0); // retry if the reservation was lost

  __isync(); // synchronize with other data transfers
}
```

Example 4-47  SPE implementation of mutex_lock

```c
#include <spe_mfcio.h>
#include <spu_intrinsics.h>

void mutex_lock(uint64_t mutex_ptr) {
  uint32_t offset, status, mask;

  volatile char buf[256], *buf_ptr;
  volatile int *lock_ptr;

  // determine the offset to the mutex word within its cache line.
  // align the effective address to a cache line boundary.
  uint32_t mfc_ea2h(mutex_ptr) & 0x7F;
  uint32_t offset = mfc_ea2h(mutex_ptr) & ~0x7F;
  mutex_ptr = mfc_hl2ea(mfc_ea2h(mutex_ptr), mutex_lo);

  // cache line align the local stack buffer.
  buf_ptr = (char*)((uint32_t)(buf) + 127) & ~127;
  lock_ptr = (volatile int*)((buf_ptr + offset);

  // setup for use possible use of lock line reservation lost events.
  // detect and discard phantom events.
  mask = spu_read_event_mask();
  spu_write_event_mask(0);

```
if (spu_stat_event_status()) {
    spu_write_event_ack( spu_read_event_status());
}
spu_write_event_mask( MFC_LL_R_LOST_EVENT );

do{  // get-and-reservation the cache line containing mutex lock word.
    mfc_getllar( buf_ptr, mutex_ptr, 128, tag_id,0,0);
    mfc_read_atomic_status();

    if (*lock_ptr) {
        // The mutex is currently locked. Wait for the lock line
        // reservation lost event before checking again.
        spu_write_event_ack( spu_read_event_status());

        status = MFC_PUTLLC_STATUS;
    } else {
        // The mutex is not currently locked. Attempt to lock.
        *lock_ptr = 1;

        // put-conditionally, the cache line containing the lock word.
        mfc_putllc( buf_ptr, mutex_ptr, 128, tag_id,0,0);
        status = mfc_read_atomic_status() & MFC_PUTLLC_STATUS;
    }
} while (status);  // retry if the reservation was lost.

spu_write_event_mask(mask);  // restore the event mask

4.5.3 Using the sync library facilities

The sync library provides several simple general purpose synchronization constructs. The supported C functions closely match those found in current traditional operating systems.

Most of the functions in the sync library are supported by both the PPE and the SPE, but a small portion of them is supported only by the SPE. The functions are all based upon the Cell/B.E. load-and-reserve and store-conditional functionality that is described in 4.5.2, “Atomic synchronization” on page 233.
To use the facilities of the sync library, the programmer must consider the following files:

- The `libsync.h`: header file contains most of the definitions.
- The `libsync.a`: library contains the implementation and should be linked to the program.
- The function specific header files define each function. Therefore, the programmer should include this header file instead of the `libsync.h` file.

For example, the programmer can include the `mutex_init.h` header file when the `mutex_lock` operation is required. In this case, the programmer adds an underline when calling the function, for example, the `mutex_lock` function when including the `mutex_init.h` file instead of `mutex_lock` when including the `libsync.h` file.

**Note:** From a performance point of view, use the function specific header files because the functions are defined as `inline`, unlike the definition of the corresponding function in the `libsync.h` file. However, a similar effect can be achieved by setting the appropriate compilation flags.

The sync library provides five subclasses of synchronization primitives:

**Atomic operations**  Atomically add or subtract a value from a 32-bit integer variable.

**Mutexes**  Routines that operate on mutex (mutual exclusion) objects and are used to ensure exclusivity. Enables the programmer to atomically “lock” the mutex before accessing a shared structure and “unlock” it upon completion.

**Condition variables**  Routines that operate on condition variables and have two main operations. When a thread calls the `wait` operation on a condition, it is suspended and waits on that condition variable signal until another thread signals (or broadcasts) the condition variable by using the `signal` operation.

**Completion variables**  Enables one thread to notify other threads that are waiting on the completion variable that the completion is true.

**Reader/writer locks**  Routines that enable a thread to lock 32-bit word variables in memory by using two types of locks. A read lock is a non-exclusive mutex that allows multiple simultaneous readers. A writer lock is an exclusive mutex that allows a single writer.
4.5.4 Practical examples of using ordering and synchronization mechanisms

In this section, we include practical examples that show how to use the storage-ordering and synchronization facilities of the Cell/B.E. processor. Most of the examples are based on the “Shared-storage ordering” chapter in the Cell Broadband Engine Programming Handbook.45

SPE writing notifications to PPE using the fenced option

The most common use of the fenced DMA is when writing back notifications as shown in the following scenario. The following actions occur on the SPU program:

1. Compute the data.
2. Issue several DMA put commands to writes the computed data back to main storage.
3. Use the fenced putf command to write a notification that the data is available.
   This notification might be to any type memory, such as main memory, I/O memory, a signal-notification or MMIO register, or the mailbox of another SPE.
4. If the program will reuse the LS buffers, it waits for the completion of all the commands issued in previous steps.

The following actions occur on the PPU program:
1. Wait for the notification (poll the notification flag).
2. Operate on the computed SPE data.

   It is guaranteed that the updated data is available in memory since the SPE uses a fence between writing the computed data and the notification.

To ensure the ordering of the DMA writing of the data (step 2) and of the notification (step 3), the notification can be sent by using a fenced DMA command. This guarantees that the notification is not sent until all previous DMA commands of the group are issued.

In this example, the writing of both the data and the notification should have the same tag ID in order to guarantee that the fence will work.

45 See note 1 on page 78.
Ordering reads followed by writes using the barrier option

A barrier option is useful when a buffer read takes multiple commands that must be performed before writing the buffer, which also takes multiple commands as shown in the following scenario:

1. Issue several `get` commands to read data into the LS.
2. Issue a single barrier `putb` to write data to the main storage from the LS. The barrier guarantees that the `putb` command and the subsequent `put` commands issued in step 3 occur only after the `get` commands of step 1 are complete.
3. Issue a basic `put` command (without a barrier) to write the data to main storage.
4. Wait for the completion of all the commands issued in the previous steps.

By using the barrier form for the first command to write the buffer (step 2), the commands can be used to put the buffer (step 2 to 3) to be queued without waiting for the completion of the `get` commands (step 1). The programmer can take advantage of this mechanism to overlap the data transfers (read and write) with computation, allowing the hardware to manage the ordering. This scenario can occur on either the SPU or PPU that uses the MFC to initiate data transfers.

The `get` and `put` commands should have the same tag ID to guarantee that the barrier option that comes with the `get` and `put` commands ensures writing the buffer just after data is read. If the `get` and `put` commands are issued by using multiple tag IDs, then an MFC `barrier` command can be inserted between the `get` and `put` command instead of using a `put` command with a barrier option for the first `put` command.

If multiple commands are used to read and write the buffer, by using the barrier option, the read and write commands can be performed in any order, which provides better performance but forces all reads to finish before the writes start.

Double buffering by using the barrier option

The barrier commands are also useful when performing double-buffered DMA transfers in which the data buffers that are used for the input data are the same as the output data buffers. Example 4-48 illustrates such a scenario.

Example 4-48   Ordering SPU reads follows by writes using barrier-option

```c
int i;
i = 0;
‘get’ buffer 0
while (more buffers) {
    ‘getb’ buffer i^1 //‘mfc_getb’ function (with barrier)
```
wait for buffer i //‘mfc_write_tag_mask’ & ‘mfc_read_tag_status_all’
compute buffer i
‘put’ buffer i    //‘mfc_put’ function
i = i^1;
}
wait buffer i    //‘mfc_write_tag_mask’ & ‘mfc_read_tag_status_all’
compute buffer i
‘put’ buffer i    //‘mfc_put’ function

In the put command at the end of each loop iteration data is written from the same local buffer to which data is later read in the beginning of next iteration’s get command. Therefore, it is critical to use a barrier for the get command to ensure that the writes complete before the reads are started and to prevent the wrong data from being written. Example 4-30 on page 162 shows the code of SPU program that implements such a double-buffering mechanism.

PPE-to-SPE communication using storage barrier instruction

For some applications, the PPE is used as an application controller that manages and distributes work to the SPEs as shown in the following scenario. This scenario also shows how a sync storage barrier instruction can be used in this case to guarantee the correct ordering:

1. The PPE writes to the main storage with the data to be processed.
2. The PPE issues a sync storage barrier instruction.
3. The PPE notifies the SPE by writing to either the inbound mailbox or one of the signal-notification registers of the SPE.
4. The SPE reads the notification and understands that the data is ready.
5. The SPE reads the data and processes it.

To make this feasible, the data storage performed in step 1 must be visible to the SPE before receiving the work-request notification (steps 3 and 4). To ensure guaranteed ordering, a sync storage barrier instruction must be issued by the PPE between the final data store in memory and the PPE write to the SPE mailbox or signal-notification register. This barrier instruction is shown as step 2.
SPEs updating shared structures using atomic operation

In some cases, several SPEs can maintain a shared structure, for example when using the following programming model:

- A list of work elements in the memory defines the work that needs to be done. Each of the elements defines one task out of the overall work that can be executed parallel with the others.
- A shared structure contains the pointer to the next work element and potentially other shared information.
- An SPE that is available to execute the next work element atomically reads the shared structure to evaluate the pointer to the next work element and updates it to point to the next element. Then it can get the relevant information and process it.

Atomic operations are useful in such cases when several SPEs need to atomically read and update the value of the shared structure. Potentially, the PPE can also update this shared structure by using atomic instructions on the PPU program. Example 4-49 illustrates such a scenario and how the SPEs can manage the work and access to the shared structure by using the atomic operations.

Example 4-49  SPEs updating a shared structure by using atomic operation

```c
// local version of the shared structure
// size of this structure is a single cache line
static volatile vector shared_var ls_var __attribute__((aligned (128)));

// effective address of the shared sturture
uint64_t ea_var;

int main(unsigned long long spu_id, unsigned long long argv){
    unsigned int status;

    ea_var = get from PPE pointer to shared structure’s effective addr.

    while (1){
        do {
            // get and lock the cache line of the shared sahred structure
            mfc_getllar((void*)&ls_var, ea_var, 0, 0);
            (void)mfc_read_atomic_status();
        }
    }
}
```
if (valus in ‘ls_var’ indicate that the work was complete){

    (comment: ‘ls_var’ may contain total # of work tasks and #
     of complete task - SPE can compare those values)
    break;
}

// else - we have a new work to do

ls_var = progress the var to point to the next work to be done

mfc_putllc((void*)&ls_var, ea_var, 0, 0);

status = mfc_read_atomic_status() & MFC_PUTLLC_STATUS;
}
}

//get data of current work, process it, and put results in memory

-------------------

4.6 SPU programming

The eight SPEs are optimized for compute-intensive applications in which a
program’s data and instruction needs can be anticipated and transferred into the
local storage by DMA. Meanwhile, the SPE computes by using previously
transferred data and instructions. However, the SPEs are not optimized for
running programs that have significant branching, such as an operating system.

In this section, we discuss the following topics:

- In 4.6.1, “Architecture overview and its impact on programming” on page 245,
  we provide an overview on the main SPE architecture features and explain
  how they affect the SPU programming.

- In 4.6.2, “SPU instruction set and C/C++ language extensions (intrinsics)” on
  page 249, we provide an overview of the SPU instruction set and the SPU
  intrinsics, which are provide a simpler high-level programming interface to
  access the SPE hardware mechanisms and assembly instructions.

- In 4.6.3, “Compiler directives” on page 256, we describe the compiler
directive that is most likely to be used when writing an SPU program.
In 4.6.4, “SIMD programming” on page 258, we explain how the programmer can explicitly exploit the SIMD instructions to the SPU.

In 4.6.5, “Auto-SIMDizing by compiler” on page 270, we describe how the programmer can use compilers to automatically convert scalar code to SIMD code.

In 4.6.6, “Working with scalars and converting between different vector types” on page 277, we describe how to work with different vector data types and how to convert between vectors and scalars and vice versa.

In 4.6.7, “Code transfer using SPU code overlay” on page 283, we explain how the programmer can use the SDK 3.0 SPU code overlay face situations in which the code is too big to fit into the local storage.

In 4.6.8, “Eliminating and predicting branches” on page 284, we explain how write efficient code when branches are required.

This section mainly covers issues that are related to writing a program that runs efficiently on the SPU while fetching instructions form the attached LS. However, in most cases, an SPU program should also interact with the associated MFC to transfer data between the main storage and communicate with other processors on the Cell/B.E. chip. Therefore, it is important that you understand the issues that are explained in the previous sections of this chapter:

- Refer to 4.3, “Data transfer” on page 110, to understand how an SPU can transfer data between the LS and main storage.
- Refer to 4.4, “Inter-processor communication” on page 178, to understand how the SPU communicates with other processors on the chip (PPE and SPEs).
- Refer to 4.5, “Shared storage synchronizing and data ordering” on page 218, to understand how the data transfer of the SPU and other processors are ordered and who the SPU can synchronize with other processors.

### 4.6.1 Architecture overview and its impact on programming

In this section, we describe the main features of SPE architectures with an emphasis on the impact that those features have on programming of the SPU applications. This section is divided into subtopics that discuss specific components of the SPU architecture.

#### Memory and data access

In this section, we summarize the main features that are related to memory and data access.
**Local storage**

The local storage offers the following features:

- From a programmer's point of view, local storage is the storage domain that the program directly refers to when doing load and store instruction or use pointers.
- The size of the LS is 256 KB, which is relatively compact. The programmer should explicitly transfer data between main memory and the LS.
- The LS holds the instructions, stack, and data (global and local variables).
- The LS is accessed directly by load and store instructions, which are deterministic, have no address translation, and have low latency.
- The LS has a 16-bytes-per-cycle load and store bandwidth and is quadword aligned only. When the programmer stores data that is smaller than 16 bytes (for example scalar), the program performs a 16-byte read, shuffles to set the alignment, and modifies the data and 16-byte store, which is not efficient.

**Main storage and DMA**

The main storage and DMA support the following features:

- Access to the main storage is done by the programmer explicitly issuing a DMA transfer between the LS and main storage.
- The effective address of the main storage data should be supplied by the program that runs on the PPE.
- DMA transfer is done asynchronously with program execution, so that the programmer can overlap between data transfer and computation.
- Data of 16 bytes is transferred per cycle.

**Register file**

The register file offers the following features:

- A large register file supports 128 entries of 128-bits each.
- Unified register files, including all types (such as floating point, integers, vectors, pointers, and so on), are stored in the same registers.
- The large and unified files allow for instruction-latency hiding by using a deep pipeline without speculation.
- Big-endian data ordering is supported. The lowest-address byte and the lowest-numbered bit are the most-significant byte and bit, respectively.
**LS arbitration**
Arbitration to the LS is done according the following priorities (from high to low):

1. MMIO, DMA, and DMA list transfer element fetch
2. ECC scrub
3. SPU load/store and hint instruction prefetch
4. Inline instruction prefetch

**Instruction set and instruction execution**
In this section, we summarize the SPE instruction set and the way in which instructions are executed (pipeline and dual issue).

**Instructions set**
The following features are supported:

- The single-instruction, multiple-data (SIMD) instruction architecture that works on 128-bit vectors
- Scalar instructions
- SIMD instructions

The programmer should use the SIMD instructions as much as possible for performance reasons. They can be done by using functions that are defined by the SDK language extensions for C/C++ or by using the auto-vectorization feature of the compiler.

**Floating-point operations**
The instructions are executed as follows:

- Single-precision instructions are performed in 4-way SIMD fashion and are fully pipelined. Since the instructions have good performance, we recommend that the programmer use them as the application allows.

- Double-precision instructions are performed in 4-way SIMD fashion, are only partially pipelined, and stall dual issues of other instructions. The performance of these instructions makes the Cell/B.E. processor less attractive for applications that have massive use of double-precision instructions.

- The data format follows the IEEE standard 754 definition, but the single precision results are not fully compliant with this standard (different overflow and underflow behavior, support only for truncation rounding mode, different denormal results). The programmer should be aware that, in some cases, the computation results will not be identical to IEEE Standard 754
Branches
Branches are handled as follows:

➤ There is no branch prediction cache. Branches are assumed to be not taken. If a branch is taken, a stall occurs, which can have a negative effect on performance.

➤ Special branch hint commands can be used in the code to direct the hardware that a coming branch will be taken and, therefore, avoids the stall.

➤ There are no hint intrinsics. Instead programmers can improve branch prediction by using either the `__builtin_expect` compiler directive or the feedback directed optimization that is supported by the IBM XL compilers or FDPRPro.

Pipeline and dual issue
Pipeline and dual issue occur as follows:

➤ The SPE has two pipelines, named even (pipeline 0) and odd (pipeline 1). Whether an instruction goes to the even or odd pipeline depends on its instruction type.

➤ The SPE issues and completes up to two instructions per cycle, one in each pipeline.

➤ Dual issue occurs when a fetch group has two issuable instructions with no dependencies in which the first instruction can be executed on the even pipeline and the second instruction can be executed on the odd pipeline.

➤ Advanced programmers can write low-level code that fully uses the two pipelines by separating instructions that have data dependencies or that go to the same pipeline. That is, they can use an instruction that goes to the pipeline between them.

➤ In many cases, the programmer relies on the compiler or other performance tools (for example FDPRPro) to use the two pipelines. Therefore, we recommend doing analysis of the results either statically, for example by using the SPU static timing tool or Code Analyzer tool, or by using profiling, for example by loading the FDPRPro data into the Code Analyzer tool.
Features that are not supported by the SPU
The SPU does not support many of the features that are provided in most general purpose processors:

- There is no direct (SPU-program addressable) access to the main storage. The SPU accesses main storage only by using the DMA transfers of the MFC.
- There is no direct access to system control, such as page-table entries. The PPE privileged software provides the SPU with the address-translation information that its MFC needs.
- With respect to access by its SPU, the LS is unprotected and has untranslated storage.

4.6.2 SPU instruction set and C/C++ language extensions (intrinsics)

The SPU Instruction Set Architecture (ISA) is fully documented in the *Synergistic Processor Unit Instruction Set Architecture Version 1.2* document. The SPU ISA operates primarily on SIMD vector operands, both fixed-point and floating-point, with support for some scalar operands.

Another recommended source of information is the “SPU instruction set and intrinsics” chapter in the *Cell Broadband Engine Programming Handbook*, which provides a table of all the supported instructions and their latency.

The SDK provides a rich set of language extensions for C/C++ that define the SIMD data types and intrinsics that map to one or more assembly-language instructions into C-language functions. By using these extension, the programmer has convenient and productive control over code performance without needing to perform assembly-language programming.

From a programmer's point of view, we make the following recommendations:

- Use SIMD operations where possible because they provide the maximum performance, which can be up to four times for a single-precision float or 32-bit integers, or 16 times for 8-bit characters, faster than a scalar processor. Refer to 4.6.4, “SIMD programming” on page 258, in which we further discuss this topic.

- Use any scalar operations on the C/C++ code and the compiler maps them to one or more SIMD operations, such as read-modify-write, if the appropriate scalar assembly instruction does not exist. The programmer must try to minimize those operations, for example to use them only for control. Their performance is not as good as the SIMD ones and, in some cases, not as good as executing similar commands on an ordinary scalar machine.

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46 See note 39 on page 224.
47 See note 1 on page 78.
SPU Instruction Set Architecture
The SPU ISA operates primarily on SIMD 128-bit vector operands, both fixed-point and floating-point. The architecture supports some scalar operands.

The ISA has 204 instructions, which are grouped into several classes according to their functionality. Most of the instructions are mapped into either generic intrinsics or specific intrinsics that can be called as C functions from the program. For a full description of the instructions set, refer to the Synergistic Processor Unit Instruction Set Architecture Version 1.2 document.48

The ISA provides a reach set of SIMD operations that can be performed on 128-bit vectors of several fixed-point or floating-point elements. Instructions are available to access any of the MFC channels in order to initiate DMA transfers or to communicate with other processors.

In the following sections, we provide additional information about some of the main types of instructions.

Memory access SIMD operations
Load and store instructions are performed on the LS memory and uses 32 bits LS address. The instruction operates on 16 bytes elements which are quadword aligned. The SPU can perform a one such instruction in every cycle and their latency is about 6 cycles.

Channels access
A set of instructions is provided to access the MFC channels. The instructions can be used to initiate the DMA data transfer, communicate with other processors, access the SPE decrementer, and more. Refer to 4.6, “SPU programming” on page 244, in which we discuss the SPU interface with the MFC channel further.

SIMD operations
ISA SIMD instructions provide a reach set of operations (such as logical, arithmetical, casting, load and store, and so on) that can be performed on 128-bit vectors of either fixed-point or floating-point values. The vectors can contain various sizes of variables, such as 8, 16, 32 or 64 bits.

The performance of the program can be significantly effected by the way the SIMD instructions are used. For example, using SIMD instructions on 32-bit variables (single-precision floating point or 32-bit integer) can speed up the program by at least four times compared to the equivalent scalar program. In every cycle, the instruction works on four different elements in parallel, since there are four 32-bit variables for one 128 vector.

48 See 39 on page 224.
Figure 4-5 shows an example of the SPU SIMD add instruction of four 32-bit vector elements. This instruction simultaneously adds four pairs of floating-point vector elements, which are stored in registers VA and VB, and produces four floating-point results, which are written to register VC.

![Figure 4-5  SIMD add instruction](image)

**Scalar related instructions**

The ISA also provides instructions to access scalars. A set of store assist instructions is available to help store bytes, halfwords, words, and double words in the 128-bit vector registers. Similarly, instructions are provided to extract such scalars from the vector registers. Rotate instructions are also available and can be used to move data into the appropriate locations in the vector. A programmer can use the instructions whenever there is a need to operate on a specific element from a given vector, for example, to summarize the elements of one vector.

In addition, the instructions are often used by the compiler. Whenever the high level C/C++ function operates on scalars, the compiler translates it into a set of 16-byte read, modify, and write operations. In this process, the compiler uses the store assist and extract instruction to access the appropriate scalar element.

The ISA provides instructions that use or produce scalar operands or addresses. In this case, the values are set in the preferred slot in the 128-bit vector registers as illustrated in Figure 4-6 on page 252. The compiler can use the scalar store assist and extract instructions when a nonaligned scalar is used to shift it into the preferred slot.

To eliminate the need for such shift operations, the programmer can explicitly define the alignment of frequently used scalar variables, so that they will be in the preferred slot. The compiler optimization and after link optimization tools, such as
FDPRPro, that come with the SDK also try to help in this process by statically aligning scalar variables to the preferred slot.

**SIMD ‘cross-element’ shuffle instructions**

The ISA provides a set of shuffle instructions for reorganizing data in a given vector, which is very useful in SIMD programming. In one instruction, the programmer can reorder all the vector elements into an output vector. A less efficient alternative is to perform a series of several scalar-based instructions to extract the scalar from a vector and store it in the appropriate location in a vector.

Figure 4-7 shows an example instruction. Bytes are selected from vectors VA and VB based on the byte entries in control vector VC. The control vector entries are indices of bytes in the 32-byte concatenation of VA and VB. While the shuffle operation is purely byte oriented, it can also be applied to more than byte vectors, for example, vectors of floating points or 32-bit integers.
SPU C/C++ language extensions (intrinsics)
A large set of SPU C/C++ language extensions (intrinsics) makes the underlying
SPU ISA and hardware features conveniently available to C programmers:

- Intrinsics are essentially inline assembly-language instructions in the form of
  C-language function calls.
- Intrinsics can be used in place of assembly-language code when writing in the
  C or C++ languages.
- A single intrinsic maps one or more assembly-language instructions.
- Intrinsics provide the programmer with explicit control of the SPE SIMD
  instructions without directly managing registers.
- Intrinsics provide the programmer with access to all MFC channels as well as
  other system registers, for example the decremener and SPU state
  save/restore register.

For a full description of the extensions, see the C/C++ Language Extensions for
Cell Broadband Engine Architecture document.49

Directory of system header file: The SPU intrinsics are defined in the
spu_intrinsics.h system header file, which should be included if the
programmer wants to use them. The directory in which this file is located
varies depending on which compiler is used. When using GCC, the file is in
/usr/lib/gcc/spu/4.1.1/include/. When using GCC XLC, the file is in
/opt/ibmcmp/xlc/cbe/9.0/include/.

The SDK compiler supports these intrinsics to emit efficient code for the SPE
architecture, similar to using the original assembly instructions. The techniques
that are used by the compilers to generate efficient code include register
coloring, instruction scheduling (dual-issue optimization), loop unrolling and auto
vectorization, up-stream placement of branch hints, and more.

For example, an SPU compiler provides the intrinsic t=spu_add(a,b) as a
substitute for the assembly-language instruction fa rt,ra,rb. The compiler
generates a floating-point add instruction fa rt, ra, rb for the SPU intrinsic
t=spu_add(a,b), assuming that t , a, and b are vector float variables.

The PPU and SPU instruction sets have similar, but distinct, SIMD intrinsics. You
must understand the mapping between the PPU and SPU SIMD intrinsics when
developing applications on the PPE that will eventually be ported to the SPEs.
Refer to 4.1.1, “PPE architecture and PPU programming” on page 78, in which
we further discuss this issue.

49 See note 2 on page 80.
**Intrinsic data types**

Many of the intrinsics can accept parameters from different types, but the intrinsic name remains the same. For example, the `spu_add` function can add two `signed int` vectors into one output `signed int` vector, or add two float vectors (single precision) into one output float vector, and a few other types of vectors.

The translation from function to instruction depend on the data type of the arguments. For example, `spu_add(a,b)` can translate to a floating add or a signed int add depending on the input parameters.

Some operations cannot be performed on all data types. For example multiply using `spu_mul` can be performed only on floating point data types. Refer to the *C/C++ Language Extensions for Cell Broadband Engine Architecture* document, which provides detailed information about all the intrinsics, including the data type that is supported for each of them.\(^5\)

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**Recommendation:** The programmer must be familiar with this issue early in the development stage while defining the program’s data types. Doing so can prevent problems later during the development of the algorithm, in case a crucial operation is not supported on the chosen data types.

---

**Intrinsics classes**

SPU intrinsics are grouped into the following classes:

- **Specific intrinsics**

  These intrinsics have a one-to-one mapping with a single assembly-language instruction and are provided for all instructions except some branch and interrupt-related ones. All specific intrinsics are named by using the SPU assembly instruction that are prefixed by the string, `si_`. For example, the specific intrinsic that implements the `stop` assembly instruction is `si_stop`.

  Programmers rarely need these intrinsics since all of them are mapped into generic intrinsics, which are more convenient.

- **Generic and built-in intrinsics**

  These intrinsics map to one or more assembly-language instructions as a function of the type of input parameters and are often implemented as compiler built-ins. Intrinsics of this group are useful and cover almost all the assembly-language instructions including the SIMD ones. Instructions that are not covered are naturally accessible through the C/C++ language semantics.

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\(^5\) See note 2 on page 80.
All of the generic intrinsics are prefixed by the string `spu_`. For example, the intrinsic that implements the `stop` assembly instruction is named `spu_stop`.

- **Composite and MFC-related intrinsics**
  These convenience intrinsics are constructed from a sequence of specific or generic intrinsics. The intrinsics are further discussed in other sections in this chapter that discuss DMA data transfer and inter-processor communication using the MFC.

**Intrinsics: Functional types**
SPU generic intrinsics, which construct the main class of intrinsics, are grouped into the several types according to their functionality:

- **Constant formation (example: `spu_splats`)**
  Replicates a single scalar value across all elements of a vector of the same type.

- **Conversion (example: `spu_convtf`, `spu_convts`)**
  Converts from one type of vector to another. Using those intrinsics is the correct approach to do cast between two vectors of different types.

- **Scalar (example: `spu_insert`, `spu_extract`, `spu_promote`)**
  Allows programmers to efficiently coerce scalars to vectors, or vectors to scalars, enabling them to easily perform operations between vectors and scalars.

- **Shift and rotate (example: `spu_rlqbyte`, `spu_rlqw`)**
  Shifts and rotates the elements within a single vector.

- **Arithmetic (example: `spu_add`, `spu_madd`, `spu_nmadd`)**
  Performs arithmetic operations on all the elements of the given vectors.

- **Logical (example: `spu_and`, `spu_or`)**
  Performs a logical operation on an entire vector.

- **Byte operations (example: `spu_absd`, `spu_avg`)**
  Performs operations between bytes of the same vector.

- **Compare, branch and halt (example: `spu_cmpeq`, `spu_cmpgt`)**
  Performs different operations to control the flow of the program.

- **Bits and masks (example: `spu_shuffle`, `spu_sel`)**
  Perform a bitwise operation such as counting the number of bits equal 1 or the number of leading zeros.
Control (example: spu_stop, spu_ienable, spu_idisable)
Performs several control operations such as stopping and signaling the PPE and controlling interrupts.

Channel Control (example: spu_readch, spu_writech)
Reads from and writes to the MFC channels.

Synchronization and Ordering (example: spu_dsync)
Synchronizes and orders data transfer as related to external components.

In addition, the composite intrinsics contain intrinsics (spu_mfcdma32, spu_mfcdma64, spu_mfcstat) that enable the programmer to issue DMA commands to the MFC and check their status.

In the following section, we discuss some of the more useful intrinsics. For a list that summarizes all the SPU intrinsics, refer to Table 18 in the Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial document.51

4.6.3 Compiler directives

Like compiler intrinsics, compiler directives are crucial programming elements. In this section, we summarize the more important directives for SPU programming.

The aligned attribute
The aligned attribute is important in Cell/B.E. programming and is used to ensure proper alignment of variables in the program.

The aligned attribute is used in the following cases:

- To ensure proper alignment of the DMA source or destination buffer. A 16-byte alignment is mandatory for data transfer of more than 16 bytes, while a 128-byte alignment is optional but provides better performance.
- To ensure proper alignment of the scalar. Whenever a scalar is used, we recommend aligning it with the preferred slot to save shuffle operations while it is read or modified.

The aligned attribute for the SDK GCC and XLC implementations to align a variable into quadword (16 bytes) uses the following syntax:

```c
float factor __attribute__((aligned (16)));  
```

51 See note 3 on page 80.
The compilers currently do not support the alignment of automatic (stack) variables to an alignment that is stricter than the alignment of the stack itself (16 bytes).

**The volatile keyword**
The `volatile` keyword can be set when a variable is defined. It instructs the compiler that this variable can be changed for a reason that is not related to the program execution itself (that is, the program instructions). This prevents the compiler from doing optimizations that assume that the memory does not change unless a store instruction wrote new data to it. This type of scenario happens when a hardware component besides the processor itself can modify the variable.

For a Cell/B.E. program (either SPU or PPE), we recommended defining the buffers that are written by the DMA as volatile, such as a buffer on the LS to which a `get` command writes data. Defining the buffers ensures that buffers are not accessed by SPU load or store instructions until after DMA transfers have completed.

The `volatile` keyword for the SDK has the following syntax:

```c
volatile float factor;
```

**The `__builtin_expect` directive**
Since branch mispredicts are relatively expensive, the `__builtin_expect` directive provides a way for the programmer to direct branch prediction. The following example returns the result of evaluating `exp`, which means that the programmer expects `exp` to equal `value`:

```c
int __builtin_expect(int exp, int value)
```

The value can be a constant for a compile-time prediction or a variable used for a run-time prediction.

Refer to “Branch hint” on page 287, in which we further discuss use of this directive and provide useful code examples.

**The `_align_hint` directive**
The `_align_hint` directive helps compilers “auto-vectorize”. Although it looks like an intrinsic, it is more properly described as a compiler directive, since no code is generated as a result of using the directive. The following example informs the compiler that the pointer `ptr` points to data with a base alignment of `base`, with a byte offset from the base alignment of `offset`:

```c
_align_hint(ptr, base, offset)
```
The base alignment must be a power of two. Giving 0 as the base alignment implies that the pointer has no known alignment. The offset must be less than the base or zero. The \_align\_hint directive should not be used with pointers that are not naturally aligned.

**The restrict qualifier**

The restrict qualifier is well known in many C/C++ implementations and is part of the SPU language extension. When the restrict keyword is used to qualify a pointer, it specifies that all accesses to the object pointed to are done through the pointer, for example:

```c
void *memcpy(void * restrict s1, void * restrict s2, size_t n);
```

By specifying s1 and s2 as pointers that are restricted, the programmer is specifying that the source and destination objects (for the memory copy) do not overlap.

### 4.6.4 SIMD programming

In this section, we explain how to write SIMD operation-based programs to be run on the SPU. Specifically, we explain how the programmer can convert code that is based on scalar data types and operations to a code that is based on vector data types and SIMD operations.

We discuss the following topics:

- In “Vector data types” on page 259, we discuss the vector data types that are supported for SIMD operation.
- In “SIMD operations” on page 260, we discuss which SIMD operations are supported in the different libraries and how to use them.
- In “Loop unrolling for converting scalar data to SIMD data” on page 265, we discuss the main technique for converting scalar code to SIMD code by unrolling long loops.
- In “Data organization: AOS versus SOA” on page 267, we discuss the two main data organization methods for SIMD programming and explain how scalar code can be converted to SIMD by using the more common data organization method among the two (structure of arrays (SOA)).

An alternative to covert scalar code into SIMD code is to let the compiler perform automatically conversion of the code. This approach is called *auto-SIMDizing*, which we discuss further in 4.6.5, “Auto-SIMDizing by compiler” on page 270.
Vector data types
SPU SIMD programming operates on vector data types. Vector data types have the following main attributes:

- 128-bits (16-bytes) long
- Aligned on quadword (16-bytes) boundaries
- Different data types supported, including fixed point (for example char, short, int, signed or unsigned) and floating point (for example float and double)
- From 1 to 16 elements contained per vector depending on the corresponding type
- Are stored in memory similar to an array of the corresponding data types (For example, the vector of an integer is like an array of four 32-bit integers.)

To use the data types, the programmer must include the spu_intrinsics.h header file.

In general, the vector data types share a lot in common with ordinary C language scalar data types:

- Pointers to vector types can be defined as well as operations on those pointers. For example, if the pointer `vector float *p` is defined, then `p+1` points to the next vector (16-bytes) after that pointed to by `p`.
- Arrays of vectors can be defined as well as operations on those arrays. For example, if the array `vector float p[10]` is defined, then `p[3]` is the third variable in this array.

The vector data types can be used in two different formats:

- Full names, which are a combination of the data type of the elements that this vector consists of, together with vector prefix (for example `vector signed int`)
- Single token typedefs (for example `vec_int4`), which are more recommended since they are shorter and are compatible with using the same code for PPE SIMD programming
Table 4-21 summarizes the different data types that are supported by the SPU including both the full and the corresponding single token typedefs.

<table>
<thead>
<tr>
<th>Vector data type</th>
<th>Single-token typedef</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>vector unsigned char</td>
<td>vec_uchar16</td>
<td>Sixteen 8-bit unsigned characters</td>
</tr>
<tr>
<td>vector signed char</td>
<td>vec_char16</td>
<td>Sixteen 8-bit signed characters</td>
</tr>
<tr>
<td>vector unsigned short</td>
<td>vec_ushort8</td>
<td>Eight 16-bit unsigned halfwords</td>
</tr>
<tr>
<td>vector signed short</td>
<td>vec_short8</td>
<td>Eight 16-bit signed halfwords</td>
</tr>
<tr>
<td>vector unsigned int</td>
<td>vec_uint4</td>
<td>Four 32-bit unsigned words</td>
</tr>
<tr>
<td>vector signed int</td>
<td>vec_int4</td>
<td>Four 32-bit signed words</td>
</tr>
<tr>
<td>vector unsigned long long</td>
<td>vec_ullong2</td>
<td>Two 64-bit unsigned double words</td>
</tr>
<tr>
<td>vector signed long long</td>
<td>vec_llong2</td>
<td>Two 64-bit signed doublewords</td>
</tr>
<tr>
<td>vector float</td>
<td>vec_float4</td>
<td>Four 32-bit single-precision floats</td>
</tr>
<tr>
<td>vector double</td>
<td>vec_double2</td>
<td>Two 64-bit double precision floats</td>
</tr>
<tr>
<td>qword</td>
<td>-</td>
<td>Quadword (16-byte)</td>
</tr>
</tbody>
</table>

**SIMD operations**

In this section, we explain how the programmer can perform SIMD operations on an SPU program vector. There are four main options to perform SIMD operations as discussed in the following sections:

- In “SIMD arithmetic and logical operators” on page 261, we explain how SDK 3.0 compilers support a vector version of some of the common arithmetic and logical operators. The operators work on each element of the vector.
- In “SIMD low-level intrinsics” on page 261, we discuss the high level C functions that support almost all the SIMD assembler instructions of the SPU. The intrinsics contain basic logical and arithmetic operations between 128-bit vectors from different types and some operations between elements of a single vector.
- In “SIMD Math Library” on page 262, we explain how the library extends the low level intrinsic and provides functions that implement more complex mathematical operations, such as root square and trigonometric operations, on 128-bit vectors.
- In “MASS and MASSV libraries” on page 264, we explain the Mathematical Acceleration Subsystem (MASS) library functions, which are similar to those
of the SIMD Math Library, but are optimized to have better performance at the
price of redundant accuracy. The MASS vector (MASSV) library functions
perform similar operations on longer vectors that have a multiple of four.

**SIMD arithmetic and logical operators**

SDK compilers support a vector version of some of the common arithmetic and
logical operators. This is the easiest way to program SIMD operations because
the syntax is identical to programming with scalar variables. When the operators
are applied on vector variables, the compiler translates it to operators that work
separately on each element of the vectors.

While the compilers support basic arithmetic, logical, and rational operators, not
all the existing operators are currently supported. If the required operator is not
supported, the programmer should use the other alternatives that are described
in the following sections.

The following operators are supported:

- Vector subscripting: []
- Unary operators: ++, --, +, -, ~
- Binary operators: +, -, *, /, unary minus, %, &, |, ^, <<, >>
- Relational operators: ==, !=, <, >, <=, >=

For more details about this subject, see the “Operator overloading for vector data
types” chapter in the C/C++ Language Extensions for Cell Broadband Engine
Architecture document.52

Example 4-50 shows a simple code that uses some SIMD operators.

**Example 4-50  Simple SIMD operator code**

```c
#include <spu_intrinsics.h>

vector float vec1={8.0,8.0,8.0,8.0}, vec2={2.0,4.0,8.0,16.0};

vec1 = vec1 + vec2;
vec1 = -vec1;
```

**SIMD low-level intrinsics**

SDK 3.0 provides a reach set of low-level specific and generic intrinsics that
support the SIMD instructions that are supported by the SPU assembler
instruction set. For example, the c=spu_add(a,b) intrinsic means the add

---

52 See note 2 on page 80.
vc, va, vb instruction). The C-level functions are implemented either internally within the compiler or as macros.

The intrinsics are grouped into several types according to their functionality, as described in “Intrinsics: Functional types” on page 255. The following groups contain the most significant SIMD operations:

- Arithmetic intrinsics, which perform arithmetic operations on all the elements of the given vectors (for example spu_add, spu_madd, spu_nmadd, ...)
- Logical intrinsics, which perform logical operations on all the elements of the given vectors (spu_and, spu_or, ...)
- Byte operations, which perform operations between bytes of the same vector (for example spu_absd, spu_avg, ...)

The intrinsics support different data types. It is up to the compiler to translate the intrinsics to the correct assembly instruction depending on the type of the intrinsic operands.

To use the SIMD intrinsics, the programmer must include the spu_intrinsics.h header file.

Example 4-51 shows a simple code that uses low-level SIMD intrinsics.

```
Example 4-51 Simple SIMD intrinsics code

#include <spu_intrinsics.h>

vector float vec1={8.0,8.0,8.0,8.0}, vec2={2.0,4.0,8.0,16.0};

vec1 = spu_sub( (vector float)spu_splats((float)3.5), vec1);
vec1 = spu_mul( vec1, vec2);
```

**SIMD Math Library**

While SIMD intrinsics contain various basic mathematical functions that are implemented by corresponding SIMD assembly instructions, more complex mathematical functions are not supported by those intrinsics. The SIMD Math Library is provided with SDK 3.0 and addresses this issue by providing a set of functions that extend the SIMD intrinsics and support additional common mathematical functions. The library, like the SIMD intrinsics, operates on short 128-bit vectors from different types (for example single precision float, 32-bit integer) are supported. Which vector types are supported depends on the specific function.
The SIMD Math Library provides functions for the following categories:

- **Absolute value and sign functions**
  Remove or extract the signs from values

- **Classification and comparison functions**
  Return boolean values from comparison or classification of elements

- **Divide, multiply, modulus, remainder, and reciprocal functions**
  Standard arithmetic operations

- **Exponentiation, root, and logarithmic functions**
  Functions related to exponentiation or the inverse

- **Gamma and error functions**
  Probability functions

- **Minimum and maximum functions**
  Return the larger, smaller, or absolute difference between elements

- **Rounding and next functions**
  Convert floating point values to integers

- **Trigonometric functions**
  \( \sin, \cos, \tan \) and their inverses

- **Hyperbolic functions**
  \( \sinh, \cosh, \tanh \) and their inverses

The SIMD Math Library is an implementation of most of the C99 math library (-lm) that operates on short SIMD vectors. The library functions conform as closely as possible to the specifications set by the scalar standards. However, fundamental differences between scalar architectures and the CBEA require deviations, including the handling of rounding, error conditions, floating-point exceptions, and special operands such as NaN and infinities.

Two different versions of the SIMD Math Library are available:

- **The linkable library archive** is a static library that contains all the library functions. It is more convenient to use this version to code because it only requires the inclusion of a single header file. However, it produces slower and potentially larger binaries (depending on the frequency of invocation) due to the branching instructions necessary for function calls. The function calls also reduce the number of instructions that are available for scheduling and leveraging the large SPE register file.

- **The set of inline function headers** is a set of stand-alone inline functions. This version requires extra header files to include in each math function that is
used. However, it produces faster and smaller (unless inlined multiple times) binaries, because the compiler can reduce branching and often achieves better dual-issue rates and optimization. The function names are prefixed with an underscore character (_) compared to the linkable library format. For example, the inline version of fabsf4 is _fabsf4.

To use the SIMD Math Library, the programmer must perform the following actions:

- For the linkable library archive version, include the primary
  /usr/spu/include/simdmath.h header file.
- For the linkable library archive version, link the SPU application with the
  /usr/spu/lib/libsimdmath.a library.
- For the inline functions version, include a distinct header file for each function
  that is used. Those header files are in the /usr/spu/include/simdmath directory. For example, add #include <simdmath/fabsf4.h> to use the
  _fabsf4 inline function.

In addition, some classification functions require inclusion of the math.h file.

For additional information about this library, refer to the following resources:

- For code example and additional usage instruction, see Chapter 8, “Case
  study: Monte Carlo simulation” on page 499.
- For the function calls format, see the *SIMD Math Library API Reference*,
  SC33-8335-01:

  http://www-01.ibm.com/chips/techlib/techlib.nsf/techdocs/6DFAEFED17
  9041E8725724200782367

- For function specifications, refer to the “Performance Information for the
  MASS Libraries for CBE SPU” product documentation on the Web at the
  following address:

  A400&uid=swg27009548&loc=en_US&cs=UTF-8&lang=en&rss=ct2021other

**MASS and MASSV libraries**

In this section, we discuss two libraries that are part of SDK 3.0 and implement
various SIMD functions:

- The MASS library contains functions that operate on short 128-bit vectors.
  The interface of the functions is similar to the SIMD Math Library that is
  described in “SIMD Math Library” on page 262.

- The MASS vector (MASSV) library contains functions that can operate on
  longer vectors. The vector length can be any number that is a multiple of 4.
Similar to the SIMD Math Library, the MASS libraries can be used in two different versions, the linkable library archive version and the inline functions version.

The implementation of the MASS and MASSV libraries are different from SIMD Math Library in the following aspects:

- The SIMD Math Library is focused on accuracy while the MASS and MASSV libraries are focused on having better performance. For a performance comparison between the libraries, see the “Performance Information for the MASS Libraries for CBE SPU” document.
- The SIMD Math Library has support across the entire input domain, while the MASS and MASSV libraries can restrict the input domain.
- The MASS and MASSV libraries support a subset of the SIMD Math Library functions.
- The MASSV library can work on long vectors whose length is any number that is multiple of 4.
- The functions of the MASSV library have names that are similar to the SIMDmath and MASS functions, but with a prefix of “vs.”

To use the MASS library, the programmer must perform the following actions:

- For both libraries, include the `mass_simd.h` and `simdmath.h` header files in the `/usr/spu/include/` directory in order to use the MASS functions, and include `massv.h` header files for MASSV functions.
- For both versions, link the SPU application with the `libmass_simd.a` header file for MASS functions and with the `libmassv.a` file for MASSV functions. Both files are in the `/usr/spu/lib/` directory.
- For the inline functions version, include a distinct header file for each function that is used. The header files are in the `/usr/spu/include/mass` directory. For example, include the `acosf4.h` header file to use the `acosf4` inline function.

For more information about these libraries, including the function call format and description, as well as usage instructions, see the “MASS C/C++ function prototypes for CBE SPU” product documentation on the Web at the following address:


**Loop unrolling for converting scalar data to SIMD data**

In this section, we discuss the loop unrolling programming technique, which is one of the most common methods for SIMD programming.
The process of loop unrolling related to SIMD programming involves the following actions:

- The programmer expands a loop for each new iteration, which contains several formerly old iterations (before the unrolling).
- Few operations on scalar variables on the old iteration are joined to a single SIMD operation on a vector variable in the new iteration. The vector variable contains several of the original scalar variables.

The loop unrolling can provide significant performance improvement when applied on relatively long loops in an SPU program. The improvement is approximately the factor that is equal to the number of elements in an unrolled vector. For example, unrolling a loop that operated on a single-precision float provides four times the speed since four 32-bit floats exist in the 128-bit vector.

Example 4-52 shows the loop unrolling technique. The code contains two versions of multiply between two inout arrays of float. The first version is an ordinary scalar version (the `mult_` function) and the second is loop-unrolled SIMD version (the `vmult_` function).

**Source code:** The code in Example 4-52 is included in the additional material for this book. See “SPE loop unrolling” on page 621 for more information.

In this example, we require the arrays to be quadword aligned and the array length divisible by 4 (that is four float elements in a vector of floats).

Consider the following comments regarding the *alignment* and *length* of the vectors:

- We ensure that the quadword alignment uses the `aligned` attribute, which is recommended in most cases. If this is not the case, a scalar prefix can be added to the unrolled loop to handle the first unaligned elements.
- We recommend working with the arrays whose length is divisible by 4. If this is not the case, a suffix can be added to the unrolled loop to handle the last elements.

**Example 4-52  SIMD loop unrolling**

```c
#include <spu_intrinsics.h>
#define NN 100

// multiply - scalar version
void mult_(float *in1, float *in2, float *out, int N){
    int i;
```
for (i=0; i<N; i++){
    out[i] = in1[i] * in2[i];
}

// multiply - SIMD loop-unrolled version
// assume the arrays are quadword aligned and N is divisible by 4
void vmult_(float *in1, float *in2, float *out, int N){
    int i, Nv;
    Nv = N>>2; //divide by 4;

    vec_float4 *vin1 = (vec_float4*)in1, *vin2 = (vec_float4*)in2;
    vec_float4 *vout = (vec_float4*)out;

    for (i=0; i<Nv; i++){
        vout[i] = spu_mul( vin1[i], vin2[i]);
    }
}

int main( )
{
    float in1[NN] __attribute__((aligned (16)));
    float in2[NN] __attribute__((aligned (16)));
    float out[NN];
    float vout[NN] __attribute__((aligned (16)));

    // init in1 and in2 vectors

    // scalar multiply 'in1' and 'in2' into 'out' array
    mult_(in1, in2, out, (int)NN);

    // SIMD multiply 'in1' and 'in2' into 'vout' array
    vmult_(in1, in2, vout, (int)NN);

    return 0;
}

---

**Data organization: AOS versus SOA**

In this section, we discuss the two main data organization methods for SIMD programming. We also shows how scalar code can be converted to SIMD by using the more common data organization method among the two, which is structure of arrays.
Depending on the programmer's performance requirements and code size restraints, advantages can be gained by properly grouping data in a SIMD vector. We present the two main methods to organize the data.

The first method for organizing data in the SIMD vectors is called an *array of structures* (AOS) as demonstrated in Figure 4-8. This figure shows a natural way to use the SIMD vectors to store the homogenous data values (x, y, z, w) for the three vertices (a, b, c) of a triangle in a 3-D graphics application. This organization has the name array of structures because the data values for each vertex are organized in a single structure, and the set of all such structures (vertices) is an array.

![Figure 4-8  AOS organization](image)

The second method is a *structure of arrays* (SOA), which is shown in Figure 4-9. This figure shows the SOA organization to represent the x, y, z vertices for four triangles. The data types are the same across the vector, and now their data interpretation is the same. Each corresponding data value for each vertex is stored in a corresponding location in a set of vectors. This is different from the AOS method, where the four values of each vertex are stored in one vector.

![Figure 4-9  SOA organization](image)

The AOS data-packing approach often produces small code sizes, but it typically executes poorly and generally requires significant loop-unrolling to improve its efficiency. If the vertices contain fewer components than the SIMD vector can hold (for example, three components instead of four), SIMD efficiencies are compromised.
Alternatively, when using SOA, it is usually easy to perform loop unrolling or other SIMD programming. The programmer can think of the data as though it were scalar, and the vectors are populated with independent data across the vector. The structure of an unrolled loop iteration should be similar to the scalar case. However there is the one difference of replacing the scalar operations with identical vector operations simultaneously on a few elements that are gathered in one element.

Example 4-53 illustrates the process of taking a scalar loop in which the elements are stored using the AOS method and the equivalent unrolled SOA-based loop, which has fewer iterations by four times. The scalar and unrolled SOA loops are similar and use the same + operators. The only difference is how the indexing to the data structure is performed.

**Source code:** The code in Example 4-53 is included in the additional material for this book. See “SPE SOA loop unrolling” on page 621 for more information.

```c
#define NN 20

typedef struct{ // AOS data structure - stores one element
    float x;
    float y;
    float z;
    float w;
} vertices;

typedef struct{ // SOA structure - stores entire array
    vec_float4 x[NN];
    vec_float4 y[NN];
    vec_float4 z[NN];
    vec_float4 w[NN];
} vvertices;

int main(
{
    int i, Nv=NN>>2;
    vertices vers[NN*4];
    vvertices vvers __attribute__((aligned (16)));

    // init x, y, and z elements
```
// original scalar loop - work on AOS which is difficult to SIMDized
for (i=0; i<NN; i++){
    vers[i].w = vers[i].x + vers[i].y;
    vers[i].w = vers[i].w + vers[i].z;
}

// SOA unrolled SIMDized loop
for (i=0; i<Nv; i++){
    vvers.w[i] = vvers.x[i] + vvers.y[i];
    vvers.w[i] = vvers.w[i] + vvers.z[i];
}
return 0;

The subject of different SIMD data organization is further discussed in the “Converting scalar data to SIMD data” chapter in the Cell Broadband Engine Programming Handbook.53

4.6.5 Auto-SIMDizing by compiler

In this section, we discuss the auto-SIMDizing support of the Cell/B.E. GCC and XLC compilers. Auto-SIMDizing is the process in which a compiler automatically merges scalar data into a parallel-packed SIMD data structure. The compiler performs this process by first identifying parallel operations in the scalar code, such as loops. The compiler then generates SIMD versions of them, for example by automatically performing loop unrolling. During this process, the compiler performs all analysis and transformations that are necessary to fulfill alignment constraint.

From a programmer's point of view, in some cases, there is no need to perform explicit translation of scalar code to SIMD as explained in 4.6.4, “SIMD programming” on page 258. Instead, the programmer can write ordinary scalar code and instruct the compiler to perform auto-SIMDizing and translate the high level scalar code into SIMD data structures and SIMD assembly instructions.

However, at this point, there are limitations on the capabilities of the compilers in translating certain scalar code to SIMD. Not any scalar code that theoretically can be translated into a SIMD will eventually be translated by the compilers.

53 See note 1 on page 78.
Therefore, a programmer must have knowledge of the compiler limitations, so that the programmer can choose one of the following options:

- Write code in a way that is supported by the compilers for auto-SIMDizing.
- Recognize places in the code where auto-SIMDizing is not realistic and perform explicit SIMD programming in those places.

In addition, the programmer must monitor the compiler auto-SIMDizing results in order to verify in which places the auto-SIMDizing was successful and in which cases it failed. The programmer can perform an iterative process of compiling with auto-SIMDizing option enabled, debugging the places where auto-SIMDizing failed, re-writing the code in those cases, and then re-compiling it.

To activate compiler auto-SIMDization, the programmer must specify the optimization level depending on the compiler:

- When using XLC, use optimization level `-O3 -qhot` or higher.
- When using GCC, use optimization level `-O2 -ftree-vectorize` or higher.

We discuss the following topics:

- In “Coding for effective auto-SIMDization” on page 271, we explain how to write code that enables the compiler to perform effective auto-SIMDization and the limitations of the compilers in auto-SIMDizing other types of code.
- In “Debugging the compiler’s auto-SIMDization results” on page 273, we explain how the program can debug the compiler auto-SIMDization results in order to know whether it was successful. If it was not successful, the compiler provides information about the potential problems, so that the programmer can re-write the code.

**Coding for effective auto-SIMDization**

In this section, we explain how to write code that enables the compiler to perform effective auto-SIMDization. We also discuss the limitations of the compilers in the auto-SIMDization of scalar code. By knowing the limitations, the programmer can identify the places where auto-SIMDization is not possible. In such places, the programmer must explicitly translate to SIMD code.

**Organizing algorithms and loops**

The programmer must organize loops, so that they can be auto-SIMDized, and structure algorithms to reduce dependencies, by using the following guidance:

- The inner-most loops are the ones that can be SIMDized.
- Do not manually unroll the loops.
Use the `for` loop construct since it is the only one that can be auto-SIMDized. The `while` construct cannot be auto-SIMDized.

The number of iterations must have the following characteristics:
- A constant (preferred using `#define` directive) and not a variable
- More than three times the number of elements per vector

Shorter loops might also be SIMDizable, but it depends on their alignment and the number of statements in the loop.

Avoid the use of a `break` or `continue` statement inside a loop.

Avoid function calls within loops because they are not SIMDizable. Instead, use either inline functions or macros, or enable inlining by the compiler and possibly add an inlining directive to make sure that it happens. Another alternative is to distribute the function calls into a separate loop.

Avoid operations that do not easily map onto vector operations. In general, all operations, except branch, hint-for-branch, and load, are capable of being mapped.

Use the select operation for conditional branches within the loop. Loops that contain `if-then-else` statements might not always be SIMDizable. Therefore, use the C language colon question-mark (`:`) operator, which causes the compiler to SIMDize this section by using the select bits instruction.

Avoid aliasing problems, for example by using the `restrict` qualified pointers (illustrated in Example 4-56 on page 276). This qualifier when applied to a data pointer indicates that all access to this data is performed through this pointer and not through another pointer.

Loops with inherent dependences are not SIMDizable, as illustrated in Example 4-54 on page 274.

Keep the memory access-pattern simple:
- Do not use an array of structures, for example:
  ```c
  for (i=0; i<N; i++)  a[i].s = x;
  ```
- Use a constant increment. For example, do not use:
  ```c
  for (i=0; i<N; i+=incr)  a[i] = x;
  ```

**Organizing data in memory**

The programmer should lay out data in memory so that operations on it can be easily SIMDized by using the following guidance:

- Use stride-one accesses, which are memory access patterns in which each element in a list is accessed sequentially. Non-stride-one accesses are less efficiently SIMDized, if at all. Random or indirect accesses are not SIMDizable.
- Use arrays and not pointer arithmetic in the application to access large data structures.
- Use global arrays that are statically declared.
- Use global arrays that are aligned to 16-byte boundaries, for example using the `aligned` attribute. In general, lay out the data to maximize 16-byte aligned accesses.
- If there is more than a single misaligned store, distribute them into a separate loop. (Currently the vectorizer peels the loop to align a misaligned store.)
- If it is not possible to use aligned data, use the `alignx` directive to indicate to the compiler what the alignment is, for example:
  ```c
  #pragma alignx(16, p[i+n+1]);
  ```
- If it is known that arrays are disjoint, use the `disjoint` directive to indicate to the compiler that the arrays specified by the pragma are not overlapping, for example:
  ```c
  #pragma disjoint(*ptr_a, b)
  #pragma disjoint(*ptr_b, a)
  ```

**Mix of data types**
The mix of data types within code sections that can be potentially be SIMDized (that is, loops with many iterations) can present problems. While the compiler might succeed in SIMDizing those sections, the programmer must try to avoid such a mix of data types and keep a single data type within those sections.

**Scatter-gather**
Scatter-gather refers to a technique for operating on sparse data, using an index vector. A **gather operation** takes an index vector and loads the data that resides at a base address that is added to the offsets in the index vector. A **scatter operation** stores data back to memory, using the same index vector.

The Cell/B.E. processor’s SIMD architecture does not directly support scatter-gather in hardware. Therefore, the best way to extract SIMD parallelism is to combine operations on data in adjacent memory addresses into vector operations. This means that the programmer can use scatter-gather to bring the data into a continuous area in the local storage and then sequentially loop on the elements of this area variable. By doing so, the programmer can enable the compiler to SIMDize this loop.

**Debugging the compiler’s auto-SIMDization results**
The XLC compiler enables the programmer to debug the compiler’s auto-SIMDization results by using the `-qreport` option. The result is a list of high-level transformation that are performed by the compiler and that includes
everything from unrolling and loop interchange, to SIMD transformations. A transformed “pseudo source” is also presented.

Reports are generated for all loops that are considered for SIMDization. Specifically successful candidates are reported. In addition, if SIMDization was not possible, the reasons that prevented it are also reported.

This debugging feature enables the programmer to quickly identify opportunities for speedup. It provides feedback to the user explaining why loops are not vectorized. While those messages are not always trivial to understand, they help the programmer to rewrite the relevant sections to allow SIMD vectorization.

Similarly, the GCC compiler can provide debug information about the auto-SIMDizing process by using the following options:

- `-ftree-vectorizer-verbose=[X]`

  This option dumps information about which loops were vectorized, which were not, and why (X=1 least information, X=6 all information). The information is dumped to stderr unless the following flag is used.

- `-fdump-tree-vect`

  This option dumps information into `<C file name>.c.t##.vect`.

- `-fdump-tree-vect-details`

  This option is equivalent to setting the combination of the first two flags:
  `-fdump-tree-vect -ftree-vectorizer-verbose=6`

In this rest of this section, we show how to debug code that might not be SIMDized as well as other code that can be successfully SIMDized. We illustrate this by using the XLC debug features with the `-qreport` option enabled.

Example 4-54 shows an SPU code of a program named `t.c`, which is hard to SIMDize because of dependencies between sequential iterations.

**Example 4-54  A nonSIMDized loop**

```c
extern int *b, *c;

int main()
{
    for (int i=0; i<1024; ++i)
        b[i+1] = b[i+2] - c[i-1];
}
```
The code is then compiled with -qreport option enabled by using the following command:

```
spuxlc -c -qhot -qreport t.c”, in t.lst
```

Example 4-55 shows the t.lst file that is generated by the XLC compiler. It contains the problems in SIMDizing the loop and the transformed “pseudo source.”

**Example 4-55  Reporting of SIMDization problems**

```
1586-535 (I) Loop (loop index 1) at t.c <line 5> was not SIMD
vectorized because the aliasing-induced dependence prevents SIMD
vectorization.
1586-536 (I) Loop (loop index 1) at t.c <line 5> was not SIMD
vectorized because it contains memory references with non-vectorizable
alignment.
1586-536 (I) Loop (loop index 1) at t.c <line 6> was not SIMD
vectorized because it contains memory references ((char *)b +
(4)*($.CIV0 + 1))) with non-vectorizable alignment.
1586-543 (I) <SIMD info> Total number of the innermost loops considered<br/"1">. Total number of the innermost loops SIMD vectorized <"0">.

3 | long main()
    {
5 |    if (!1) goto lab_5;
   $.CIV0 = 0;
6 |    $.ICM.b0 = b;
   $.ICM.c1 = c;
5 |    do { /* id=1 guarded */ /* ~4 */
     /* region = 8 */
     /* bump-normalized */
6 |     $.ICM.b0[$.CIV0 + 1] = $.ICM.b0[$.CIV0 + 2] -
$.ICM.c1[$.CIV0 - 1];
5 |     $.CIV0 = $.CIV0 + 1;
7 | } while ((unsigned) $.CIV0 < 1024u); /* ~4 */
lab_5:
   rstr = 0;
```
The following messages are other examples of report problems with performing auto-SIMDization:

- Loop was not SIMD vectorized because it contains operation which is not suitable for SIMD vectorization.
- Loop was not SIMD vectorized because it contains function calls.
- Loop was not SIMD vectorized because it is not profitable to vectorize.
- Loop was not SIMD vectorized because it contains control flow.
- Loop was not SIMD vectorized because it contains unsupported vector data types.
- Loop was not SIMD vectorized because the floating point operation is not vectorizable under -qstrict.
- Loop was not SIMD vectorized because it contains volatile reference.

Example 4-56 shows an SPU code that is similar to Example 4-55 on page 275 but with correcting SIMD inhibitors.

*Example 4-56  A SIMDized loop*

```c
extern int * restrict b, * restrict c;

int main()
{
    // __alignx(16, c);     Not strictly required since compiler
    // __alignx(16, b);     inserts runtime alignment check

    for (int i=0; i<1024; ++i)
        b[i] = b[i] - c[i];
}
```
Example 4-57 shows the output t.lst file after compiling with the -qreport option enabled. The example reports successful auto-SIMDizing and the transformed “pseudo source.”

Example 4-57  Reporting of SIMDization problems

1586-542 (I) Loop (loop index 1 with nest-level 0 and iteration count 1024) at t.c <line 9> was SIMD vectorized.
1586-542 (I) Loop (loop index 2 with nest-level 0 and iteration count 1024) at t.c <line 9> was SIMD vectorized.
1586-543 (I) <SIMD info> Total number of the innermost loops considered <"2">. Total number of the innermost loops SIMD vectorized <"2">.
4 | long main()
   { 
       $.ICM.b0 = b;
       $.ICM.c1 = c;
       $.CSE2 = $.ICM.c1 - $.ICM.b0;
       $.CSE4 = $.CSE2 & 15;
       if (!(! $.CSE4)) goto lab_6;
   ...

4.6.6 Working with scalars and converting between different vector types

In this section, we explain how to convert between different types of vectors and how to work with scalars in the SIMD environment of the SPE. This section includes the following topics:

► In “Converting between different vector types” on page 277, we explain how to perform correct and efficient conversion between vectors of different types.

► In “Scalar overlay on SIMD instructions” on page 278, we discuss how to use scalars in SIMD instructions, include how to format them into a vector data type, and explain how to extract the results scalar from the vectors.

► In “Casting between vectors and scalar” on page 280, we describe how to cast vectors into an equivalent array of scalars and vice versa.

Converting between different vector types

Casts from one vector type to another vector type must be explicit and can be done using normal C-language casts. However, none of these casts performs any data conversion, and the bit pattern of the result is the same as the bit pattern of the argument that is cast.
Example 4-58 shows a casting approach between vectors that we do not recommend. This method usually does not provide the results that are expected by the programmer, because the integer variable \texttt{i\_vector} is assigned with a single precision float \texttt{f\_vector} variable which has different format. That is, the casting does not convert the bit pattern of the float to the integer format.

\hspace{1cm} \textit{Example 4-58 \ Method for casting between vectors - Not recommended}

\begin{verbatim}
// BAD programming example
vector float f_vector;
vector int i_vector;

i_vector = (vector int)f_vector;
\end{verbatim}

Instead, we recommend that the programmer perform casting between vectors by using special intrinsics that convert between different data types of vectors including modify the bit pattern to the required type. The casting involves the following conversion intrinsics:

\begin{itemize}
  \item \texttt{spu\_convtf} converts a signed or unsigned integer vector to a float vector.
  \item \texttt{spu\_convts} converts a float vector to a signed integer vector.
  \item \texttt{spu\_convtu} converts a float vector to an unsigned integer vector.
  \item \texttt{spu\_extend} extends an input vector to an output vector whose elements are two times larger than the input vector's elements. For example, the short vector is extended to the int vector, the float vector is extended to double, and so on.
\end{itemize}

\textbf{Scalar overlay on SIMD instructions}

The SPU loads and stores one quadword at a time. When instructions use or produce scalar (sub quadword) \texttt{i} operands (including addresses), the value is kept in the preferred scalar slot of a SIMD register. The fact that the scalar should be in the specific preferred slots requires extra instructions whenever a scalar is used as part of a SIMD instruction:

\begin{itemize}
  \item When a scalar is loaded in order to be a parameter of a SIMD instruction, it should be rotated to the preferred slot before being executed.
  \item When a scalar should be modified by a SIMD instruction, it should be loaded, rotated to the preferred slot, modified by the SIMD instruction, rotated back to its original alignment, and stored in memory.
\end{itemize}

Obviously these extra rotating instructions reduce performance, making vector operations on scalar data inefficient.
Making such scalar operations more efficient requires the following static technique:

1. Use the **aligned** attribute and extra padding if necessary to statically align the scalar to the preferred slot. Refer to “The aligned attribute” on page 256 regarding use of this attribute.

2. Change the scalars to quadword vectors. Doing so eliminates the three extra instructions that are associated with loading and storing scalars, which reduces the code size and execution time.

In addition, the programmer can use one of the SPU intrinsics to efficiently promote scalars to vectors, or vectors to scalars:

- `spu_insert` inserts a scalar into a specified vector element.
- `spu_promote` promotes a scalar to a vector that contains the scalar in the element that is specified by the input parameter. Other elements of the vector are undefined.
- `spu_extract` extracts a vector element from its vector and returns the element as a scalar.
- `spu_splats` replicate a single scalar value across all elements of a vector of the same type.

These instructions are efficient. By using them, the programmer can eliminate redundant loads and stores. One example for using the instructions is to cluster several scalars into vectors, load multiple scalars at one instruction using a quadword memory, and perform a SIMD operation that will operate on all the scalars at once.

There are two possible implementations for such a mechanism:

- Use extract or insert intrinsics. Cluster several scalars into a vector using `spu_insert` intrinsics, perform SIMD operations on them, and extract them back to their scalar shape using `spu_extract` intrinsic.

  Example 4-59 on page 280 shows an SPU program that implements this mechanism. Even this simple case is more efficient than multiplying the scalar vectors one by one using ordinary scalar operations. Obviously, if more SIMD operations are performed on the constructed vector, the performance overhead of creating the vector and extracting the scalars becomes negligible.

- Use the unions that perform casting between vectors and scalar arrays. See Example 4-60 on page 281 and Example 4-61 on page 282.
**Source code:** The code in Example 4-59 is included in the additional material for this book. See “SPE scalar-to-vector conversion using insert and extract intrinsics” on page 622 for more information.

**Example 4-59  Cluster scalars into vectors**

```c
#include <spu_intrinsics.h>
int main( )
{
    float a=10,b=20,c=30,d=40;
    vector float abcd;
    vector float efgh = {7.0,7.0,7.0,7.0};

    // initiate ‘abcd’ vector with the values of the scalars
    abcd = spu_insert(a, abcd, 0);
    abcd = spu_insert(b, abcd, 1);
    abcd = spu_insert(c, abcd, 2);
    abcd = spu_insert(d, abcd, 3);

    // SIMD multiply the vectors
    abcd = spu_mul(abcd, efgh);

    // do many other SIMD operations on ‘abcd’ and ‘efgh’ vectors

    // extract back the ‘multiplied’ scalar from the computed vector
    a = spu_extract(abcd, 0);
    b = spu_extract(abcd, 1);
    c = spu_extract(abcd, 2);
    d = spu_extract(abcd, 3);

    printf("a=%f, b=%f, c=%f, d=%f\n",a,b,c,d);
    return 0;
}
```

**Casting between vectors and scalar**

The SPU vector data types are kept in memory in a continuous 16-byte area whose address is also 16-bytes aligned. Pointers to vector types and non-vector types can, therefore, be cast back and forth to each other. For the purpose of aliasing, a vector type is treated as an array of its corresponding element type. For example, a vector float can be cast to float* and vice versa. If a pointer is cast to the address of a vector type, the programmer must ensure that the address is 16-byte aligned.
Casts between vector types and scalar types are *not allowed*. On the SPU, the `spu_extract`, `spu_insert`, and `spu_promote` generic intrinsics or the specific casting intrinsics can be used to efficiently achieve the same results.

In some cases, it is essential to perform SIMD computation on vectors but also perform computations between different elements of the same vector. A convenient programming approach is to define the casting unions of either the vectors or an array of scalars as explained in Example 4-60.

**Source code:** The code in Example 4-60 and Example 4-61 on page 282 is included in the additional material for this book. See “SPE scalar-to-vector conversion using unions” on page 622 for more information.

An SPU program that can use the casting unions is shown in Example 4-61 on page 282. The program uses the unions to perform a combination of SIMD operations on the entire vector and scalar operations between the vector elements.

While the scalar operations are easy to program, they are not efficient from a performance point of view. Therefore, the programmer must try to minimize the frequency in which they happen and use them only if there is not a simple SIMD solution.

**Example 4-60  Header file for casting between scalars and vectors**

```c
// vec_u.h file

#include <spu_intrinsics.h>

typedef union {
    vector signed char c_v;
    signed char c_s[16];

    vector unsigned char uc_v;
    unsigned char uc_s[16];

    vector signed short s_v;
    signed short s_s[8];

    vector unsigned short us_v;
    unsigned short us_s[8];

    vector signed int i_v;
    signed int i_s[4];
};
```

The code in Example 4-60 and Example 4-61 on page 282 is included in the additional material for this book. See “SPE scalar-to-vector conversion using unions” on page 622 for more information.
Example 4-61  Cluster scalars into vectors using a casting union

```c
#include <spu_intrinsics.h>
#include "vec_u.h"  // code from Example 4-60 on page 281 show

int main() {
  vec_float a __attribute__((aligned (16)));  
  vec_float b __attribute__((aligned (16)));  

  // do some SIMD operations on `a` and `b` vectors

  // perform some operations between scalar of specific vector
  a.s[0] = 10;
  a.s[1] = a.s[0] + 10;

  // initiate all `b` elements to be 7
  b.v = spu_splats( (float)7.0 );

  // SIMD multiply the two vectors
```
a.v = spu_mul(a.v, b.v);

// do many other different SIMD operations on ‘a’ and ‘b’ vectors

// extract back the scalar from the computed vector

printf("a0=%f, a1=%f, a2=%f, a3=%f\n", a.s[0], a.s[1], a.s[2], a.s[3]);

return 0;
}

4.6.7 Code transfer using SPU code overlay

In this section, we provide a brief overview on the SPU overlay facility that handles cases in which the entire SPU code is too big to fit the in the LS. We take into account that the 256 KB of the LS should also store the data, stack, and heap. The overlays can be used in other circumstances. For example, performance might be improved if the size of the data areas can be increased by moving rarely used functions to overlays.

An overlay is a program segment that is not loaded into an LS before the main program begins to execute, but is instead left in main storage until it is required. When the SPU program calls code in an overlay segment, this segment is transferred to the LS where it can be executed. This transfer usually overwrites another overlay segment that is not immediately required by the program.

The overlay feature is supported on SDK 3.0 for SPU programming but not for PPU programming.

The overlay is based on the following main principles:

- The linker that generates the overlays as two or more code segments can be mapped to the same physical address in the LS.
- The linker also generates call stubs and associated tables for overlay management. Instructions to call functions in overlay segments are replaced by branches to these call stubs.
- At execution time when a call is made from an executing segment to another segment, the system determines from the overlay tables whether the requested segment is already in the LS. If not, this segment is loaded dynamically by using a DMA command and can overlay another segment that was loaded previously.
XL compilers can assist in the construction of the overlays based upon the call graph of the application.

For a detailed description of this facility, including instructions on how to use it and a usage example, see the “SPU code overlays” chapter in the *Cell Broadband Engine Programming Handbook.*

### 4.6.8 Eliminating and predicting branches

The SPU hardware assumes the sequential instruction flow. This means that unless explicitly defined otherwise, all branches are not taken. Correctly predicted branches execute in one cycle, but a mispredicted branch (conditional or unconditional) incurs a penalty of 18 to 19 cycles, depending on the address of the branch target. Considering the typical SPU instruction latency of two to seven cycles, mispredicted branches can seriously degrade program performance. The branch instructions also restrict a compiler’s ability to optimally schedule instructions by creating a barrier on instruction reordering.

The most effective way to reduce the impact of branches is to eliminate them by using the first three methods as listed as follows. The second most effective method for reducing the impact of branches is to use a branch hint, which is presented last in the list:

- In “Function inlining”, we explain how this method defines the functions as inline and avoids the branch when a function is called and another branch when it is returned.
- In “Loop unrolling” on page 285, we explain how this method removes loops or reduces the number of iterations in a loop in order to reduce the number of branches (that appears at the end of the loop).
- In “Branchless control flow statement” on page 287, we explain how this method uses `spu_sel` intrinsics to replace a simple control statement.
- In “Branch hint” on page 287, we discuss the hint-for branch instructions. If the software speculates that the instruction branches to a target path, a branch hint is provided. If a hint is not provided, the software speculates that the branch is not taken. That is, the instruction execution continues sequentially.

**Function inlining**

The function-inlining technique can be used to increase the size of basic blocks (sequences of consecutive instructions without branches). This technique eliminates the two branches that are associated with function-call linkage, the branch for function-call entry and the branch indirect for function-call return.

---

54 See note 1 on page 78.
To use function inlining, the programmer can use either of the following techniques:

- Explicitly add the `inline` attribute to the declaration of any function that the programmer wants to inline. When recommended, one case is for functions that are short. Another case is for functions that have a small number of instances in the code but are often executed in run time, for example, when they appear inside a loop.

- Use the compiler options for automatic inlining of the appropriate functions. Table 4-22 lists such options of the GCC compiler.

Over-aggressive use of inlining can result in larger code, which reduces the LS space that is available for data storage or, in extreme cases, is too large to fit in the LS.

**Table 4-22  GCC options for functions inlining**

<table>
<thead>
<tr>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>-finline-small-functions</code></td>
<td>Integrates functions into their callers when their body is smaller than the expected function call code, so that the overall size of program gets smaller. The compiler heuristically decides which functions are simple enough to be worth integrating this way.</td>
</tr>
<tr>
<td><code>-finline-functions</code></td>
<td>Integrates all simple functions into their callers. The compiler heuristically decides which functions are simple enough to be worth integrating this way. If all calls to a given function are integrated, and the function is declared static, then the function is normally not output as assembler code in its own right.</td>
</tr>
<tr>
<td><code>-finline-functions-called-once</code></td>
<td>Considers all static functions that are called once for inlining into their caller even if they are not marked inline. If a call to a given function is integrated, then the function is not output as assembler code in its own right.</td>
</tr>
<tr>
<td><code>-finline-limit=n</code></td>
<td>By default, the GCC limits the size of the functions that can be inlined. This flag allows control of this limit for functions that are explicitly marked as inline.</td>
</tr>
</tbody>
</table>

**Loop unrolling**

Loop unrolling is another technique that can be used to increase the size of basic blocks (sequences of consecutive instructions without branches), which increases scheduling opportunities. It eliminates branches by decreasing the number of loop iterations.

If the number of loop iterations is a small constant, then we recommend removing the loop in order to eliminate branches in the code. Example 4-62 on page 286 provides a similar code example.
Example 4-62  Removing a short loop to eliminate branches

// original loop
for (i=0; i<3; i++) x[i]=y[i];

// can be removed and replaces by
x[0]=y[0];
x[1]=y[1];
x[2]=y[2];

If the number of loops is bigger, but the loop iterations are independent of each other, the programmer can reduce the number of loops and work on several items in each iteration as illustrated in Example 4-63. Another advantage of this technique is that it usually improves dual issue utilization. The loop-unrolling technique is often used when moving from scalar to vector instructions.

Example 4-63  Long loop unrolling for eliminating branches

// original loop
for (i=0; i<300; i++) x[i]=y[i];

// can be unrolled to
for (i=0; i<300; i+=3){
    x[i] =y[i];
    x[i+1]=y[i+1];
    x[i+2]=y[i+2];
}

Automatic loop unrolling can be performed by the compiler when the optimization level is high enough or when one of the appropriate options is set, for example -funroll-loops, -funroll-all-loops.

Typically, branches that are associated with the loop with a relatively large number of iterations are inexpensive because they are highly predictable. In this case, a non-predicted branch usually occurs only in the first and last iterations.

Similar to function inlining, over-aggressive use of loop unrolling can result in code that reduces the LS space available for data storage or, in an extreme case, is too large to fit in the LS.
Branchless control flow statement

The select-bits (selb) instruction is the key to eliminating branches for simple control flow statements such as if and if-then-else constructs. An if-then-else statement can be made branchless by computing the results of both the then and else clauses and by using select-bits intrinsics (spu_sel) to choose the result as a function of the conditional. If computing both results costs less than a mispredicted branch, then a performance improvement is expected.

Example 4-64 demonstrates the use of the spu_sel intrinsic to eliminate branches in a simple if-then-else control block.

**Example 4-64 Branchless if-then-else control block**

```c
// a,b,c,d are vectors

// original if-else control block
if (a>b) c +=1;
else    d = a+b;

// optimized spu_sel based code that eliminates branches but provides
// similar functionality.
select  = spu_cmpgt(a,b);
c_plus_1 = spu_add(c,1);
a_plus_b = spu_add(a,b);

c = spu_sel(c, c_plus_1, select);
d = spu_sel(a_plus_b, d, select);
```

Branch hint

The SPU supports branch prediction through a set of hint-for branch (HBR) instructions (hbr, hbra, and hbrr) and a branch-target buffer (BTB). These instructions support efficient branch processing by allowing programs to avoid the penalty of taken branches.

The hint-for branch instructions provide advanced knowledge about future branches such as the address of the branch target, the address of the actual branch instruction, and the prefetch schedule (when to initiate prefetching instructions at the branch target).

The hint-for branch instructions have no program-visible effects. They provide a hint to the SPU about a future branch instruction, with the intention that the information is to improve performance by prefetching the branch target.
If the software provides a branch hint, the software speculates that the instruction
branches to the branch target. If a hint is not provided, the software speculates
that the branch is not taken. If the speculation is incorrect, the speculated branch
is flushed and prefetched. It is possible to sequence multiple hints in advance of
multiple branches.

As with all programmer-provided hints, use care when using branch hints
because, if the information provided is incorrect, performance might degrade.
There are immediate and indirect forms for this instruction class. The location of
the branch is always specified by an immediate operand in the instruction.

A common use for a branch hint is in the end-of-loop branches when it is
expected to be correct. Such a hint is correct for all loop iterations besides the
last one.

A branching hint should be present soon enough in the code. A hint that
precedes the branch by at least eleven cycles plus four instruction pairs is
minimal. Hints that are too close to the branch do not affect the speculation after
the branch.

A common approach to generating static branch prediction is to use expert
knowledge that is obtained either by feedback-directed optimization techniques
or by using linguistic hints that are supplied by the programmer.

There are many arguments against profiling large bodies of code, but most SPU
code is not like that. SPU code tends to be well-understood loops. Thus,
obtaining realistic profile data should not be time consuming. Compilers should
be able to use this information to arrange code to increase the number of
fall-through branches (that is, conditional branches are not taken). The
information can also be used to select candidates for loop unrolling and other
optimizations that tend to unduly consume LS space.

Programmer-directed hints can also be used effectively to encourage compilers
to insert optimally predicted branches. Even though there is some anecdotal
evidence that programmers do not use them often, when they do use them, the
result is wrong. This is likely not the case for SPU programmers. SPU
programmers generally know a great deal about performance and are highly
motivated to generate optimal code.

The SPU C/C++ Language Extension specification defines a compiler directive
mechanism for branch prediction. The __builtin_expect directive allows
programmers to predicate conditional program statements. Example 4-65 on
page 289 demonstrates how a programmer can predict that a conditional
statement is false (that is, a is not larger than b).
Example 4-65  Predicting a false conditional statement

if(__builtin_expect((a>b),0))
    c += a;
else
    d += 1;

Not only can the __builtin_expect directive be used for static branch prediction, it can also be used for dynamic branch prediction. The return value of __builtin_expect is the value of the exp argument, which must be an integral expression. For dynamic prediction, the value argument can be either a compile-time constant or a variable. The __builtin_expect function assumes that exp equals a value. Example 4-66 shows code for a static prediction.

Example 4-66  Static branch prediction

if (__builtin_expect(x, 0)) {
    foo(); /* programmer doesn't expect foo to be called */
}

Example 4-67 shows a dynamic prediction.

Example 4-67  Dynamic branch prediction

cond2 = ... /* predict a value for cond1 */
...
cond1 = ...
if (__builtin_expect(cond1, cond2)) {
    foo();
}
cond2 = cond1; /* predict that next branch is the same as the previous*/

Compilers might require limiting the complexity of the expression argument because multiple branches can be generated. When this situation occurs, the compiler issue a warning message if the program’s branch expectations are ignored.

4.7 Frameworks and domain-specific libraries

In this section, we discuss the high-level frameworks for development and execution of parallel applications on the Cell/B.E. platform and domain-specific libraries that are provided by SDK 3.0.
The high-level frameworks provide an alternative to using the lower level libraries. The lower level libraries enable the programmer to have full control over the hardware mechanisms, such as DMA, mailbox, and SPE thread, and are discussed in other sections this chapter.

The two main purposes for the high level frameworks are to reduce the development time of programming an Cell/B.E. application and to create an abstract layer that hides from the programmer the specific features of the CBEA. In some cases, the performance of the application that uses those frameworks is similar to programming that uses the lower level libraries. Given the fact that development time is shorted and the code is more architecture independent, using the framework in those cases is preferred. However, in general, using the lower level libraries can provide better performance since the programmer can tune the program to the application-specific requirements.

We discuss the main frameworks that are provided with SDK 3.0 in the first two sections:

- In “Data Communication and Synchronization” on page 291, we discuss DaCS, which is an API and a library of C callable functions that provides communication and synchronization services among tasks of a parallel application running on a Cell/B.E. system. Another version of DaCS provides similar functionality for a hybrid system and is discussed in 7.2, “Hybrid Data Communication and Synchronization” on page 449.

- In “Accelerated Library Framework” on page 298, we discuss the ALF, which offers a framework for implementing the function off-load model on a Cell/B.E. system. It uses the PPE as the program control and SPEs as functions off-load accelerators. A hybrid version is also available and is discussed in 7.3, “Hybrid ALF” on page 463.

In addition to these SDK 3.0 frameworks, a growing number of high level frameworks for Cell/B.E. programming are being developed by companies other than IBM or by universities. Refer to 3.1.4, “The Cell/B.E. programming frameworks” on page 41, for a brief discussion of some of those frameworks.

The domain-specific libraries aim to assist Cell/B.E. programmers by providing reusable functions that implement a set of common algorithms and mathematical operators, such as Fast Fourier Transform (FFT), Monte Carlo, Basic Linear Algebra Subprograms (BLAS), matrix, and vector operators. We discuss these libraries in 4.7.3, “Domain-specific libraries” on page 314. In this section, we briefly describe some of the main libraries that are provided by SDK 3.0.

The functions that these libraries implement are optimized specifically for the Cell/B.E. processor and can reduce development time in cases where the developed application uses similar functions. In such cases, the programmer can use the corresponding library to implement those functions. Alternatively, they
can use the library code as a starting point and customize it to the specific requirements of the application. (The libraries are open source.)

4.7.1 Data Communication and Synchronization

DaCS is an API and a library of C callable functions that provides communication and synchronization services among tasks of a parallel application running either on a Cell/B.E. system or a hybrid system. Hybrid specific issues are discussed in 7.2, “Hybrid Data Communication and Synchronization” on page 449. In the rest of this discussion, the actual implementation, Cell/B.E. or hybrid, is of no importance because we describe only the concepts and the API calls.

DaCS can be used to implement various types of dialogs between parallel tasks by using common parallel programming mechanisms, such as message passing, mailboxes, mutex, and remote memory accesses. The only assumption is that there is a master task and slave tasks, a host element (HE), and accelerator elements (AE) in DaCS terminology. This is to be contrasted with MPI, which treats all tasks as equal. The goal of DaCS is to provide services for applications by using the host/accelerator model, where one task subcontracts lower-level tasks to perform a given piece of work. One model might be an application that is written by using MPI communication at the global level with each MPI task connected to accelerators that communicate with DaCS as illustrated in Figure 4-10 on page 292.
As shown in Figure 4-10, five MPI tasks exchange MPI messages and use DaCS communication with their accelerators. No direct communication occurs between the accelerators that report to a different MPI task.
DaCS also supports a hierarchy of accelerators. A task can be an accelerator for a task that is higher in the hierarchy. It can also be a host element for lower-level accelerators as shown in Figure 4-11.

As shown in Figure 4-11, the host element H0 is accelerated by two accelerators (A/H0, A/H1), which in turn are accelerated by four and three accelerators.

A DaCS program does not need to be an MPI program nor use a complex multi-level hierarchy. DaCS can be used for an application that consists of a single host process and its set of accelerators.

The main benefit of DaCS is to offer abstractions (message passing, mutex, and remote memory access) that are more common to application programmers than the DMA model, which is probably better known by system programmers. It also hides the fact that the host element and its accelerators might not be running on the same system.
**DaCS concepts**
To understand DaCS, it is important to understand the concepts as presented in the following sections.

**DaCS elements (HE/AE)**
A task in a DaCS program is identified by a couple, <DaCS Element, Process ID>. The DaCS Element (DE) identifies the accelerator or host element, which can be a PPE, an SPE, or another system in a hybrid configuration. In general, a given DE can have multiple processes running. Therefore, a Process ID is required to uniquely identify a participating DaCS task.

A DE can be either an HE, an AE, or both in the case of a multi-level hierarchy. An HE reserves an AE for its exclusive use and creates and terminates processes to run on it.

**DaCS groups**
In DaCS, the groups that are created by an HE and an AE can only join a group that was previously created by their HE. The groups are used to enable synchronization (barrier) between tasks.

**DaCS remote memory regions**
An HE can create an area of memory that is to be shared by its AE. A region can be shared in read-only or read-write mode. After the different parties are set up to share a remote memory segment, the data is accessed by each DE by using a put or get primitive to move data back and forth between its local memory and the shared region. The data movement primitives also support DMA lists to enable gather/scatter operations.

**DaCS mutex**
An HE can create a mutex that an AE agrees to share. After the sharing is explicitly set up, the AE can use lock-unlock primitives to serialize accesses to shared resources.

**DaCS communication**
Apart from the put and get primitives in remote memory regions, DaCS offers two other mechanisms for data transfer: mailboxes and message passing using basic send and recv functions. These functions are asymmetric like the data in mailboxes and send and recv functions between HE and AE only.

**DaCS wait identifiers**
The data movement functions in DaCS, the put and get primitives and the send and recv functions are asynchronous and return immediately. DaCS provides functions to wait for the completion of a previously issued data transfer request.
These functions use a wait identifier that must be explicitly reserved before being used. The functions should also be released when they are no longer in use.

**DaCS services**
The functions in the API are organized in groups as explained in the following sections.

**Resource and process management**
An HE uses functions in the resource and process management group to query the status and number of available AEs and to reserve them for future use. When an AE has been reserved, it can be assigned some work with the `dacs_de_start()` function. In a Cell/B.E. environment, the work given to an AE is an embedded SPE program, where in a hybrid environment, it is a Linux binary.

**Group management**
Groups are required to operate synchronizations between tasks. Currently, only barriers are implemented.

**Message passing**
Two primitives are provided for sending and receiving data by using a message passing model. The operations are nonblocking and the calling task must wait later for completion. The exchanges are point-to-point only, and one of the endpoints must be an HE.

**Mailboxes**
Mailboxes provide efficient message notification mechanism for small 32-bit data between two separate processes. In the case of the Cell/B.E. processor, mailboxes are implemented in the hardware by using an interrupt mechanism for communication between the SPE and the PPE or other devices.

**Remote memory operations**
Remote memory operations is a mechanism for writing or reading directly to or from memory in a remote processes address space. In MPI, this type of data movement is called *one-sided communication*.

**Synchronization**
Mutexes might be required to protect the remote memory operations and serialize access to shared resources.

**Common patterns**
The group, remote memory, and synchronization services are implemented with a consistent set of create, share/accept, and use and release/destroy primitives.
In each case, the HE is the initiator and the AE is invited to share what the HE has prepared for them as shown in Table 4-23.

**Table 4-23  Patterns for sharing resources in DaCS**

<table>
<thead>
<tr>
<th>HE side</th>
<th>AE side</th>
</tr>
</thead>
<tbody>
<tr>
<td>Creates the resource.</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Invites each AE to share the resource and wait for confirmation from each AE, one by one.</td>
<td>Accepts the invitation to share the resource.</td>
</tr>
<tr>
<td>Uses the resource. The HA can take part in the sharing, but it is not mandatory.</td>
<td>Uses the resource.</td>
</tr>
<tr>
<td>Destroys the shared resource. Waits for each AE to indicate that it does not use the resource anymore.</td>
<td>Releases the resource. Signals to the HE that the resource is no longer used.</td>
</tr>
</tbody>
</table>

For the remote memory regions, refer to Table 4-24.

**Table 4-24  Remote memory sharing primitives**

<table>
<thead>
<tr>
<th>HE side</th>
<th>AE side</th>
</tr>
</thead>
<tbody>
<tr>
<td>dacs_remote_mem_create()</td>
<td>Not applicable</td>
</tr>
<tr>
<td>dacs_remote_mem_share()</td>
<td>dacs_remote_mem_accept()</td>
</tr>
<tr>
<td>Not applicable</td>
<td>dacs_put(), dacs_get()</td>
</tr>
<tr>
<td>dacs_remote_mem_destroy()</td>
<td>dacs_remote_mem_release()</td>
</tr>
</tbody>
</table>

For the groups, refer to Table 4-25.

**Table 4-25  Group management primitives**

<table>
<thead>
<tr>
<th>HE side</th>
<th>AE side</th>
</tr>
</thead>
<tbody>
<tr>
<td>dacs_group_init()</td>
<td>Not applicable</td>
</tr>
<tr>
<td>dacs_group_add_member()</td>
<td>dacs_group_accept()</td>
</tr>
<tr>
<td>dacs_group_close()</td>
<td></td>
</tr>
<tr>
<td>This marks the end of the group creation.</td>
<td></td>
</tr>
<tr>
<td>Not applicable</td>
<td>dacs_barrier_wait()</td>
</tr>
<tr>
<td>dacs_group_destroy()</td>
<td>dacs_group_leave()</td>
</tr>
</tbody>
</table>
For mutexes, refer to Table 4-26.

**Table 4-26 Mutexes primitives**

<table>
<thead>
<tr>
<th>HE side</th>
<th>AE side</th>
</tr>
</thead>
<tbody>
<tr>
<td>dacs_mutex_init()</td>
<td>Not applicable</td>
</tr>
<tr>
<td>dacs_mutex_share()</td>
<td>dacs_mutex_accept()</td>
</tr>
<tr>
<td>Not applicable</td>
<td>dacs_mutex_lock()</td>
</tr>
<tr>
<td></td>
<td>dacs_mutex_unlock()</td>
</tr>
<tr>
<td></td>
<td>dacs_mutex_trylock()</td>
</tr>
<tr>
<td>dacs_mutex_destroy()</td>
<td>dacs_mutex_release()</td>
</tr>
</tbody>
</table>

**Annotated DaCS example**

We provide an example program that illustrates the use of most of the API functions. The program source code is well documented so that by reading through it, you will clearly understand the DaCS API functions and how they need to be paired between the HE and AE.

**Source code:** The DaCS code example is part of the additional material for this book. See “Data Communication and Synchronization programming example” on page 617 for more details.

The DaCS libraries are fully supported by the debugging and tracing infrastructure that is provided by the IBM SDK for Multicore Acceleration. The sample code can be built with the debug and trace types of the DaCS library.

**Usage notes and current limitations**

DaCS provides services for data communication and synchronization between an HE and its AE. It does not tie the application to a certain type of parallelism, and any parallel programming structure can be implemented.

In the current release, no message passing between AEs is allowed and complex exchanges either require more HE intervention or must be implemented by using the shared memory mechanisms (remote memory and mutexes). A useful extension allows AE-to-AE messages. Some data movement patterns (pipeline, ring of tasks) might be easier to implement in DaCS. However, we can always call libspe2 functions directly from within a DaCS task to implement custom task synchronizations and data communications, but this technique is not a supported by the SDK.
4.7.2 Accelerated Library Framework

The ALF offers a framework for implementing the function offload model. The functions that were previously running on the host are now offloaded and accelerated by one or more accelerators. In the ALF, the host and accelerator node types can be specified by the API and are general enough to allow various implementations. In the current implementations, the accelerator functions always run on the SPE of a Cell/B.E. system and the host applications run either on the PPE of a Cell/B.E. system on the officially supported cell of the ALF version or on an x86_64 node in the alpha version of the hybrid model.

ALF overview

With the ALF, the application developer is required to divide the application in two parts: the control part and the computational kernels. The control part runs on the host, which subcontracts accelerators to run the computational kernels. These kernels take their input data from the host memory and write back the output data to the host memory.

The ALF is an extension of the subroutine concept, with the difference that input arguments and output data have to move back and forth between the host memory and the accelerator memory, similar to the Remote Procedure Call (RPC) model. The input and output data might have to be further divided into blocks to be made small enough to fit the limited size of the accelerator’s memory. The individual blocks are organized in a queue and are meant to be independent of each other. The ALF run time manages the queue and balances the work between the accelerators. The application programmer only has to put the individual pieces of work in the queue.

Let us suppose that we have an MPI application that we want to accelerate by offloading the computational routines onto a multitude of Cell/B.E. SPEs. When accelerated by using the ALF, each MPI task still exists as an MPI task, working in sync with the other MPI tasks and performing the necessary message passing. But now, instead of running the computational parts between MPI calls, it only orchestrates the work of the accelerator tasks that it will allocate for its own use.

Each MPI task must know about the other MPI tasks for synchronization and message passing. However, an accelerator task is not required to know anything about its host task nor about its siblings nor about the accelerators running on behalf of foreign MPI tasks. An accelerator task has no visibility to the outside world. It only answers to requests. It is fed with input data, does some processing, and the output data it produces is sent back.

There is no need for an accelerator program to know about its host program because the ALF run time handles all the data movement between the accelerator memory and the host memory on behalf of the accelerator task. The
ALF run time does the data transfer by using various tricks, exploiting DMA, double or triple buffering and pipelining techniques that the programmer does not need to learn about. The programmer must only describe, generally at the host level, the layout of the input and output data in host memory that the accelerator task will work with.

The ALF gives a lot of flexibility to manage accelerator tasks. It supports the Multiple Program Multiple Data (MPMD) model in two ways:

- A subset of accelerator nodes can run task A providing the computational kernel ckA, while another subset runs task B providing the kernel ckB.
- A single accelerator task can perform only a single kernel at an one time. There are ways that the accelerator can load a different kernel after execution starts.

The ALF can also express dependencies between tasks by allowing for complex ordering of tasks when synchronization is required.

**The ALF run time and programmer’s tasks**

The ALF run time provides the following services from the application programmer’s perspective:

- At the host level:
  - Work blocks queue management
  - Load balancing between accelerators
  - Dependencies between tasks

- At the accelerator level:
  - Optimized data transfers between the accelerator memory and the host memory, exploiting the data transfer list used to describe the input and output data

On the host side, the application programmer must make calls to the ALF API to perform the following actions:

- Create the tasks
- Split the work into work blocks, that is describing the input and output data for each block
- Express the dependencies between tasks if needed
- Put the work blocks in the queue

On the accelerator side, the programmer must only write the computational kernels. As we see later, this is slightly over simplified because the separation of duties between the host programs and the accelerator program can become blurred in the interest of performance.
ALF architecture

Figure 4-12 (from the Accelerated Library Framework for Cell Broadband Engine Programmer’s Guide and API Reference) summarizes how ALF works.55

Near the top is the host view with presumably large memory areas for figuring the input and output data for the function that is to be accelerated. In the middle, lies the data partitioning where the input and output are split into smaller chunks, small enough to fit in the accelerator memory. They are “work blocks.” At the bottom, the accelerator tasks process one work block at a time. The data transfer part between the host and accelerator memory is handled by the ALF run time.

Figure 4-13 on page 301 shows the split between the host task and the accelerator tasks. On the host side, we create the accelerator tasks, create the work blocks, enqueue the blocks, and wait for the tasks to collectively empty the work block queue. On the accelerator task, for each work block, the ALF run time fetches the data from the host memory, calls the user provided computational kernel, and sends the output data back to the host memory.

55 You can find the Accelerated Library Framework for Cell Broadband Engine Programmer’s Guide and API Reference on the Web at the following address:
A simplified view of an accelerator task workflow

To illustrate how the ALF run time works on the accelerator side, Example 4-68 shows simplified pseudo code of the accelerator task. This program is similar to what ALF provides. An application programmer only needs to register the routines that are called by the call_user_routine. This is obviously not the real ALF source code, but it shows what it does in a simplistic way.

Example 4-68 A pseudo code for the accelerator workflow

// This SPE program is started by the alf_task_create() call on the
// host program.
int main()
{
    // If the user provided a context setup routine, call it now
    call_user_routine(task_context_setup);

    // Enter the main loop, we wait for new work blocks
    while (more_work_blocks()) {
        while(multi_use_count) { // more on this later
            get_user_data(work_block_parm);

            // If the user wants to have the data partitioning on
            // the accelerator, call the routine he gave us to do so.
            if(accelerator_data_partitioning) {
                call_user_routine(input_prepare);
            } // end if
        } // end while
    } // end while
} // end main

Figure 4-13 The host and accelerator sides
// The input data list is ready. Fetch the items from the
// host into the task input buffer.
get_input_data();

// We have the input data. Let’s call the user function.
call_user_routine(compute_kernel);

// If the output data partitioning is to be run on the
// accelerator, call the user provided routine.
if(accelerator_data_partitioning) {
    call_user_routine(output_prepare);
}

// The output data list is ready, scatteer the data back to
// the host memory.
put_output_data();

// We are about to leave. If we were asked to merge the context,
// do it now with the user provided routine.
call_user_routine(task_context_merge);

---

Description of the ALF concepts
ALF manipulates the following entities:

- Computational kernels,
- Tasks
- Work blocks
- Data partitioning
- Data sets

We describe each topic in more detail in the sections that follow.

**Computational kernel**
The ALF is used to offload the computation kernels of the application from the
host task to accelerator tasks running on the accelerator nodes, which are the
SPEs in our case. A computational kernel is a function that requires input, does
some processing, and produces output data. Within the ALF, a computational
kernel function has a given prototype to which application programmers must
conform. In the simplest case, an application programmer must implement a
single routine, which is the one that does the computation. In the most general
case, up to five functions might need to be written:

- The compute kernel
- The input data transfer list prepare function
- The output data transfer list prepare function
- The task context setup function
- The task context merge function

The full prototypes are given in Example 4-69 and taken from the alf_accel.h
file, which an accelerator program must include. The ALF run time fills in the
buffers (input, output, context) before calling the user-provided function. From an
application programmer’s perspective, the function gets called after the run time
has filled in all the necessary data, transferring the data from host memory to
accelerator memory, that is, the local storage. The programmers do not have to
be concerned with that. They are given pointers to where the data has been
made available. This is similar to what the shell does when the main() function of
a Linux program is called. The runtime system (the exec() system call set to run
in this case) has filled in the char *argv[] array to use.

**Example 4-69 The accelerator functions prototypes**

```c
/* This is the compute kernel. It is called for every work block.*/
/* Some arguments may be NULL. For example, the inout buffer may */
/* not be used. */

/* The task context data pointed to by p_task_context is filled */
/* at task startup only, not for every work block, but we want to */
/* be able to use this state data from inside the compute kernel */
/* every time we get called. */

/* The current_count and total_count are 0 and 1 and can be ignored */
/* for single-use work blocks. See later for multi-use work blocks. */

/* The data pointed to by p_parm_context is filled every time we */
/* process a work block. In the case of a multi-use work block, */
/* it can be used to store data that is common to multi-use blocks */
/* invocations. */
int (*compute_kernel) (  
    void *p_task_context,  
    void *p_parm_context,  
    void *p_input_buffer,  
    void *p_output_buffer,  
    void *p_inout_buffer,  
    int current_count,  
    int total_count);
```
The two routines below are used when we do the data
partitioning on the accelerator side, possibly because
this requires too much work for the PPE to keep up with
the SPEs. If we stick to host data partitionning, we do
not define these routines.

The area pointed to by `p_dtl` is given to us by the ALF runtime.
We will use this pointer as a handle and pass it as an argument to
the functions we will call to add entries to the list of items
that need to be brought in (resp. out) before (resp. after) the
compute kernel is called. The `parm_context` data may contain
information required to compute the data transfer lists.

```c
int (*input_list_prepare or output_list_prepare) (         
    void *p_task_context,
    void *p_parm_context,
    void *p_dtl,
    int current_count,
    int total_count);
```

The `task_context_setup` function is called at task startup time.
This function can be used to prepare the necessary environment
to be ready when the work blocks will be sent to us.

```c
int (*task_context_setup) (                             
    void *p_task_context);
```

The `task_context_merge` function is called at task exit time. It can
be used for reduction operations. We update our task context data
`(p_task_context)` by applying a reduction operation between this
data and the incoming context data that is filled for us by
the runtime.

```c
int (*task_context_merge) (                              
    void *p_context_to_merge,
    void *p_task_context);
```
The computational kernel functions must be registered to the ALF run time in order for them to be called when a work block is received. This is accomplished by using export statements that usually come at the end of the accelerator source code. Example 4-70 on page 305 shows the typical layout for an accelerator task.

Example 4-70  Export statements for accelerator functions

```c
#include <alf_accel.h>

int foo_comp_kernel(...)
{
    // statements here...
}

int foo_input_prepare(...)
{
    // statements here
}

int foo_output_prepare(...)
{
    // statements here
}

int foo_context_setup(...)
{
    // statements here
}

int foo_context_merge(...)
{
    // statements here
}

ALF_ACCEL_API_LIST_BEGIN
    ALF_ACCEL_EXPORT_API("foo_compute",foo_comp_kernel);
    ALF_ACCEL_EXPORT_API("foo_input_prepare",foo_input_prepare);
    ALF_ACCEL_EXPORT_API("foo_output_prepare",foo_output_prepare);
    ALF_ACCEL_EXPORT_API("foo_context_setup",foo_context_setup);
    ALF_ACCEL_EXPORT_API("foo_context_merge",foo_context_merge);
ALF_ACCEL_API_LIST_END
```

It is important to understand that we do not write a `main()` program for the accelerator task. The ALF run time runs the `main()` function and calls the functions upon receiving a work block.
**Tasks and task descriptors**

A _task_ is a ready-to-be-scheduled instantiation of an accelerator program, an SPE here. The tasks are created and finalized on the host program. A task is created by the `alf_task_create()` call. Before calling the task creation routine, the programmer must describe it, which is done by setting the attributes of a task descriptor. Example 4-71 shows the task descriptor, using a sample of pseudo C code.

**Example 4-71  A task descriptor**

```c
//

struct task_descriptor {

    // The task context buffer holds status data for the task.
    // It is loaded at task started time and can be copied back
    // at task unloading time. It’s meant to hold data that is
    // kept across multiple invocations of the computational kernel
    // with different work blocks

    struct task_context {
        task_context_buffer [SIZE];
        task_context_entries [NUMBER];
    };

    // These are the names of the functions that this accelerator task
    // implements. Only the kernel function is mandatory. The context
    // setup and merge, if specified, get called upon loading and
    // unloading of the task. The input and output data transfer list
    // routines are called when the accelerator does the data
    // partitioning itself.

    struct accelerator_image {
        char *compute_kernel_function;
        char *input_dtl_prepare_function;
        char *output_dtl_prepare_function;
        char *task_context_setup_function;
        char *task_context_merge_function;
    };

    // Where is the data partitioning actually run ?

    enum data_partition
        {HOST_DATA_PARTITION, ACCEL_DATA_PARTITION};

    // Work blocks. A work block has an input and output buffer.
    // These can overlap if needed. A block can also have parameters
    // and a context when the block is a multi-use block
```
struct work_blocks {
    parameter_and_context_size;
    input_buffer_size
    output_buffer_size;
    overlapped_buffer_size;
    number_of_dtl_entries;
};

// This is required so that the ALF runtime can work out
// the amount of free memory space and therefore how much
// multi-buffering can be done.
accelerator_stack_size;
};

The task context is a memory buffer that is used for two purposes:

- Store persistent data across work blocks. For example, the programmer loads
  state data that is to be read every time we process a work block. It contains
  work block “invariants.”
- Store data that can be reduced between multiple tasks. The task context can
  be used to implement all-reduce (associative) operations such as min, max,
  or global sum.

**Work blocks**

A work block is a single invocation of a task with a given set of input, output, and
parameter data. Single-use work blocks are processed only once, and multi-use
work blocks are processed up to total_count times.

The input and output data description of single-use work blocks can be
performed either at the host level or the accelerator level. For multi-use work
blocks, data partitioning is always performed at the accelerator level. The current
count of the multi-use work block and the total count are passed as arguments
every time the input list preparation routine, the compute kernel, and the output
data preparation routine are called.

With the multi-use blocks, the work block creation loop that was running on the
host task is now performed jointly by all the accelerator tasks that this host has
allocated. The only information that a given task is given to create, on the fly, the
proper input and output data transfer lists is the work block context buffer, the
current and total counts. The previous host loop is now parallelized across the
accelerator tasks which balance the work automatically between themselves.
The purpose of the multi-use work blocks is to make sure that the PPE, which
runs the host tasks, does not become a bottleneck, too busy creating work blocks
for the SPEs.
The work blocks queue is managed by the ALF run time, which balances the work across the accelerators. API calls influence the way that the ALF work blocks are allocated if the default mechanism is not satisfactory.

**Data partitioning**

Data partitioning ensures that each work block gets the right data to work with. The partitioning can be performed at the host level or accelerator level. We use a data transfer list, consisting of multiple entries of type `<start address, type, count>` that describe from where, in the host memory, we must gather data to be sent to the accelerators. The API calls differ whether you use the host or accelerator data partitioning.

**Data sets**

At the host level, data sets are created that assign attributes to the data buffers that are to be used by the accelerator tasks. A memory region can be described as read-only, read-write, or write-only. This information gives hints to the ALF run time to help improve the data movement performance and scheduling.

**The memory layout of an accelerator task**

Memory on the accelerator is a limited resource. It is important to understand how the various data buffers are organized to tune their definitions and usage. Also, the ALF run time has more opportunities to use clever multi-buffering techniques if it has more room left after the user data has been loaded into the accelerator memory.

Figure 4-14 shows the memory map of an ALF program. The user code contains the computational kernels and optionally the input/output data transfer list functions and context setup/merge functions.
As shown in Figure 4-15, five pointers to data buffers are passed to the computational kernels (see Example 4-69 on page 303). Various combinations are possible, depending on the use of overlapped buffers for input and output. In the simplest case, no overlap exists between the input and output buffers.

![Diagram of memory map without overlapped input/output buffer]

No overlapped input and output buffers. p_inout is undefined.

*Figure 4-15  Memory map without overlapped input/output buffer*
The input and output data buffers can overlap entirely as shown in Figure 4-16.

Overlapped input and output buffers only. 
$p_{\text{input}}$ and $p_{\text{output}}$ are undefined.

*Figure 4-16  Memory map with a single input/output overlapped buffer*
In the most general case, three data buffer pointers can be defined as shown in Figure 4-17.

![Diagram showing memory map with all five data pointers defined](image)

**API description**

The API has two components: the host API and the accelerator. A host program must include the `alf.h` file, and an accelerator program must include the `alf_accel.h` file.
Table 4-27 lists the main host API functions.

**Table 4-27  The host side of the ALF API**

<table>
<thead>
<tr>
<th>Groups</th>
<th>Functions</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Framework</td>
<td>alf_init</td>
<td>▶ Initializes the ALF</td>
</tr>
<tr>
<td></td>
<td>alf_query_system_info</td>
<td>▶ Queries the various system information</td>
</tr>
<tr>
<td></td>
<td>alf_num_instances_set</td>
<td>▶ Sets the maximum number of accelerators</td>
</tr>
<tr>
<td></td>
<td>alf_exit</td>
<td>▶ Exits the ALF</td>
</tr>
<tr>
<td></td>
<td>alf_task_desc_create</td>
<td>▶ Creates a task descriptor</td>
</tr>
<tr>
<td></td>
<td>alf_task_desc_set_int32</td>
<td>▶ Sets a task descriptor parameter</td>
</tr>
<tr>
<td></td>
<td>alf_task_desc_set_int64</td>
<td>▶ Sets a task descriptor parameter 64 bit</td>
</tr>
<tr>
<td></td>
<td>alf_task_desc_ctx_entry_add</td>
<td>▶ Adds an entry in the task context</td>
</tr>
<tr>
<td></td>
<td>alf_task_desc_destroy</td>
<td>▶ Destroys the context</td>
</tr>
<tr>
<td></td>
<td>alf_task_create</td>
<td>▶ Creates a task</td>
</tr>
<tr>
<td></td>
<td>alf_task_finalize</td>
<td>▶ Makes the task runnable</td>
</tr>
<tr>
<td></td>
<td>alf_task_destroy</td>
<td>▶ Terminates a task</td>
</tr>
<tr>
<td></td>
<td>alf_task_wait</td>
<td>▶ Waits for a task termination</td>
</tr>
<tr>
<td></td>
<td>alf_taskdepends_on</td>
<td>▶ Expresses a task dependency</td>
</tr>
<tr>
<td></td>
<td>alf_wb_create</td>
<td>▶ Creates a work block</td>
</tr>
<tr>
<td></td>
<td>alf_wb_parm_add</td>
<td>▶ Adds a work block parameter</td>
</tr>
<tr>
<td></td>
<td>alf_wb_dtl_begin</td>
<td>▶ Starts a data transfer list</td>
</tr>
<tr>
<td></td>
<td>alf_wb_dtl_entry_add</td>
<td>▶ Adds an entry to the list</td>
</tr>
<tr>
<td></td>
<td>alf_wb_dtl_end</td>
<td>▶ Closes the data transfer list</td>
</tr>
<tr>
<td></td>
<td>alf_wb_enqueue</td>
<td>▶ Queues the work block for execution</td>
</tr>
<tr>
<td></td>
<td>al_dataset_create</td>
<td>▶ Creates a dataset structure</td>
</tr>
<tr>
<td></td>
<td>alf_dataset_buffer_add</td>
<td>▶ Adds a buffer and type to the data set</td>
</tr>
<tr>
<td></td>
<td>alf_task_dataset_associate</td>
<td>▶ Associates the data set with a task</td>
</tr>
<tr>
<td></td>
<td>alf_dataset_destroy</td>
<td>▶ Destroys the dataset structure</td>
</tr>
</tbody>
</table>

The accelerator API is much leaner because it includes only a few functions to perform the data partitioning. See Table 4-28.

**Table 4-28  The accelerator side of the ALF API**

<table>
<thead>
<tr>
<th>Groups</th>
<th>Functions</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Framework</td>
<td>alf_accel_num_instances</td>
<td>▶ Number of accelerators</td>
</tr>
<tr>
<td></td>
<td>alf_accel_instance_id</td>
<td>▶ Personal rank</td>
</tr>
<tr>
<td>Data partitioning</td>
<td>ALF_ACCEL_DTL_BEGIN</td>
<td>▶ Starts a data transfer list</td>
</tr>
<tr>
<td></td>
<td>ALF_ACCEL_DTL_ENTRY_ADD</td>
<td>▶ Adds to the list</td>
</tr>
<tr>
<td></td>
<td>ALF_ACCEL_DTL_END</td>
<td>▶ Closes the list</td>
</tr>
</tbody>
</table>
**ALF optimization tips**
Apart from tuning the computational kernel itself and ensuring that the amount of work per data communication is maximized, it can be beneficial to tune the data movement part. To do so, explore the following techniques:

- Data partitioning on the accelerator side
- Multi-use work blocks

These techniques lower the workload on the host task, which otherwise might not be able to keep up with the speed of the accelerators, thus becoming a bottleneck for the whole application. Also, using data sets on the host side and using overlapped input and output buffers whenever possible gives more flexibility to the ALF run time to optimize the data transfers.

**ALF application development notes**
When designing an ALF strategy for an application, a trade-off is necessary to decide on the granularity of the computational kernels. Consider the following forces:

- The ability to extract independent pieces of work
- The computation to communication ratio
- The memory constraints imposed by the SPE

From an application development perspective, the host-accelerator model allows two different types of programmers to work on the same project. For example, the developer named Sam can concentrate on the high level view, implementing the application algorithm, managing MPI tasks, and making sure they synchronize and communicate when needed. Sam is also responsible for writing the “contract” between the task and the accelerator tasks and describing the input and output data as well as the required operation. Then the developer named Gordon focuses on implementing the computational kernels according to the specifications and tunes these kernels.

The examples in the following sections show the type of work that is involved when accelerating applications with the ALF. Of particular interest in this respect are the matrix_add and matrix_transpose examples.

**A guided tour of the ALF examples provided in the SDK**
Before embarking on the acceleration of an application by using the ALF, we recommend that you look at the examples provided by the IBM SDK for Multicore Acceleration. The examples come with two rpms:

- alf-examples-source
- alf-hybrid-examples-source
The hybrid version includes non-hybrid rpms too. The examples are described in Table 4-29 and are presented in suggested order of use.

### Table 4-29 ALF examples in the IBM SDK for Multicore Acceleration

<table>
<thead>
<tr>
<th>Example</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>hello_world</td>
<td>Simple, minimal ALF program</td>
</tr>
</tbody>
</table>
| matrix_add    | This example gives the steps that were taken to enable and tune this application with the ALF and includes the following successive versions:  
  - scalar: The reference version  
  - host_partition: First ALF version, data partitioning on the host  
  - host_partition_simd: The compute kernel is tuned using SIMD  
  - accel_partition: Data partitioning performed by the accelerators  
  - dataset: Use of the dataset feature  
  - overlapped_io: Use of overlapped input and output buffers |
| PI            | Shows the use of task context buffers for global parameters and reduction operations |
| pipe_line     | Shows the implementation of a pipeline by using task dependencies and task context merge operations |
| FFT16M        | Shows multi-use work blocks and task dependencies                            |
| BlackScholes  | Pricing model: Shows how to use multi-use work blocks                        |
| matrix_transpose | Like matrix_add, shows the steps going from a scaler version to a tuned ALF version and includes the following successive versions:  
  - scalar: the reference version  
  - STEP1a: Using ALF and host data partitioning  
  - STEP1b: Using accelerator data partitioning  
  - STEP2: Using a tuned SIMD computational kernel |
| inout_buffer  | Shows the use of input/output overlapped buffers                              |
| task_context  | Shows the use of the task context buffer for associative reduction operations (min, max), a global sum, and a storage for a table lookup. |
| inverse_matrix_ovl | Shows the use of function overlay, data sets                                  |

### 4.7.3 Domain-specific libraries

In this section, we discuss the main domain specific libraries that are part of SDK 3.0. These libraries aim to assist Cell/B.E. programmers by providing reusable functions that implement a set of common algorithms and mathematical operators. The functions are optimized specifically to the Cell/B.E. processor by exploiting its unique architecture, for example, running parallel on several SPEs and using SIMD instructions.
A software developer who starts developing an application for the Cell/B.E. processor (or ports an existing application) can first check whether some parts of its application are already implemented in one of the SDK’s domain specific libraries. If it is, then using the corresponding library can provide an easy solution to save development efforts.

Most of those libraries are open source. Therefore, even if the exact functionality required by the developed application is not implemented, the programmer can use those functions as a reference and be customized and tailored for developing the application specific functions.

In the next four sections, we provide a brief description of the following libraries:

- Fast Fourier Transform library
- Monte Carlo libraries
- Basic Linear Algebra Subprograms library
- Matrix, large matrix, and vector libraries

While we found the libraries to be the most useful, SDK 3.0 provides several other libraries. The *Example Library API Reference* document discusses the additional libraries and provides detailed description of some of the libraries that are discussed in this chapter (and are described only briefly).\(^{56}\)

### Fast Fourier Transform library

The FFT prototype library handles a wide range of FFTs and consists of the following parts:

- An API for the following routines used in single precision:
  - 1D of 2D FFT
  - FFT Real -> Complex 1D
  - FFT Complex-Complex 1D
  - FFT Complex -> Real 1D
  - FFT Complex-Complex 2D for frequencies (from 1000x1000 to 2500x2500)

  The implementation manages sizes up to 10000 and handles multiples of 2, 3, and 5 as well as powers of those factors, plus one arbitrary factor as well. User code running on the PPU uses the Cell/B.E. FFT library by calling one of the streaming functions. An SPU version is also available.

- Power-of-two-only 1D FFT code for complex-to-complex single and double precision processing
  It is supported on the SPU only.

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\(^{56}\) The *Example Library API Reference* is available on the Web at the following address:
Both parts of the library run by using a common interface that contains an initialization and termination step, and an execution step that can process “one-at-a-time” requests (streaming) or entire arrays of requests (batch). The latter batch mode is more efficient in applications in which several distinct FFT operation might be executed one after the other. In this case, the initialization and termination steps are done only once for the FFT execution. Initialization is done before the first execution, and termination is done after the last execution.

Both the FFT transform and inverse FFT transform are supported by this library.

To retrieve more information about this library, the programmer can do either of the following actions:

- Enter the following command on a system where the SDK is installed:
  
  man /opt/cell/sdk/prototype/usr/include/libfft.3

- Read the “Fast Fourier Transform (FFT) library” chapter in the Example Library API Reference document.\(^{57}\)

Another alternative library that implements FFT for the Cell/B.E. processor is the FFTW library. The Cell/B.E. implementation of this library is currently available only as an alpha preview release. For more information, refer to the following Web address:

http://www.fftw.org/cell/

**Monte Carlo libraries**

The Monte Carlo libraries are a Cell/B.E. implementation of random number generator (RNG) algorithms and transforms. The objective of this library is to provide functions that are needed to perform Monte Carlo simulations.

The library contains the following items:

- Four RNG algorithms: Hardware-generated, Kirkpatrick-Stoll, Mersenne Twister, and Sobol
- Three distribution transformations: Box-Muller, Moro’s Inversion, and Polar method
- Two Monte Carlo simulation samples: Calculations of pi and the volume of an n-dimensional sphere

For a detailed description of this library and how to use, refer to the Monte Carlo Library API Reference Manual document.\(^{58}\)

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57 See note 56 on page 315.

Basic Linear Algebra Subprograms library

The BLAS library is based upon a published standard interface (described in the *BLAS Technical Forum Standard* document)\(^{59}\) for commonly used linear algebra operations in high-performance computing (HPC) and other scientific domains. It is widely used as the basis for other high quality linear algebra software, for example LAPACK and ScaLAPACK. The Linpack (HPL) benchmark largely depends on a single BLAS routine (DGEMM) for good performance.

The BLAS API is available as standard ANSI C and standard FORTRAN 77/90 interfaces. BLAS implementations are also available in open-source (netlib.org). Based on its functionality, BLAS is divided into three levels:

- Level 1 routines are for scalar and vector operations.
- Level 2 routines are for matrix-vector operations.
- Level 3 routines are for matrix-matrix operations.

Each routine has four versions:

- Real single precision
- Real double precision
- Complex single precision
- Complex double precision

The BLAS library in SDK 3.0 supports only real single precision and real double precision versions.

All single-precision and double-precision routines in the three levels of standard BLAS are supported on the PPE. These are available as PPE APIs and conform to the standard BLAS interface.

Some of the routines have been optimized by using the SPEs and show a marked increase in performance in comparison to the corresponding versions that are implemented solely on the PPE. The optimized routines have an SPE interface in addition to the PPE interface. However, the SPE interface does not conform to the standard BLAS interface and provides a restricted version of the standard BLAS interface.

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\(^{59}\) You can find the *BLAS Technical Forum Standard* document on the Web at the following address: [http://www.netlib.org/blas/blast-forum/](http://www.netlib.org/blas/blast-forum/)
The single precision versions of these routines have been further optimized for maximum performance by using various features of the SPE (for example SIMD, Dual Issue, and so on):

- Level 1:
  - SSCAL, DSCAL
  - SCOPY, DCOPY
  - ISAMAX, IDAMAX
  - SAXPY, DAXPY
  - SDOT, DDOT

- Level 2:
  - SGEMV, DGEMV (TRANS='No transpose' and INCY=1)

- Level 3:
  - SGEMM, DGEMM
  - SSYRK, DSYRK (Only for UPLO='Lower' and TRANS='No transpose')
  - STRSM, DTRSM (Only for SIDE='Right', UPLO='Lower', TRANS='Transpose' and DIAG='Non-Unit')

For a detailed description of this library and how to use it, refer to the BLAS Programmer's Guide and API Reference document.60

Matrix, large matrix, and vector libraries
SDK 3.0 provides three libraries that implement various linear operation on matrixes and vectors:

- Matrix library
- Large matrix library
- Vector library

The matrix library consists of various utility libraries that operate on 4x4 matrices as well as quaternions. The library is supported on both the PPE and SPE. In most cases, all 4x4 matrices are maintained as an array of four 128-bit SIMD vectors, while both single-precision and double-precision operands are supported.

The large matrix library consists of various utility functions that operate on large vectors and large matrices of single precision floating-point numbers. The size of the input vectors and matrices are limited by the local storage size of the SPE. This library is currently only supported on the SPE.

60 The BLAS Programmer's Guide and API Reference is available at the following address:
The matrix and large matrix libraries support different matrix operations such as multiplying, adding, transpose, and inverse. Similar to the SIMDmath and MASS libraries, the libraries can be used either as a linkable library archive or as a set of inline function headers. For more details, see “SIMD Math Library” on page 262 and “MASS and MASSV libraries” on page 264.

The vector library consists of a set of general purpose routines that operate on vectors. This library is supported on both the PPE and SPE.

For a detailed description of the three libraries and how to use them, see the “Matrix library,” “Large matrix library,” and “Vector library” chapters in the Example Library API Reference document.\(^{61}\)

### 4.8 Programming guidelines

In this section, we provide a general collection of programming guidelines and tips that cover different aspects of Cell/B.E. programming. This section heavily relies on information from the following resources:

- “Maximizing the power of the Cell Broadband Engine processor: 25 tips to optimal application performance”, by Daniel A. Brokenshire \(^{14}\)
- “Cell/B.E. Programming Gotchas! or Common Rookie Mistakes”, Michael Perrone, IBM Watson Research Center
  

- The “General SPE programming tips” chapter in the Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial\(^ {62}\)

- The “SPE programming tips” chapter in the Cell Broadband Engine Programming Handbook\(^ {63}\)

In addition, refer to the following sources of information for high performance programming in the CBEA:

- Cell Broadband Engine Architecture forum at IBM developerWorks:
  

- Cell Performance Web site:
  
  [http://www.cellperformance.com](http://www.cellperformance.com)

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\(^{61}\) See 56 on page 315.

\(^{62}\) See note 3 on page 80.

\(^{63}\) See note 1 on page 78.
4.8.1 General guidelines

Follow these general guidelines for Cell/B.E. programming:

- Offload as much work onto the SPEs as possible. Use the PPE as the control processor and the SPE to perform all the heavy computational lifting.

- Exploit Cell/B.E. parallelism:
  - Use multithreading so that parallel tasks run on separate SPE threads.
  - Try to avoid using more threads than physical SPEs because context switching consumes a fair amount of time.
  - Do not spawn SPE threads for each scalar code hot spot since thread creation overhead reduces performance.

- Choose a partitioning and work allocation strategy that minimizes atomic operations and synchronization events.

- Use the appropriate programming model to your application. For more information, see 3.3, “Deciding which parallel programming model to use” on page 53.

- Choose the fixed point data types carefully:
  - Floating point
    As most processors, the SPE has better performance when performing SIMD operations on single-precision floating point variables compared to double-precision ones. Alternatively, double-precision operations are more accurate. Therefore, we recommend that you use double-precision types only in case the accuracy of single-precision floating point variables is not sufficient.
  - Fixed point
    Similarly to floating point, 32-bit integers have better performance than 64-bit integers. In addition, specifically for multiply, 16-bit fixed-point integers have better performance than 32-bit fixed-point integers.

- Use the volatile keyword for the declaration of DMA buffers in order to instruct the compiler, not to reorder software memory access to those buffers and DMA requests and waiting for completion. Usage of the volatile keyword can significantly impact the compiler’s ability to order buffer access and coalesce multiply loads.
4.8.2 SPE programming guidelines

In this section, we briefly summarize the programming guidelines for optimizing the performance of SPE programs. The intention is to address programming issues that are related only to programming the SPU itself, without interacting with external components, such as PPE, other SPEs, or main storage.

Since almost any SPE program needs to interact with external components, we recommend that you become familiar with the programming guidelines in 4.8, “Programming guidelines” on page 319.

General

- Avoid over usage of 128-byte alignment. Consider cases in which alignment is essential, such as for data transfers that are performed often, and use redundant alignment, for example 16 bytes, for other cases. There are two main reasons why 128-byte alignment can reduce the performance:
  - The usage of 128-byte alignment requires the global definition of the variables. This causes the program to use more registers and reduces the number of free registers which also reduces performance, for example increased loads and stores, increased stack size, and reduced loop unrolling. Therefore, if only redundant alignment is required, for example a 16-byte alignment, then you can use the variables as local, which can significantly increase performance.
  - The usage of 128-byte alignment increases the code size because of the padding that is added by the compiler to make the data aligned.

- Avoid writing recursive SPE code that uses a lot of stack variables because they can cause stack overflow errors. The compilers provide support for runtime stack overflow checking that can be enabled during application debugging of such errors.

Intrinsics

- Use intrinsics to achieve machine-level control without needing to write assembly language code.

- Understand how the intrinsics map to assembly instructions and what the assembly instructions do.

Local storage

- Design for the LS size. The LS holds up to 256 KB for the program, stack, local data structures, heap, and DMA buffers. You can do a lot with 256 KB, but be aware of this size.
In case the code is too large too fit into the LS (taking into account that the data should also reside in the LS), use the overlays mechanism. See 4.6.7, “Code transfer using SPU code overlay” on page 283.

Use code optimization carefully since it can increase the code size, for example function inline and loop unrolling.

Loops

If the number of loop iterations is a small constant, then consider removing the loop altogether.

If the number of loop iterations is variable, consider unrolling the loop as long as the loop is relatively independent. That is, an iteration of the loop does not depend upon the previous iteration. Unrolling the loops reduces dependencies and increases dual-issue rates. By doing so, compiler optimization can exploit the large SPU register file.

SIMD programming

Exploit SIMD programming as much as possible, which can increase the performance of your application in several ways, especially in regard to computation bounded programs.

Consider using the compiler auto-SIMDizing feature. This feature can convert ordinary scalar code into SIMD code. Be aware of the compiler limitations and on the code structures that are supported for auto-SIMDizing. Try to code according to the limitations and structures. For more information, see 4.6.5, “Auto-SIMDizing by compiler” on page 270.

An alternative is to explicitly do SIMD programming by using intrinsics, SIMD libraries, and supported data types. For more information, see 4.6.4, “SIMD programming” on page 258.

Not all SIMD operations are supported by all data types. Review the operations that are critical to your application and verify in which data types they are supported.

Choose an SIMD strategy that is appropriate for the application that is under development. The two common strategies are AOS and SOA:

- Array-of-structure organization
  From a programmer’s point of view, you can have more-efficient code size and simpler DMA needs, but SIMDize is more difficult. From a computation point of view, this organization can be less efficient, but depends on the specific application.

- Structure-of-arrays organization
From a programmer’s point of view, this organization is usually easier to SIMDize, but the data must be maintained in separate arrays or the SPU must shuffle the AOS data into an SOA form.

If the data is in AOS organization, consider runtime converting the AOS data to SOA, performing the calculations, and converting the results back. For more information, see “Data organization: AOS versus SOA” on page 267.

In general, note the following guidelines regarding SIMD programming:

– Use of auto-SIMDizing requires less development time, but in most cases, the performance is inferior compared to explicit SIMD programming, unless the program can perfectly fit into the code structures that are supported by the compiler for auto-SIMDizing.

– On the contrary, correct SIMD programming provides the best performance, but requires non-negligible development effort.

**Scalars**

– Because SPUs only support quadword loads and stores, scalar loads and stores (less than a quadword) are slow, with long latency.

– Align the scalar loads that are often used with the quadword address to improve the performance of operations that are done on those scalars.

– Cluster scalars into groups and load multiple scalars at a time by using quadword memory access. Later use extract or insert intrinsics to explicitly move between scalars and vector data types, which eliminates redundant loads and stores.

For details, see 4.6.6, “Working with scalars and converting between different vector types” on page 277

**Branches**

– Eliminate nonpredicted branches by using select bit intrinsics (spu_sel).

– For branches that are highly predicted, use the __builtin_expect directive to explicitly do direct branch prediction. Compiler optimization can add the corresponding branch hint in this case.

– Inline functions are often called by explicitly defining them as inline in the program, or use compiler optimization.

– Use feedback-directed optimization, for example by using the FDPRPro tool.

For details, see 4.6.8, “Eliminating and predicting branches” on page 284.
Multiplies

- Avoid integer multiplies on operands that are greater than 16 bits in size. The SPU supports only a 16-bit x16-bit multiply. A 32-bit multiply requires five instructions (three 16-bit multiplies and two adds).

- Keep array elements sized to a power of 2 to avoid multiplies when indexing.

- When multiplying or dividing by a factor that is power of 2, instead of using the ordinary operators, use the shift operation (corresponding intrinsics for vectors and << and >> operator for scalars).

- Cast operands to an unsigned short prior to multiplying. Constants are of type int and require casting. Use a macro to explicitly perform 16-bit multiplies to avoid inadvertent introduction of signed extends and masks due to casting.

Dual issue

- Choose intrinsics carefully to maximize dual-issue rates or reduce latencies.

- Dual issue occurs if a pipe-0 instruction is even-addressed, a pipe-1 instruction is odd-addressed, and there are no dependencies (operands are available).

- Manually insert nops to align instructions for dual issue when writing non-optimizing assembly programs. In other cases, the compilers automatically insert nops when needed.

- Use software pipeline loops to improve dual-issue rates as described in the “Loop unrolling and pipelining” chapter in the Cell Broadband Engine Programming Handbook.64

- Understand the fetch and issue rules to maximize the dual-issue rate. The rules are explained in the “SPU pipelines and dual-issue rules” chapter in the Cell Broadband Engine Programming Handbook.

- Avoid over usage of the odd pipeline for load instructions, which can cause instruction starvation. This can happen, for example, on a large matrix transpose on SPE when there are many loads on an odd pipeline and minimal usage of the even pipeline for computation. A similar case can happen for the dot product of large vectors. To solve this problem, the programmer can add more computation on the data that is being loaded.

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64 See note 1 on page 78.
4.8.3 Data transfers and synchronization guidelines

In this section, we provide a summary of the programming guidelines for performing efficient data transfer and synchronization on the Cell/B.E. program:

- Choose the transfer mechanism that fits your application data access pattern:
  - If the pattern is predictable, for example sequential access array or matrix, use explicit DMA requests to transfer data. The requests can be implemented with SDK core libraries functions.
  - If the pattern is random or unpredictable, for example a sparse matrix operation, consider using the software manage cache, especially if there is a high ratio of data re-use, for example the same data or cache line is used in different iterations of the algorithm.

- When the core libraries for explicitly initiating DMA transfer:
  - Follow the supported and recommended values for the DMA parameters. See “Supported and recommended values for DMA parameters” on page 116 and “Supported and recommended values for DMA-list parameters” on page 117.
  - DMA throughput is maximized if the transfers are at least 128 bytes, and transfers greater than or equal to 128 bytes should be cache-line aligned (aligned to 128 bytes). This refers to the data transfer size and source and destination addresses as well.
  - Overlap DMA data transfer with computation by using a double-buffering or multibuffering mechanism. See 4.3.7, “Efficient data transfers by overlapping DMA and computation” on page 159.
  - Minimize small transfers. Transfers of less than one cache line consume a bus bandwidth that is equivalent to a full cache-line transfer.

- When explicitly using software managed cache, try to exploit the unsafe asynchronous mode because it can provide significantly better results than the unsafe synchronous mode. A double mechanism can also be implemented by using this safe mode.

- Uniformly distribute memory bank accesses. The Cell/B.E. memory subsystem has 16 banks, interleaved on cache line boundaries. Addresses of 2 KB apart access the same bank. System memory throughput is maximized if all memory banks are uniformly accessed.
Use SPE-initiated DMA transfers rather than PPE-initiated DMA transfers. There are more SPEs than PPEs (only one), and the PPE can enqueue only eight DMA requests, where each SPE can enqueue 16 DMA requests. In addition, the SPE is much more efficient at enqueuing DMA requests.

Use a kernel with large 64 KB base pages to reduce page table and TLB thrashing. If significantly large data sets are accessed, consider using huge pages instead. See 4.3.8, “Improving the page hit ratio by using huge pages” on page 166.

For applications that are memory bandwidth-limited, consider using NUMA. See 4.3.9, “Improving memory access using NUMA” on page 171. NUMA is recommended in a system of more than one Cell/B.E. node (such as QS20 and QS21) such as in the following two common cases:

- Only one Cell/B.E. node is used. Since access latency is slightly lower on node 0 (Cell/B.E. 0) as compared to node 1 (Cell/B.E. 1), use NUMA to allocate memory and processor on this node.

- More than one Cell/B.E. node is used (for example, using the two nodes of QS21) and the data and tasks execution can be perfectly divided between nodes. In this case, NUMA can be used to allocate memory on both nodes and exploit the aggregated memory bandwidth. The processor on node 0 primarily accesses memory on this node, and the same happens for node 1.

DMA transfers from main storage have high bandwidth with moderate latency, where transfers from the L2 have moderate bandwidth with low latency. For that reason, consider the effect of whether the updated data is stored in the L2 versus on the main memory:

- When SPEs access large data sets, make sure that it is not on the L2. This can be done, for example, by making sure that the PPE does not access the data set before the SPEs do.

- When the SPEs and PPE must share short messages, such as a notification or status, we recommend that they do so on the L2. The sharing can be done, for example, by the PPE accessing the data before the SPEs, which ensures that the system creates a copy of this data on the L2.

- You can also have control over the L2 behavior by using the __dcbf function to flush a data cache block and the __dcbst function to store the data cache block in a cache.

- Most applications are better off not trying to over manage the PPE cache hierarchy.
Exploit the on-chip data transfer and communication mechanism by using LS-to-LS DMA transfers when sharing data between SPEs and using mailboxes, signal notification registers for small data communications, and synchronization. The reason is that the EIB provides significantly more bandwidth than system memory (in the order of 10 or more).

Be aware that when the SPEs receive a DMA put data transfer completion, the local MFC completed the transaction from its side but not unnecessarily that the data is already stored in memory. Therefore, it might not be accessible yet for other processors.

Use the explicit command to force data ordering when sharing data between SPEs and the PPE and between SPEs to themselves because the CBEA does not guarantee such ordering between the different storage domains. Coherency is guaranteed on each of the memory domains separately:

- The DMA can be re-ordered compared to the order in which the SPE program initiates the corresponding DMA commands. Explicit DMA ordering commands must be issued to force ordering.
- Use fence or barrier DMA commands to order DMA transfers within a tag group.
- Use a barrier command to order DMA transfers within the queue.
- Minimize the use of ordering such commands because they have a negative effect on the performance.
- See 4.5, “Shared storage synchronizing and data ordering” on page 218, for more information and 4.5.4, “Practical examples of using ordering and synchronization mechanisms” on page 240, for practical scenarios.

Use affinity to improve the communication between SPEs (for example LS-to-LS DMA data transfer, mailbox, and signals). See 4.1.3, “Creating SPEs affinity by using a gang” on page 94, for more information.

Minimize the use of atomic, synchronizing, and data-ordering commands because they can add significant overhead.

Atomic operations operate on reservation blocks that correspond to 128-byte cache lines. As a result, place synchronization variables in their own cache line so that other non-atomic loads and stores do not cause inadvertent lost reservations.
4.8.4 Inter-processor communication

Use the following recommended methods for inter-processor communication:

► PPE-to-SPE communication:
  – PPE writes to the SPE inbound mailbox.
  – The SPE performs a blocking read of its inbound mailbox.

► SPE-to-PPE communication:
  – The SPE writes to system memory, which also invalidates the corresponding cache line in L2.
  – The PPE polls L2.

► SPE-to-SPE communication:
  – The SPE writes to the inbound mailbox of remote SPE, signal notification registers, or LS.
  – The SPE polls its inbound mailbox or signals the notification registers or LS.

Avoid having the PPE wait for the SPEs to complete by polling the SPE outbox mailbox.
Chapter 5. Programming tools and debugging techniques

In this chapter, we introduce and explore the plethora of available development tools for the Cell Broadband Engine Architecture (CBEA). We give special attention to the potential tool management issues that might arise in a heterogeneous architecture. For these reasons, we dedicate a large portion of this chapter to the debugger and available debugging techniques or tools, with a focus on both error detection and performance analysis.

The following topics are discussed:

- 5.1, “Tools taxonomy and basic time line approach” on page 330
- 5.2, “Compiling and building executables” on page 332
- 5.3, “Debugger” on page 345
- 5.4, “IBM Full System Simulator” on page 354
- 5.5, “IBM Multicore Acceleration Integrated Development Environment” on page 362
- 5.6, “Performance tools” on page 375
5.1 Tools taxonomy and basic time line approach

The design of the Cell Broadband Engine (Cell/B.E.) system presents many challenges for software development. With nine cores, multiple Instruction Set Architectures (ISAs) and non-coherent memory, the Cell/B.E. system imposes challenges to compiling, debugging and performance tuning. Therefore, it is important to understand the basic concepts of how the tools interlock with the operating system in order to achieve success in development.

5.1.1 Dual toolchain

The Cell/B.E. processor is a heterogeneous multiprocessor because the Synergistic Processor Elements (SPEs) and the Power Processor Element (PPE) have different architectures, disjoint address spaces, and different models of memory and resource protection. The PPE can run a virtual-memory operating system, so that it can manage and access all system resources and capabilities.

In contrast, the synergistic processor units (SPUs) are not intended to run an operating system. SPE programs can access the main-storage address space, called the effective address (EA) space, only indirectly through the direct memory access (DMA) controller in the memory flow controller (MFC).

The two processor architectures are different enough to require two distinct toolchains for software development.

Application binary interface

Application binary interface (ABI) establishes the set of rules and conventions to ensure portability of code and compatibility between code generators, linker and runtime libraries. Typically, the ABI states rules regarding data types, register usage, calling conventions, and object formats.

The toolchains for both the PPE and SPEs produce object files in the Executable and Linking Format (ELF). The ELF is a flexible, portable container for re-locatable, executable, and shared object (dynamically linkable) output files of assemblers and linkers. The terms PPE-ELF and SPE-ELF are used to differentiate between ELF for the two architectures.

Cell/B.E. Embedded SPE Object Format (CESOF) is an application of PPE-ELF that allows PPE executable objects to contain SPE executables. See Figure 5-1 on page 331. To ease the development of combined PPE-SPE multiprocessor programs, the Cell/B.E. operating-system model uses the CESOF and provides SPE process-management primitives.
Programmers often keep in mind a heterogeneous model of the Cell/B.E. processor when dividing an application program into concurrent threads. By using the CESOF format and, for example, the Linux operating-system thread application programming interfaces (APIs), programmers can focus on application algorithms. They do not have to spend time managing basic tasks such as SPE process creation and global variable sharing between SPE and PPE threads. From an application developer's point of view, it is important to note that such a mechanism also enables the access of PPE variables from the SPE code.
5.1.2 Typical tools flow

The typical tools usage pattern should be similar to the flow shown in Figure 5-2.

Figure 5-2  Typical development cycle

Throughout the remainder of this chapter, in addition to exploring the relevant characteristics of each tool, we show similar flows that relate the development cycle and where each tool fits within the cycle. We guide you where and when to use each tool.

5.2 Compiling and building executables

In this section, we explore the IBM Software Development Kit (SDK) 3.0 capabilities for compiling and optimizing executables and managing the build process in the Cell/B.E. environment. Figure 5-3 highlights the compile stage of the process.

Figure 5-3  Compile

5.2.1 Compilers: gcc

The GNU toolchain contains the GCC C-language compiler (GCC compiler) for the PPU and the SPU. For the PPU, this toolchain replaces the native GCC compiler on PowerPC platforms and is a cross-compiler on X86.
This release of the GNU toolchain includes a GCC compiler and utilities that optimize code for the Cell/B.E. processor:

- The spu-gcc compiler for creating an SPU binary
- The ppu-embedspu (and ppu32-embedspu) tool that enables an SPU binary to be linked with a PPU binary into a single executable program
- The ppu-gcc (and ppu32-gcc) compiler

Creating a program

In the following scenario, we create the executable program, called simple, that contains the SPU code, simple_spu.c, and PPU code, simple.c:

1. Compile and link the SPE executable as shown in Example 5-1.

   **Example 5-1  Compiling SPE code**

   ```bash
   #/usr/bin/spu-gcc -g -o simple_spu simple_spu.c
   ```

2. (Optional) Run the `embedspu` command to wrap the SPU binary into a CESOF linkable file that contains additional PPE symbol information. See Example 5-2.

   **Example 5-2   Embedding SPE code**

   ```bash
   #/usr/bin/ppu32-embedspu simple_spu simple_spu simple_spu-embed.o
   ```

3. Compile the PPE side and link it together with the embedded SPU binary (Example 5-3).

   **Example 5-3   Linking**

   ```bash
   #/usr/bin/ppu32-gcc -g -o simple simple.c simple_spu-embed.o -lspe2
   ```

   Alternatively compile the PPE side and link it directly with the SPU binary (Example 5-4). The linker invokes `embedspu` and uses the file name of the SPU binary as the name of the program handle struct.

   **Example 5-4   Implicitly linking**

   ```bash
   /usr/bin/ppu32-gcc -g -o simple simple.c simple_spu -lspe2
   ```
Fine tuning: Command-line options
Each of the GNU compilers offers Cell/B.E. relevant options, to either enable architecture specific options or further optimize the generated binary code.

**ppu-gcc options**
The ppu-gcc compiler offers the following options:

- **-mcpu=cell**
  This option selects the instruction set architecture to generate code for, either Cell/B.E. or PowerXCell. Code that is compiled with `-march=celledp` can use the new instructions and does not run on the Cell/B.E. processor.
  `-march=celledp` also implies `-mtune=celledp`.

- **-m32**
  Selects the 32-bit option. The ppu32-gcc defaults to 32 bit.

- **-m64**
  Selects the 64-bit option. The ppu-gcc defaults to 64 bit.

- **-maltivec**
  This option enables code generation that uses Altivec vector instructions (the default in ppu-gcc).

**spu-gcc options**
The spu-gcc compiler offers the following options:

- **-march=cell | celledp**
  Selects between the CBEA and the PowerXCell architecture, as well as its registers, mnemonics, and instruction scheduling parameters.

- **-mtune=cell | celledp**
  Tunes the generated code for either the Cell/B.E. or PowerXCell architecture. It mostly affects the instruction scheduling strategy.

- **-mfloat=accurate | fast**
  Selects whether to use the fast fused-multiply operations for floating point operations or to enable calls to library routines that implement more precise operations.

- **-mdouble=accurate | fast**
  Selects whether to use the fast fused-multiply operations for floating point operations or to enable calls to library routines that implement more precise operations.

- **-mstdmain**
  Provides a standard argv/argc calling convention for the main SPU function.
-fpic
-mwarn-reloc
-merror-reloc

Generates position independent code and indicates a warning if the resulting code requires load-time relocations.

-msafe-dma

Controls whether load and store instructions have not moved past DMA operations, by compiler optimizations.

-munsafe-dma

Controls whether load and store instructions have not moved past DMA operations, by compiler optimizations.

-ffixed-<reg>
-mfixed-range=<reg>-<reg>

Reserve specific registers for user application.

**Language options**

The GNU GCC compiler offers the following language extensions to provide access to specific Cell/B.E. architectural features, from a programmer's point of view:

- Vectors

  GNU compiler language support offers the vector data type for both PPU and SPU, with support for arithmetic operations. Refer to 4.6.4, “SIMD programming” on page 258, for more details.

- Intrinsics

  The full set of AltiVec and SPU intrinsics are available. Refer to 4.6.2, “SPU instruction set and C/C++ language extensions (intrinsics)” on page 249, for more details.

**Optimization**

The GNU compiler offers mechanisms to optimize code generation specifically to the underlying architecture. For a complete reference of all optimization-related options, consult the GCC manual on the Web at the following address:

http://gcc.gnu.org/onlinedocs/gcc-4.1.2/gcc/

In particular, refer to 3.10 “Options that control optimization” at the following address:

Predefined generic options

The -O family of options offers a “predefined” generic set of optimization options:

**O0 (default)**  No optimization shortest compilation time; best results when debugging.

**-O1 (-O)**  Default optimization moderately increased compilation time.

**-O2**  Heavy optimization significantly increased compilation time; no optimizations with potentially adverse effects.

**-O3**  Optimal execution time can increase code size; might make debugging difficult.

**-Os**  Optimal code size can imply slower execution time than -O3.

By default, the GCC compiler generates completely unoptimized code. To activate optimization, use one of the -O, -O2, or -O3 flags. Usually, -O3 is the best choice. However -O3 activates such optimizations as automatic inlining that can be undesirable in certain circumstances. In such cases, use the -O2 flag. There might be a few cases where source code was heavily optimized by the programmer for the SPU, in which even -O2 generated worse code than just -O. While these cases are rare, when in doubt, test both options.

**Tip:** When in doubt, use the following set of compiler options to generate optimized code for SPE:

```
-O3 -funroll-loops -fmodulo-sched -ftree-vectorize -ffast-math
```

Flags for special optimization passes

GCC supports a number of specific optimization passes that are not implemented by default at any optimization level (-O...). These can be selected manually where appropriate. Some of the following options might be of particular interest on the SPE:

- **-funroll-loops**
  
  “Unroll” loops by duplicating the loop body multiple times. This option can be helpful on the SPE because it reduces the number of branches. However, the option can increase code size.

- **-fmodulo-sched**

  A special scheduling pass (Swing Modulo Scheduling, also known as *software pipelining*) attempts to arrange instructions in loops to minimize pipeline stalls. This option is usually beneficial on SPEs.

- **-ffast-math**

  By using this option, the compiler can optimize floating-point expressions without preserving exact Institute of Electrical and Electronics Engineers
(IEEE) semantics. For example, with this option, the vectorizer can change the order of computation of complex expressions.

**Compiler directives: Function inlining**

Function inlining is an optimization technique where performance is achieved by replacing function calls with their explicit set of instructions (or body). The formal parameters are replaced with arguments.

The benefits of such a technique are to avoid function call overhead and to keep the function as a whole for combined optimization. The disadvantages are an increase in code size and compilation time.

The compiler offers the following choices for function inlining:
- By explicit declaration, with both the “inline” keyword in function declarations (Example 5-5) and by defining C++ member functions inside the class body

```
Example 5-5  Inline keyword usage
...
static inline void swap(int a, int b) {
    int tmp;
    tmp = a;
    a = b;
    b = tmp;
}
```

- Activated by compiler option -finline-functions, where the compiler applies heuristics over the function call, considering size and complexity

**Compiler directives: Others**

See 4.6.3, “Compiler directives” on page 256, for more information about the usage of other available compiler directives such as volatile, aligned, builtin_expect, align_hint, and restrict.

**Auto-vectorization**

The auto-vectorization feature, which is enabled by the -ftree-vectorize switch, automatically detects situations in the source code where loop-over-scalar instructions can be transformed into loop-over-vector instructions. This feature is usually beneficial on the SPE. In cases where the compiler manages to transform a loop that is performance-critical to the overall application, a significant speedup can be observed.
To help the vectorizer detect loops that are safe to transform, you must follow some general rules when writing loops:

- Use countable loops (known number of iterations).
- Avoid function calls or "break"/"continue" in the loop.
- Avoid aliasing problems by using the C99 "restrict" keyword where appropriate.
- Keep memory access patterns simple.
- Operate on properly aligned memory addresses whenever possible.

If your code has loops that you think should be vectorized, but are not, you can use the `-ftree-vectorizer-verbose=[X]` option to determine why this occurs. X=1 is the least amount of output, and X=6 yields the largest amount of output.

Refer to 4.6.5, “Auto-SIMDizing by compiler” on page 270, for more information about this topic.

**Profile-directed feedback optimization**

Although considered an advanced optimization technique, by using *profile-directed feedback optimization*, the compiler can tune generated code according to the behavior measured during the execution of trial runs.

To use this approach, perform the following steps:

1. Build the application with the `-fprofile-generate` switch to generate an instrumented executable.
2. Run the generated instrumented application on a sample input data set, which results in a profile data file.
3. Use the `-fprofile-use` switch (instead of `-fprofile-generate`). The compiler incorporates feedback from the profile that is run to generate an optimized final executable.

Figure 5-4 on page 339 illustrates the profile-directed feedback process.
5.2.2 Compilers: xlc

IBM XL C/C++ for Multicore Acceleration for Linux is an advanced, high-performance cross-compiler that is tuned for the CBEA. The XL C/C++ compiler, which is hosted on an x86, IBM PowerPC technology-based system, or a BladeCenter QS21, generates code for the PPU or SPU. The compiler requires the GCC toolchain for the CBEA, which provides tools for cross-assembling and cross-linking applications for both the PPE and SPE.

For full documentation regarding the IBM XL C/C++ compiler, refer to the XL C/C++ Library at the following address:


Optimization
The IBM XL C/C++ introduces several innovations, especially with regard to the optimization options. We discuss the general concepts that are involved and provide some useful tips.
The XL compiler also offers the “predefined” optimization options as shown in Figure 5-5.

![Optimization Levels](image)

**Figure 5-5   XL Optimization levels**

**Levels 2 and 3**
- The -O2 level brings comprehensive low-level optimizations while keeping partial support for debugging:
  - Global assignment of user variables to registers
  - Strength reduction and effective usage of addressing modes
  - Elimination of unused or redundant code
  - Movement of invariant code out of loops
  - Scheduling of instructions for the target machine
  - Some loop unrolling and pipelining
  - Visible externals and parameter registers at procedure boundaries
  - Additional program points for storage visibility created by the Snapshot™ pragma/directive
  - Parameters forced to memory on entry, by the -qkeepparam option, so that they can be visible in a stack trace
The -O3 level has extensive optimization, but might introduce precision trade-offs:

- Deeper inner loop unrolling
- Loop nest optimizations, such as unroll-and-jam and interchange (-qhot subset)
- Better loop scheduling
- Additional optimizations allowed by -qnostrict
- Widened optimization scope (typically the whole procedure)
- No implicit memory usage limits (-qmaxmem=-1)
- Reordering of floating point computations
- Reordering or elimination of possible exceptions, for example divide by zero, overflow

To achieve the most of the level 2 and 3 optimizations, consider the following guidance:

- Ensure that your code is standard-compliant and, if possible, test and debug your code without optimization before using -O2.
- With regard to the C code, ensure that pointer use follows type restrictions (generic pointers should be char* or void*), and verify if all shared variables and pointers to same are marked as volatile.
- Try to be uniform, by compiling as much of your code as possible with -O2. If you encounter problems with -O2, consider using -qalias=noansi or -qalias=nostd rather than turning off optimization.
- Use -O3 as much code as possible. If you encounter problems or performance degradations, consider using -qstrict, -qcompact, or -qnohot along with -O3 where necessary. If you still have problems with -O3, switch to -O2 for a subset of files or subroutines, but consider using -qmaxmem=-1, -qnostrict, or both.

**High order transformations (-qhot)**

High order transformations are supported for all languages. The usage is specified as follows:

- qhot=[no]vector | arraypad=[n] | [no]simd
The general optimization gain involves the following areas:

- Transforms (for example, interchange, fusion, and unrolling) loop nests at a high level to optimize the following items:
  - Memory locality (reduced cache or translation look-aside buffer (TLB) misses)
  - Usage of hardware prefetch
  - Loop computation balance (typically ld/st versus float)

- Optionally transforms loops to exploit Mathematical Acceleration Subsystem (MASS) vector library (for example, reciprocal, sqrt, or trig); might result in slightly different rounding

- Optionally introduces array padding under user control; potentially unsafe if not applied uniformly

- Optionally transforms loops to exploit the Vector/SIMD Multimedia Extension (VMX) VMX/SIMD unit

The -qhot option is designed to be used with other optimization levels, such as -O2 and -O3, since it has a neutral effect if no optimization opportunities exist.

Sometimes you might encounter a long unacceptable compilation time or performance degradation, which can be solved by the combined use of -qhot=novector, -qstrict, or -qcompact along with -qhot.

Other times you might encounter unacceptably long compilation times or performance degradation, which can be solved by the combined use of -qhot=novector, -qstrict, or -qcompact along with -qhot. As with any optimization option, try disabling them selectively, if needed.

**Link-time optimization (-qipa)**

The XL compiler also offers a “link-time” optimization option:

```
-qipa[=level=n | inline= | fine tuning]
```

The link-time optimization can be enabled per compile unit (compile step) or on the whole program (compile and link), where it expands its reach to the whole final artifact (executable or library).

The following options can be explored by this feature:

```
level=0 Program partitioning and simple inter procedural optimization
level=1 Inlining and global data mapping
level=2 Global alias analysis, specialization, inter procedural data flow
```
**inline** Precise user control of inlining

**fine tuning** Specify library code behavior, tune program partitioning, or read commands from a file

Although `-ipa` works when building executables or shared libraries, make sure that you compile main and exported functions with `-qipa`. Again, try to apply this option as much as possible.

**Levels 4 and 5**
Optimization levels 4 (-O4) and 5 (-O5) automatically apply all previous optimization level techniques (-O3). Additionally, it includes its own “packages” options:

- `-qhot`
- `-qipa`
- `-qarch=auto`
- `-qtune=auto`
- `-qcach=auto`
- (In -O5 only) `-qipa=level=2`

**Vectorization (VMX and SIMD)**
The XL compiler support two modes for exploitation for the vector features in the CBEA:

- **User driven**
  The code is explicitly ported to use vector types and intrinsics, as well as alignment constrains.

- **Automatic Vectorization (SIMDization)**
  The compiler tries to automatically identify parallel operations across the scalar code and generates vector versions of them. The compiler also performs all necessary transformations to resolve any alignment constrains. This mode requires, at least, optimization level -O3 `-qhot`.

Although the compiler does a through analysis to produce the best fit auto-vectorized code, still the programmer can influence the overall process, making it more efficient. Consider the following more relevant tips:

- **Loop structure**
  - Inline function calls inside innermost loops
  - Automatically (-O5 more aggressive, use inline pragma/directives)
Data alignment
- Align data on 16-byte boundaries:
  __attribute__((aligned(16))
- Describe pointer alignment, which can be placed anywhere in the code, preferably close to the loop:
  _alignx(16, pointer)
- Use -O5, which enables inter-procedural alignment analysis

Pointer aliasing
- Refine pointer aliasing:
  #pragma disjoint(*p, *q) or restrict keyword

Obviously, if you already manually unrolled any of the loops, it becomes more difficult for the SIMDization process. Even in that case, you can manually instruct the compiler to skip those loops:

#pragma nosimd (right before the innermost loop)

5.2.3 The build environment

In /opt/cell/sdk/buildutils, top-level makefiles control the build environment for all of the examples. Most of the directories in the libraries and examples contain a makefile for that directory and everything in it. All of the examples have their own makefile, but the common definitions are in the top-level makefiles. Table 5-1 shows examples of configurable makefile features. The build environment makefiles are documented in /opt/cell/sdk/buildutils/README_build_env.txt.

Table 5-1 Makefile example configurable features

<table>
<thead>
<tr>
<th>Environment variable</th>
<th>Description</th>
<th>Example value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CFLAGS_[gcc</td>
<td>xlc]</td>
<td>Passes additional compilation flags to the compiler</td>
</tr>
<tr>
<td>LDFLAGS_[gcc</td>
<td>xlc]</td>
<td>Passes additional linker flags to the linker</td>
</tr>
<tr>
<td>CC_OPT_LEVEL</td>
<td>Overrides specifically the compiler optimization level</td>
<td>-O0</td>
</tr>
<tr>
<td>INCLUDE</td>
<td>Additional include paths to the compiler</td>
<td>-I/usr/spu/lib</td>
</tr>
<tr>
<td>IMPORTS</td>
<td>Additional libraries to be imported by the linker</td>
<td>-lnuma -lpthread</td>
</tr>
</tbody>
</table>
Changing the environment
The environment variables in the /opt/cell/sdk/buildutils/make.* files are used to determine which compiler is used to build the examples. The /opt/cell/sdk/buildutils/cellsdk_select_compiler script can be used to switch the compiler. This command has the following syntax:

```
/opt/cell/sdk/buildutils/cellsdk_select_compiler [xlc | gcc]
```

In this command, the xlc flag selects the XL C/C++ compiler and the gcc flag selects the GCC compiler. The default, if unspecified, is to compile the examples with the GCC compiler. After selecting a particular compiler, that same compiler is used for all future builds, unless it is specifically overwritten by the shell environment variables SPU_COMPILER, PPU_COMPILER, PPU32_COMPILER, or PPU64_COMPILER.

5.3 Debugger

The debugger is a tool to help find and remove problems in your code. In addition to fixing problems, the debugger can help you understand the program, because it typically gives you memory and registers contexts, stack call traces, and step-by-step execution. Figure 5-6 highlights the debug stage of the process.

![Figure 5-6 Debug](image)

5.3.1 Debugger: gdb

GDB is the standard command-line debugger that is available as part of the GNU development environment. GDB has been modified to allow debugging in a Cell/B.E. processor environment. Debugging in a Cell/B.E. processor environment is different from debugging in a multithreaded environment, because threads can run either on the PPE or on the SPE.
Three versions of GDB can be installed on a BladeCenter QS21 blade server:

- `gdb`, which is installed with the Linux operating system for debugging PowerPC applications. You should not use this debugger for Cell/B.E. applications.
- `ppu-gdb`, which is for debugging PPE code or for debugging combined PPE and SPE code. This is the combined debugger.
- `spu-gdb`, which is for debugging SPE code only. This is the standalone debugger.

In this section, we describe how to debug Cell/B.E. software by using the new and extended features of the GDB that is supplied with SDK 3.0.

**Setup and considerations**

The linker embeds all the symbolic and additional information that is required for the SPE binary within the PPE binary, so that it is available for the debugger to access when the program runs. Use the `-g` option when compiling both SPE and PPE code with either the GCC or XLC compiler. The `-g` option adds debugging information to the binary, which then enables GDB to look up symbols and show the symbolic information.

When you use the top-level makefiles of the SDK, you can specify the `-g` option on compilation commands by setting the `CC_OPT_LVL` makefile variable to `-g`.

**Debugging PPE code**

There are several ways to debug programs that are designed for the Cell/B.E. processor. If you have access to Cell/B.E. hardware, you can debug directly by using `ppu-gdb`. You can also run the application under `ppu-gdb` inside the simulator. Alternatively, you can debug remotely.

Regardless of the method that you choose, after you start the application under `ppu-gdb`, you can use the standard GDB commands that are available to debug the application. For more information, refer to the GDB user manual, which is available from the GNU Web site at the following address:

http://www.gnu.org/software/gdb/gdb.html

**Debugging SPE code**

Standalone SPE programs or spulets are self-contained applications that run entirely on the SPE. Use `spu-gdb` to launch and debug standalone SPE programs in the same way as you use `ppu-gdb` on PPE programs.
**Debugging multi-threaded code**

Typically a simple program contains only one thread. For example, a PPU “hello world” program is run in a process with a single thread and the GDB attaches to that single thread.

On many operating systems, a single program can have more than one thread. With the ppu-gdb program, you can debug programs with one or more threads. The debugger shows all threads while your program runs, but whenever the debugger runs a debugging command, the user interface shows the single thread involved. This thread is called the *current thread*. Debugging commands always show program information from the point of view of the current thread.

For more information about GDB support for debugging multithreaded programs, see “Debugging programs with multiple threads” and “Stopping and starting multi-thread programs” in the GDB user manual, which is available from the GNU Web site at the following address:

http://www.gnu.org/software/gdb/gdb.html

The `info threads` command shows the set of threads that are active for the program. The `thread` command can be used to select the current thread for debugging.

**Debugging architecture**

On the Cell/B.E. processor, a thread can run on either the PPE or on an SPE at any given point in time. All threads, both the main thread of execution and secondary threads that are started by using the pthread library, start execution on the PPE. Execution can switch from the PPE to an SPE when a thread executes the `spe_context_run` function. See *SPE Runtime Management Library Version 2.2*, SC33-8334-01, which is available on the Web at the following address:


Conversely, a thread that currently executes on an SPE can switch to use the PPE when executing a library routine that is implemented via the PPE-assisted call mechanism. See the *Cell BE Linux Reference Implementation Application Binary Interface Specification* document for details.

When you choose a thread to debug, the debugger automatically detects the architecture that the thread is currently running on. If the thread is currently running on the PPE, the debugger uses the PowerPC architecture. If the thread is currently running on an SPE, the debugger uses the SPE architecture. A thread that is currently executing code on an SPE can also be referred to as an *SPE thread*. 
To see which architecture the debugger is using, use the following command:

```
show architecture
```

**Using scheduler-locking**

Scheduler-locking is a feature of GDB that simplifies multithread debugging by enabling you to control the behavior of multiple threads when you single-step through a thread. By default, scheduler-locking is off, which is the recommended setting.

In the default mode where scheduler-locking is off, single-stepping through one particular thread does not stop other threads of the application from running, but allows them to continue to execute. This applies to both threads that execute on the PPE and SPE. This is not always what you expect or want when debugging multithreaded applications. The threads that execute in the background can affect global application state asynchronously in ways that can make it difficult to reliably reproduce the problem that you are debugging. If this is a concern, you can turn on scheduler-locking. In active mode, all other threads remain stopped while you debug one particular thread. Another option is to set scheduler-locking to *step*, which stops other threads while you are single-stepping the current thread, but lets them execute while the current thread is freely running.

If scheduler-locking is turned on, there is the potential for deadlocking where one or more threads cannot continue to run. Consider, for example, an application that consists of multiple SPE threads that communicate with each other through a mailbox. Let us assume that you single-step one thread across an instruction that reads from the mailbox, and that mailbox happens to be empty at the moment. This instruction (and thus the debugging session) blocks until another thread writes a message to the mailbox. However, if scheduler-locking is on, the other thread remains stopped by the debugger because you are single-stepping.

In this situation, none of the threads can continue, and the whole program stalls indefinitely. This situation cannot occur when scheduler-locking is off, because all other threads continue to run while the first thread is single-stepped. Ensure that you enable scheduler-locking only for applications where such deadlocks cannot occur.

There are situations where you can safely set scheduler-locking on, but you should do so only when you are sure there are no deadlocks.

The **scheduler-locking** command has the following syntax:

```
set scheduler-locking <mode>
```
In this command, \textit{mode} has one of the following values:

- off
- on
- step

You can check the scheduler-locking mode with the following command:

\texttt{show scheduler-locking}

\textbf{Threads and per-frame architecture}

The most significant design change introduced by the combined debugger is the switch from a per-thread architecture selection to a per-frame selection. The new combined debugger for SDK 3.0 eliminates the GDB notion of a “current architecture.” It removes the global notion of a current architecture per thread. In practice, the architecture depends on the current frame. The frame is another fundamental GDB notion. It refers to a particular level in the function call sequence (stack back-trace) on a specific thread.

The architecture selection per-frame notion allows the Cell/B.E. back ends to represent the flow of control that switches from PPE code to SPE code and back. The full back-trace can represent the following program state, for example:

1. (Current frame) PPE libc printf routine
2. PPE libspe code implementing a PPE-assisted call
3. SPE newlib code that issued the PPE-assisted call
4. SPE user application code that called printf
5. SPE main
6. PPE libspe code that started SPE execution
7. PPE user application code that called \texttt{spe_context_run}
8. PPE main

Therefore, any thread of a combined Cell/B.E. application executes either on the PPE or an SPE at the time that the debugger interrupted execution of the process that is currently being debugged. This determines the main architecture that GDB uses when examining the thread. However, during the execution history of that thread, execution may have switched between architectures one or multiple times. When looking at the thread’s stack backtrace (using the \texttt{backtrace} command), the debugger reflects those switches. It shows stack frames that belong to both the PPE and SPE architectures.

When you choose a particular stack frame to examine by using the \texttt{frame}, \texttt{up}, or \texttt{down} commands, the debugger switches its notion of the current architecture as appropriate for the selected frame. For example, if you use the \texttt{info registers} command to look at the selected frame’s register contents, the debugger shows the SPE register set if the selected frame belongs to an SPE context. It shows the PPE register set if the selected frame belongs to PPE code.
Breakpoints

Generally speaking, you can use the same procedures to debug code for the Cell/B.E. processor as you would for PowerPC code. However, some existing features of GDB and one new command can help you to debug in the Cell/B.E. processor multithreaded environment as described in the following sections.

Setting pending breakpoints

Breakpoints stop programs from running when a certain location is reached. You set breakpoints with the `break` command, followed by the line number, function name, or exact address in the program.

You can use breakpoints for both PPE and SPE portions of the code. In some instances, however, GDB must defer insertion of a breakpoint because the code that contains the breakpoint location has not yet been loaded into memory. This occurs when you want to set the breakpoint for code that is dynamically loaded later in the program. If ppu-gdb cannot find the location of the breakpoint, it sets the breakpoint to `pending`. When the code is loaded, the breakpoint is inserted and the pending breakpoint deleted. You can use the `set breakpoint` command to control the behavior of GDB when it determines that the code for a breakpoint location is not loaded into memory.

The breakpoint pending command uses the following syntax:

```
set breakpoint pending <on off auto>
```

In this command, note the following explanations:

- `on` specifies that GDB should set a pending breakpoint if the code for the breakpoint location is not loaded.
- `off` specifies that GDB should not create pending breakpoints and `break` commands for a breakpoint location that is not loaded result in an error.
- `auto` specifies that GDB should prompt the user to determine if a pending breakpoint should be set if the code for the breakpoint location is not loaded. This is the default behavior.

Multiple defined symbols

When debugging a combined Cell/B.E. application that consists of a PPE program and more SPE programs, multiple definitions of a global function or variable with the same name can exist. For example, both the PPE and SPE programs define a global main function. If you run the same SPE executable simultaneously within multiple SPE contexts, all its global symbols show multiple instances of definition. This might cause problems when attempting to refer to a specific definition from the GDB command line, for example when setting a breakpoint. It is not possible to choose the desired instance of the function or variable definition in all cases.
To catch the most common usage cases, GDB uses the following rules when looking up a global symbol:

- If the command is issued while currently debugging PPE code, the debugger first attempts to look up a definition in the PPE executable. If none is found, the debugger searches all currently loaded SPE executables and uses the first definition of a symbol with the given name it finds.

- When referring to a global symbol from the command line while currently debugging SPE context, the debugger first attempts to look up a definition in that SPE context. If none is found there, the debugger continues to search the PPE executable and all other currently loaded SPE executables and uses the first matching definition.

**Architecture specific commands**
In addition to promoting changes to some of the common gdb debugger commands behavior, the Cell/B.E. SDK 3.0 introduces a set of new commands to better accommodate both PPE and SPE code needs.

*set spu stop-on-load*

The new *set spu stop-on-load* command stops each thread before it starts running on the SPE. While *set spu stop-on-load* is in effect, the debugger automatically sets a temporary breakpoint on the main function of each new SPE thread immediately after it is loaded. You can use the *set spu stop-on-load* command to do this instead of simply issuing a *break main* command, because the latter is always interpreted to set a breakpoint on the main function of the PPE executable.

**Note:** The *set spu stop-on-load* command has no effect in the SPU standalone debugger spu-gdb. To let an SPU standalone program proceed to its “main” function, you can use the *start* command in spu-gdb.

The *spu stop-on-load* command has the following syntax, where mode is either *on* or *off*:

```
set spu stop-on-load <mode>
```

To check the status of the *spu stop-on-load*, use the following command:

```
show spu stop-on-load
```
**Info SPU commands**

In addition to the `set spu stop-on-load` command, the ppu-gdb and spu-gdb programs offer an extended set of the standard GDB `info` commands:

- `info spu event`
- `info spu signal`
- `info spu mailbox`
- `info spu dma`
- `info spu proxydma`

If you are working in GDB, you can access help for these new commands. To access help, type the `help info spu` command, followed by the `info spu` subcommand name, which displays full documentation. Command name abbreviations are allowed if they are unambiguous.

**info spu event**

The `info spu event` command displays the SPE event facility status. Example 5-6 shows the output.

*Example 5-6   Output of the info spu event command*

```
(gdb) info spu event
Event Status 0x00000000
Event Mask   0x00000000
```

**info spu signal**

The `info spu signal` command displays the SPE signal notification facility status. Example 5-7 shows the output.

*Example 5-7   Output of the info spu signal command*

```
(gdb) info spu signal
Signal 1 not pending (Type Or)
Signal 2 control word 0x30000001 (Type Or)
```

**info spu mailbox**

The `info spu mailbox` command displays the SPE mailbox facility status. Only pending entries are shown. Entries are displayed in the order of processing. That is, the first data element in the list is the element that is returned on the next read from the mailbox. Example 5-8 on page 353 shows the output.
Example 5-8  Output of the info spu mailbox command

```
(gdb) info spu mailbox
SPU Outbound Mailbox
0x00000000
SPU Outbound Interrupt Mailbox
0x00000000
SPU Inbound Mailbox
0x00000000
0x00000000
0x00000000
0x00000000
```

### info spu dma

The **info spu dma** command displays the MFC DMA status. For each pending DMA command, the opcode, tag, and class IDs are shown, followed by the current effective address, local store address, and transfer size (updated as the command is processed). For commands that use a DMA list, the local store address and size of the list are shown. The “E” column indicates commands flagged as erroneous by the MFC. Figure 5-7 shows the output.

![Output of the info spu dma command](image)

*Figure 5-7  Output of the info spu dma command*
**info spu proxydma**

The `info spu proxydma` command displays the MFC Proxy-DMA status. Figure 5-8 shows the output.

![Output of the info spu proxydma command](image)

---

**5.4 IBM Full System Simulator**

The IBM Full System Simulator is a software application that emulates the behavior of a full system that contains a Cell/B.E. processor. You can start a Linux operating system on the simulator and run applications on the simulated operating system. The simulator also supports the loading and running of statically-linked executable programs and standalone tests without an underlying operating system. Figure 5-9 shows the simulator flow.

![Simulator flow](image)

---
The simulator infrastructure is designed for modeling processor and system-level architecture at levels of abstraction, which vary from functional to performance simulation models with a number of hybrid fidelity points in between:

- **Functional-only simulation**
  
  This simulation models the program-visible effects of instructions without modeling the time it takes to run these instructions. Functional-only simulation assumes that each instruction can be run in a constant number of cycles. Memory access is synchronous and is also performed in a constant number of cycles. This simulation model is useful for software development and debugging when a precise measure of execution time is not significant. Functional simulation proceeds much more rapidly than performance simulation, and therefore, is useful for fast-forwarding to a specific point of interest.

- **Performance simulation**
  
  For system and application performance analysis, the simulator provides performance simulation (also referred to as *timing simulation*). A performance simulation model represents internal policies and mechanisms for system components, such as arbiters, queues, and pipelines. Operation latencies are modeled dynamically to account for both processing time and resource constraints. Performance simulation models have been correlated against hardware or other references to acceptable levels of tolerance.

  The simulator for the Cell/B.E. processor provides a cycle-accurate SPU core model that can be used for performance analysis of computational-intense applications. The simulator for SDK 3.0 provides additional support for performance simulation, which is described in the *IBM Full-System Simulator Users Guide and Performance Analysis* document.

  The simulator can also be configured to fast forward the simulation, by using a functional model, to a specific point of interest in the application and to switch to a timing-accurate mode to conduct performance studies. This means that various types of operational details can be gathered to help you understand real-world hardware and software systems.

  See the `/opt/ibm/systemsim-cell/doc` subdirectory for complete documentation including the *IBM Full-System Simulator Users Guide and Performance Analysis* document.
5.4.1 Setting up and bringing up the simulator

To verify that the simulator is operating correctly and then to run it, enter the following commands:

```bash
export PATH=/opt/ibm/systemsim-cell/bin:$PATH
systemsim -g
```

The systemsim script found in the simulators bin directory launches the simulator. The -g parameter starts the graphical user interface (GUI).

You can use the GUI of the simulator to gain a better understanding of the CBEA. For example, the simulator shows two sets of the PPE state. This is because the PPE processor core is dual-threaded, and each thread has its own registers and context. You can also look at the state of the SPEs, including the state of their MFC.

Access to the simulator image

By default, the simulator does not write changes back to the simulator system root (sysroot) image. This means that the simulator always begins in the same initial state of the sysroot image. When necessary, you can modify the simulator configuration, so that any file changes made by the simulated system to the sysroot image are stored in the sysroot disk file. This way, they are available to subsequent simulator sessions.

To specify that you want to update the sysroot image file with any changes made in the simulator session, change the newcow parameter on the `mysim bogus disk init` command in `.systemsim.tcl` to `rw` (specifying read/write access) and remove the last two parameters. The following changed line is from `.systemsim.tcl`:

```tcl
mysim bogus disk init 0 $sysrootfile rw
```

When running the simulator with read/write access to the sysroot image file, ensure that the file system in the sysroot image file is not corrupted by incomplete writes or a premature shutdown of the Linux operating system running in the simulator. In particular, ensure that Linux writes any cached data to the file system before exiting the simulator. You do this by typing the following command in the Linux console window just before you exit the simulator:

```bash
sync ; sync
```

Selecting the architecture

Many of the tools provided in SDK 3.0 support multiple implementations of the CBEA. These include the Cell/B.E. processor and a future processor. This future processor is a CBEA-compliant processor with a fully pipelined, enhanced double-precision SPU.
The processor supports five optional instructions to the SPU ISA:

- DFCEQ
- DFCGT
- DFCMEQ
- DFCMEQ
- DFCMGT

Detailed documentation for these instructions is provided in version 1.2 (or later) of the SPU ISA specification. The future processor also supports improved issue and latency for all double-precision instructions.

The simulator also supports simulation of the future processor. The simulator installation provides a tcl run script to configure it for such simulation. For example, the following sequence of commands starts the simulator that is configured for the future processor with a GUI:

```bash
export PATH=$PATH:/opt/ibm/systemsim-cell/bin
systemsim -g -f config_edp_smp.tcl
```

### 5.4.2 Operating the simulator GUI

The simulator GUI offers a visual display of the state of the simulated system, including the PPE and the eight SPEs. You can view the values of the registers, memory, and channels, as well as the performance statistics. The GUI also offers an alternate method of interacting with the simulator.

The main GUI window has two basic areas:

- The vertical panel on the left
- The rows of buttons on the right

The vertical panel represents the simulated system and its components. The rows of buttons are used to control the simulator.

To start the GUI from the Linux run directory, enter the following command:

```bash
PATH=/opt/ibm/systemsim-cell/bin:$PATH; systemsim -g
```

The simulator then configures the simulator as a Cell/B.E. system and shows the main GUI window, which is labeled with the name of the application program. When the GUI window first is displayed, click the Go button to start the Linux operating system.
The simulation panel
When the main GUI window first opens, the vertical panel contains a single folder labeled *mysim*. To see the contents of *mysim*, click the plus sign (+) in front of the folder icon.

When the folder is expanded, you see the following contents:
- A PPE (labeled PPE0:0:0 and PPE0:0:1)
- The two threads of the PPE
- Eight SPEs (SPE0... SPE7)

The folders that represent the processors can be further expanded to show the viewable objects, as well as the options and actions that are available.

**PPE components**
Five PPE components are visible in the expanded PPE folder:
- PCTrack
- PCCCore
- GPRegs
- FPRegs
- PCAddressing

You can view the general-purpose registers (GPRs) and the floating-point registers (FPRs) separately by double-clicking the GPRegs and the FPRegs folders respectively. As data changes in the simulated registers, the data in the windows is updated, and registers that have changed state are highlighted.

The PPE Core window (PPCCore) shows the contents of all the registers of the PPE, including the VMX registers.

**SPE components**
The SPE folders (SPE0 ... SPE7) each have ten subitems. Five of the subitems represent windows that show data in the registers, channels, and memory:
- SPUTrack
- SPUCore
- SPEChannel
- LS_Stats
- SPUMemory

Two of the subitems represent windows that show state information about the MFC:
- MFC
- MFC_XLate
The last three subitems represent actions to perform on the SPE:

- SPUStats
- Model
- Load-Exec

The last three items in an SPE folder represent actions to perform, with respect to the associated SPE. The first of these is SPUStats. When the system is stopped and you double-click this item, the simulator displays program performance statistics in its own window. These statistics are collected only when Model is set to pipeline mode.

The next item in the SPE folder has one of the following labels:

- Model: instruction
- Model: pipeline
- Model: fast

The label indicates whether the simulation is in one of the following modes:

- Instruction mode for checking and debugging the functionality of a program
- Pipeline mode for collecting performance statistics on the program
- Fast mode for fast functional simulation only

You can toggle the model by double-clicking the item. You can use the Perf Models button on the GUI to display a menu for setting the simulator model modes of all of the SPEs simultaneously.

The last item in the SPE folder, Load-Exec, is used for loading an executable onto an SPE. When you double-click this item, a file-browsing window opens in which you can find and select the executable file to load.

**Simulation control buttons**

On the right side of the GUI window are five rows of buttons, which are used to manipulate the simulation process. The buttons include the following options:

- Advance Cycle

This option advances the simulation by a set number of cycles. The default is one cycle, but it can be changed by entering an integer value in the text box above the buttons, or by moving the slider next to the text box. From the drop-down menu at the top of the GUI, the user can select the time domain for cycle stepping. The time units to use for cycles are expressed in terms of various system components. The simulation must be stopped for this button to work. If the simulation is not stopped, the button is inactive.
**Go**
This option starts or continues the simulation. In the SDK\u2019s simulator, the first time the Go button is clicked, it initiates the Linux boot process. In general, the action of the Go button is determined by the startup tcl file that is in the directory from which the simulator is started.

**Stop**
This option pauses the simulation.

**Service GDB**
This option allows the external gdb debugger to attach to the running program. This button is also inactive while the simulation is running.

**Triggers/Breakpoints**
This option displays a window that shows the current triggers and breakpoints.

**Update GUI**
This option refreshes all of the GUI windows. By default, the GUI windows are updated automatically every four seconds. Click this button to force an update.

**Debug Controls**
This option displays a window of the available debug controls from which you can select the ones that should be active. When this button is enabled, corresponding information messages are displayed.

**Options**
This option displays a window in which you can select fonts for the GUI display. On a separate tab, you can enter the gdb debugger port.

**Emitters**
This option displays a window with the defined emitters, with separate tabs for writers and readers.

**Fast Mode**
This option toggles fast mode on and off. Fast mode accelerates the execution of the PPE at the expense of disabling certain system-analysis features. It is useful for quickly advancing the simulation to a point of interest. When fast mode is on, the button is displayed as unavailable. Otherwise it is displayed as active available. Fast mode can also be enabled by using the `mysim fast on` command and disabled by using the `mysim fast off` command.
Perf Models
This option displays a window in which various performance models can be selected for the various system simulator components. It provides a convenient means to set each SPU\textsuperscript{2} simulation mode to either cycle accurate pipeline mode, instruction mode, fast functional-only mode. The same capabilities are available by using the Model:instruction, Model:pipeline, Model:fast toggle menu subitem under each SPE in the tree menu at the left of the main control panel.

SPE Visualization
This option plots histograms of SPU and DMA event counts. The counts are sampled at user-defined intervals and are continuously displayed. Two modes of display are provided:
- A \texttt{scroll} view, which tracks only the most recent time segment
- A \texttt{compress} view, which accumulates samples to provide an overview of the event counts during the time elapsed

Users can view collected data in either detail or summary panels:
- The detailed, single-SPE panel tracks SPU pipeline phenomena (such as stalls, instructions executed by type, and issue events) and DMA transaction counts by type (such as gets, puts, atomics, and so forth).
- The summary panel tracks all eight SPEs for the Cell/B.E. processor, with each plot showing a subset of the detailed event count data that is available.

Process-Tree and Process-Tree-Stats
This option requires OS kernel hooks that allow the simulator to display process information. This feature is currently not provided in the SDK kernel.

SPU Modes
This option provides a convenient means to set each SPU\textsuperscript{2} simulation mode to either cycle accurate pipeline mode or fast functional-only mode. The same capabilities are available using the Model:instruction or Model:pipeline toggle menu subitem under each SPE in the tree menu at the left of the main control panel.

Event Log
This option enables a set of predefined triggers to start collecting the log information. The window provides a set of buttons that can be used to set the marker cycle to a point in the process.

Exit
This option exits the simulator and closes the GUI window.
5.5 IBM Multicore Acceleration Integrated Development Environment

The IBM SDK for Multicore Acceleration Integrated Development Environment (Cell/B.E. IDE) is built upon the Eclipse and C Development Tools (CDT) platform. It integrates the Cell/B.E. GNU toolchain, compilers, IBM Full-System Simulator for the Cell/B.E. system, and other development components in order to provide a comprehensive, user-friendly development platform that simplifies Cell/B.E. development. The SDK supports the entire development process, as shown in Figure 5-10.

![Figure 5-10](image)

*Figure 5-10  IDE supports all parts of the process*

This Cell/B.E. IDE includes the following key features:

- A C/C++ editor that supports syntax highlighting; a customizable template; and an outline window view for procedures, variables, declarations, and functions that appear in source code
- A rich visual interface for the PPE and SPE GDB
- Seamless integration of the simulator into Eclipse
- Automatic builder, performance tools, and several other enhancements

The Cell/B.E. IDE offers developers a complete solution for developing an application. The IDE is capable of managing all artifacts that are involved in the development, as well as deploying and testing them on the target environment. Typically, the developer goes from projects creation, including multiple build configurations, target environment setup, and application launching and debugging.

5.5.1 Step 1: Setting up a project

The underlying Eclipse Framework architecture offers the concept of projects as units of agglomeration for your application artifacts. A project is responsible for holding a file system structure and binding your application code with build, launch, and debug configurations, as well as non-source code artifacts.
Defining projects
The Cell/B.E. IDE leverages the Eclipse CDT framework, which is the tooling support for developing C/C++ applications. In addition to the CDT framework, you can choose whether you manage the build structure yourself or you have Eclipse automatically generate it for you. This choice is the difference respectively between the Standard Make and Managed Make options for project creation (Table 5-2).

Table 5-2  Project management options

<table>
<thead>
<tr>
<th>Project management style</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard Make C/C++</td>
<td>You are required to provide a makefile.</td>
</tr>
<tr>
<td>Managed Make C/C++</td>
<td>Eclipse auto-creates and manages the makefiles for you.</td>
</tr>
</tbody>
</table>

From the menu bar, you select File → New → Project. In the New Project Wizard window that opens (Figure 5-11), you can select Standard Make or Managed Make projects under both the C and C++ project groups.
Creating projects
After you define the style of projects that suits your needs, give your project a name. Then in the Select a type of project window (Figure 5-12), choose a project type from among the available project types for Cell/B.E. development.

Figure 5-12  Project types
Table 5-3 describes each of the available project types.

Table 5-3  Project type options

<table>
<thead>
<tr>
<th>Project type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell PPU Executable</td>
<td>Creates a PPU executable binary. This project has the capability of referencing any other SPU project binary, in order to produce a Cell/B.E. combined binary.</td>
</tr>
<tr>
<td>Cell PPU Shared Library</td>
<td>Creates a PPU shared library binary. This project has the capability of referencing any other SPU project binary, in order to produce a Cell/B.E. combined library.</td>
</tr>
<tr>
<td>Cell PPU Static Library</td>
<td>Creates a PPU static library binary. This project has the capability of referencing any other SPU project binary, in order to produce a Cell/B.E. combined library.</td>
</tr>
<tr>
<td>Cell SPU Executable</td>
<td>Creates an SPU binary. The resulting binary can be executed as a spulet or embedded in a PPU binary.</td>
</tr>
<tr>
<td>Cell SPU Static Library</td>
<td>Creates an SPU static library. The resulting library can be linked together with other SPU libraries and be embedded in an PPU binary.</td>
</tr>
</tbody>
</table>

**Project configuration**
The newly created project should be displayed in the C/C++ view, on the left side of the window. Next configure the project’s build options.

**Standard Make:** Choosing the Standard Make style of project management implies that you are responsible for maintaining and updating your makefiles. Eclipse only offers a thin run wrapper, where you can define the relevant targets within your makefile. Therefore, when you click **Build** in Eclipse, it invokes the desired makefile target, instead of the default of **all**.
Select the desired project and right-click the **Properties** option. In the properties window, select **C/C++ Build** on the left pane of the window (Figure 5-13).

![Figure 5-13  C/C++ Build options](image)

The C/C++ Build options carry all of the necessary build and compile configuration entry points, so that we can customize the whole process. Each tool that is part of the toolchain has its configuration entry point on the Tools Settings tab. After you alter or add any of the values, click **Apply** and then **OK** to trigger the automatic makefile generation process of the IDE, so that you project's build is updated immediately.
5.5.2 Step 2: Choosing the target environments with remote tools

Now that the projects are created and properly configured, we must first create and start a programming environment before we can test the program. The Cell/B.E. IDE integrates the IBM Full System Simulator for the Cell/B.E. processor and Cell/B.E. blades into Eclipse, so that you are only a few clicks away from testing your application on a cell environment.

**Simulator integration**

In the Cell Environments view at the bottom, right-click **Local Cell Simulator** and select **Create**.

The Local Cell Simulator properties window (Figure 5-14) opens, in which you can configure the simulator to meet any specific needs that you might have. You can modify an existing cell environment's configuration at any time (as long as it's not running). To modify the configuration, right-click the environment and select **Edit**. Enter a name for this simulator, such as **My Local Cell Simulator**, and then click **Finish**.

![Figure 5-14 Simulator environment window](image-url)
Cell/B.E. blade integration
In the Cell Environments view at the bottom, right-click Cell Box and select Create.

The Cell Box properties window (Figure 5-15) opens in which you can configure remote access to your Cell/B.E. blade to meet any specific needs. You can modify an existing cell environment's configuration at any time (as long as it is not running). Right-click the environment and select Edit. Enter a name for this configuration, such as My Cell Blade, and then click Finish.

Figure 5-15  Cell blade environment window

Starting the environments
At least one of the environment configurations must be started in order to access the launch configuration options. Highlight the desired environment configuration and click the green arrow button in the left corner of Cell Environments view. By
clicking this button, you activate the connection between the chosen environment and the IDE.

5.5.3 Step 3: Debugging the application

Next, a C/C++ cell application launch configuration must be created and configured for the application debugging.

Creating the launch configuration
Select Run → Debug... In the left pane of the Debug window, right-click C/C++ Cell Target Application and select New.

Configuring the launch
In the launch configuration window (Figure 5-16), in the Project field (on the Main tab), specify the project that you are going to debug. You must also specify which application, within the project.

![Launch configurations window](image)

*Figure 5-16  Launch configurations window*

Click the Target tab. On the Target tab (Figure 5-17 on page 370), you can select which remote environment you want to debug your application with. It is possible to select any of the previously configured environments, as long as they are active, that is started.
Click the **Launch** tab. On the Launch tab (Figure 5-18), you can specify any command line arguments and shell commands that must be executed before your application, after your application, or both.
Click the **Synchronize** tab (Figure 5-19), on which you can specify resources, such as input/output files, that must be synchronized with the cell environment’s file system before the application executes, after the application executes, or both. Click **New upload rule** to specify the resource or resources to copy to the cell environment before the application executes. Click **New download rule** to copy the resource or resources back to your local file system after execution. Select the **Upload rules enabled** and **Download rules enabled** boxes respectively after adding any upload or download rules.

![Figure 5-19 Synchronize tab](image)

Configure the debugger parameters. Click the **Debugger** tab (Figure 5-20 on page 372). In the Debugger field, choose **Cell PPU gdbserver**, **Cell SPU gdbserver**, or **Cell/B.E. gdbserver**. To debug only PPU or SPU programs, select **Cell PPU gdbserver** or **Cell SPU gdbserver**, respectively. The **Cell/B.E. gdbserver** option is the combined debugger, which allows for the debugging of PPU and SPU source code in one debug session.
Since this is a remote debugging session, it is important to select the correct remote side debugger (gdbserver) type, according to your application. Click the **Connection** tab (Figure 5-21), on the Debugger page.
If your application is 32 bit, choose the Cell/B.E. 32-bit gdbserver option for the gdbserver Debugger field. Otherwise, if you have a 64-bit application, select the **Cell/B.E. 64bit gdbserver** option.

If there are no pending configuration problems, click the **Apply** button and then the **Debug** button. The IDE switches to the Debug Perspective.

**Debug Perspective**

In the Eclipse Debug Perspective (Figure 5-22), you can use any of the regular debugging features such as breakpoint setting, variables inspection, registers, and memory.

![Debug perspective](image)

*Figure 5-22  Debug perspective*
Additionally, the SPU architecture `info debug` commands (see “Info SPU commands” on page 352) are also available through their respective views. Select **Window → Show View** to locate them as shown in Figure 5-23.

![Figure 5-23  Info SPU available views](image)

**Tip:** As explained in “Threads and per-frame architecture” on page 349”, the `info spu` commands only work when debugging in the SPE architecture. Use the stack frame view of Eclipse, in the upper left corner (under Debug) to precisely determine in which architecture your code is executing.
5.6 Performance tools

The Cell/B.E. SDK 3.0 has a set of performance tools. It is crucial to understand how each one relates to the other, from a time-flow perspective. Figure 5-24 highlights the optimize stage of the process.

Figure 5-24  Optimize stage

5.6.1 Typical performance tuning cycle

The major tasks to achieve a complete development cycle can be organized as follows:

- Develop/port the application for the Cell/B.E. system
  - Programming best practices
  - Knowledge from the architecture

- Compile, link, and debug
  - Compiler (gcc or xlc)
  - Linker (ld)
  - Debugger (gdb)

- Performance tuning
  - OProfile, Performance Debugging Tool (PDT), PDTR, Visual Performance Analyzer (VPA), Post-link Optimization for Linux on Power tool (FDPR-Pro), and cell-perf-counter (CPC)
Figure 5-25 summarizes the Cell/B.E. tools interlock.

The CPC tool

The cell-perf-counter (CPC) tool is for setting up and using the hardware performance counters in the Cell/B.E. processor. With these counters, you can see how many times certain hardware events occur, which is useful if you are analyzing the performance of software running on a Cell/B.E. system.

Hardware events are available from all of the logical units within the Cell/B.E. processor including the following units:

- The PPE
- The SPEs
- The interface bus
- Memory
- I/O controllers

The Cell/B.E. performance monitoring unit (PMU) provides four 32-bit counters, which can also be configured as pairs of 16-bit counters, for counting these events. The CPC tool also uses the hardware sampling capabilities of the
Cell/B.E. PMU. By using this feature, the hardware can collect precise counter data at programmable time intervals. The accumulated data can be used to monitor changes in performance of the Cell/B.E. system over longer periods of time.

**Operation**
The tool offers two modes of execution:

- **Workload mode**
  PMU counters are active only during complete execution of a workload, providing an accurate view of the performance of a single process.

- **System-wide mode**
  PMU counters monitor all processes that run on specified processors for a specified duration.

The results are grouped according to seven logical blocks, PPU, PPSS, SPU, MFC, EIB, MIC, and BEI, where each block has signals (hardware events) organized into groups. The PMU can monitor any number of signals within one group, with a maximum of two signal groups at a time.

**Hardware sampling**
The Cell/B.E. PMU provides a mechanism for the hardware to periodically read the counters and store the results in a hardware buffer. By doing so, the CPC tool can collect a large number of counter samples while greatly reducing the number of calls that the CPC tool must make into the kernel.

The following steps are performed in the hardware sampling mode:

1. Specify the initial counter values and sampling time interval.
2. Record the PMU and reset counter values after each interval.
3. Make samples available in hardware trace-buffer.

As the default behavior, hardware buffers contain the total number of each monitored signal’s hit for the specified interval, which is called **count mode**. In addition to sampling the counters and accumulating them in the buffers, the Cell/B.E. PMU offers other sampling modes:

- **Occurrence mode**
  This mode monitors one or two entire groups of signals, allowing the specifying of any signal within the desired group. It also indicates whether each event occurred at least once during each sampling interval.
Threshold mode

In this mode, each event is assigned a “threshold” value, which indicates whether each event occurred at least the specified number of times during each sampling interval.

**PPU Bookmarks**

The CPC tool offers a feature that allows finer grained tracing method for signals sampling. The PPU Bookmark mode is provided as an option to start or stop the counters when a value is written to the *bookmark register*. The triggering write can be issued from both a command line and within your application, achieving the desired sampling scope narrowing.

The chosen bookmark register must be specified as a command line option for the CPC, as well as the desired action to be performed on the counter sampling. The registers can be reached as files in the sysfs file system:

```
/sys/devices/system/cpu/cpu*/pmu_bookmark
```

**Overall usage**

The typical command line syntax for CPC is as follows:

```
cpc [options] [workload]
```

Here, the presence of the `[workload]` parameter controls whether the CPC must be run against a single application (workload mode) or against the whole running system (system-wide mode). In system-wide mode, the following options control both the duration and broadness of the sampling:

- `--cpus <CPUS>`
  Controls which processors to use, and where CPUS should be a comma separated list of processor or the keyword *all*

- `--time <TIME>`
  Controls the sampling duration time (in seconds)

**Typical options**

The following options enable the features of CPC:

- `--list-events`
  Returns the list of all possible events, grouped by logic units within the Cell/B.E. system (see Example 5-9).
Example 5-9  Possible events returned

Performance Monitor Signals

Key:
1) Signal Number:
   Digit 1 = Section Number
   Digit 2 = Subsection Number
   Digit 3,4 = Bit Number
   .C (Cycles) or .E (Events) if the signal can record
   either
2) Count Type
   C = Count Cycles
   E = Count Event Edges
   V = Count Event Cycles
   S = Count Single-Cycle Events
3) Signal Name
4) Signal Description

********************************************************************
* Unit 2: PowerPC Processing Unit (PPU)
********************************************************************

2.1 PPU Instruction Unit - Group 1 (NClk)
-----
2100V  Branch_Commit_t0 Branch instruction committed. (Thread 0)
2101E  Branch_Flush_t0 Branch instruction that caused a misprediction
       flush is committed. Branch misprediction includes: (1) misprediction
       of taken or not-taken on conditional branch, (2) misprediction of
       branch target address on bc[1] and bc[1]. (Thread 0)
2102C  Ibuf_Empty_t0 Instruction buffer empty. (Thread 0)
2103E  IERAT_Miss_t0 Instruction effective-address-to-real-address
       translation (I-ERAT) miss. (Thread 0)
2104.CC
2104.EE  IL1_Miss_Cycles_t0 Instruction cache miss cycles. Counts
       the cycles from the miss event until the returned instruction is
       dispatched or cancelled due to branch misprediction, completion
       restart, or exceptions (see Note 1). (Thread 0)
2106C Dispatch Blocked toValid instruction available for dispatch, but dispatch is blocked. (Thread 0)

2109E Instr Flushed toInstruction in pipeline stage EX7 causes a flush. (Thread 0)

2111V PPC Commit toTwo PowerPC instructions committed. For microcode sequences, only the last microcode operation is counted. Committed instructions are counted two at a time. If only one instruction has committed for a given cycle, this event will not be raised until another instruction has been committed in a future cycle. (Thread 0)

2119V Branch Commit toBranch instruction committed. (Thread 1)

2120E Branch Flush toBranch instruction that caused a misprediction flush is committed. Branch misprediction includes: (1) misprediction of taken or not-taken on conditional branch, (2) misprediction of branch target address on bclr[1] and bcctr[1]. (Thread 1)

2121C Ibff Empty toInstruction buffer empty. (Thread 1)

► --event <ID>

Specifies the event to be counted.

► --event <ID.E>

Some events allow counting of either events (E) or cycles (C).

► --event <ID:SUB>

When specifying SPU or MFC events, the desired subunit can be given (see Example 5-10).

Example 5-10 Specifying subunits for events

cpc -event 4103:1 ... # Count event 4103 on SPU 1.

► --event <ID[E][:SUB],...

Specifies multiple comma-separated events (in all of the previous forms) to be counted.
Multiple specification of the event option, allows events to be grouped in sets, with the kernel cycling through them at the interval defined by the switch-timeout option (see Example 5-11).

**Example 5-11  Multiple sets of events**

```
cpc --event 4102:0,4103:0 --event 4102:1,4103:1 \ 
   --switch-timeout 150m ....
```

Specifies the interval time for the Hardware Sampling mode. Uses a suffix on the value of “n” for nanoseconds, “u” for microseconds, or “m” for milliseconds.

**Example 5-12  Hardware Sampling mode**

```
cpc --event 4102:0=1000,4103:0=1000 --interval 10m \ 
   --sampling-mode threshold ....
```

Starts counters upon the PPU hardware-thread 0 bookmark start.

**Example 5-13  PPU Bookmarks with command line trigger**

```
cpc --events 4103 --start-on-ppu-th0-bookmark app # set-up bookmark
```
There are two choices for triggering the counters, as shown in Example 5-14 and Example 5-15.

**Example 5-14  Command line trigger**

```bash
echo 9223372036854775808 > /sys/devices/system/cpu/cpu0/pmu_bookmark
```

**Example 5-15  Program embedded trigger**

```c
#define PMU_BKMK_START (1ULL << 63)
char str[20] ;
fd = open("/sys/devices/system/cpu/cpu0/pmu_bookmark", O_WRONLY);
sprintf(str, "%llu", PMU_BKMK_START);
write(fd, str, strlen(str) + 1);
```

## 5.6.3 OProfile

OProfile for the Cell/B.E. system is a system-level profiler for Cell Linux systems and is capable of profiling all running code at low overhead. It consists of a kernel driver, a daemon for collecting sample data, and several post-profiling tools for turning data into information.

OProfile leverages the hardware performance counters of the processor to enable profiling of a wide variety of interesting statistics, which can also be used for basic time-spent profiling. All code is profiled, including hardware and software interrupt handlers, kernel modules, the kernel, shared libraries, and applications.

The OProfile tool can be used in a number of situations, for example:

- Profiling an application and its shared libraries
- Profiling interrupt handlers
- Profiling performance behavior of entire system
- Examining hardware effects

The OProfile tool requires low overhead and cannot use other highly intrusive profiling methods. In addition, the tool requires instruction-level profiles and call-graph profiles.

**Operation**

The OProfile requires root privileges and exclusive access to the Cell/B.E. PMU. For those reasons, it supports only one session at a time and no other PMU related tool (such as CPC) can be running simultaneously.
**Tools**
The tool is composed of utilities that control the profiling session and reporting. Table 5-4 lists the most relevant utilities.

**Table 5-4  OProfile relevant utilities**

<table>
<thead>
<tr>
<th>Tool</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>opcontrol</td>
<td>Configures and controls the profiling system. Sets the performance counter event to be monitored and the sampling rate.</td>
</tr>
<tr>
<td>oreport</td>
<td>Generates the formatted profiles according to symbols or file names. Supports text and XML output.</td>
</tr>
<tr>
<td>opannotate</td>
<td>Generates annotated source or assembly listings. The listings are annotated with hits per each source code line. Requires compilation with debug symbols (-g).</td>
</tr>
</tbody>
</table>

**Considerations**
The current SDK 3.0 version of OProfile for the Cell/B.E. system supports profiling on POWER processor events and SPU cycle profiling. These events include cycles as well as the various processor, cache, and memory events. It is possible to profile up to four events simultaneously on the Cell/B.E. system. There are restrictions on which PPU events can be measured simultaneously. (The tool now verifies that multiple events specified can be profiled simultaneously. In the previous release, the user had to do this verification.)

When using SPU cycle profiling, events must be within the same group due to restrictions in the underlying hardware support for the performance counters. You can use the following command to view the events and which group contains each event:

```
opalcontrol --list-events
```

**Overall process**
The `opalcontrol` utility drives the profiling process, which can be initiated at compilation time, if source annotation is desired. Both the `oreport` and `opannotator` tools are deployed at the end of the process to collect, format, and co-relate sampled information.
Step 1: Compiling (optional)

Usually, OProfile does not require any changes to the compilation options, even with regard to the optimization flags. However, to achieve a better annotation experience, relocation and debugging symbols are required in the application binary. Table 5-5 shows the required flags.

Table 5-5  Required flags

<table>
<thead>
<tr>
<th>Flag</th>
<th>Tool</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>-g</td>
<td>compiler</td>
<td>Instructs the compiler to produce debugging symbols in the resulting binary.</td>
</tr>
<tr>
<td>-Wl,q</td>
<td>linker</td>
<td>Preserves the relocation and the line number information in the final integrated executable.</td>
</tr>
</tbody>
</table>

Step 2: Initializing

As previously explained, the opcontrol utility drives the process from initialization to the end of sampling. Since the OProfile solution comprehends a kernel module, a daemon, and a collection of tools, in order to properly operate the session, both the kernel module and the daemon must be operational, and no other stale session information must be present. Example 5-16 shows the initialization procedure.

Example 5-16  Initialization procedure

```
opcontrol --deinit  # recommended to clear any previous OProfile data
opcontrol --start-daemon --no-vmlinux  # start the OProfile daemon
opcontrol --init   # perform the initialization procedure
opcontrol --reset  # sanity reset
```

--no-vmlinux: The --no-vmlinux option shown in Example 5-16 controls whether kernel profiling should also be carried on. The example here disables such profiling, which is always true when sampling for the SPUs (when there are not any kernels running on them). In case such an option is needed (for PPU only), replace the --no-vmlinux with --vmlinux=/path/to/vmlinux, where vmlinux is the running kernel's uncompressed image.
Step 3: Running
After properly cleaning up and ensuring that the kernel module and daemon are present, we can proceed with the sampling. First, we must properly set how samples should be organized, indicate the desired event group to monitor, and define the number of events per sampling as shown in Example 5-17.

Example 5-17  Adjusting event group and sampling interval

```
opcontrol --separate=all --event=SPU_CYCLES:10000
```

In Example 5-17, notice the following explanations

- **--separate**
  This option organizes samples based on the given separator. “lib” separates dynamically linked library samples per application. “kernel” separates kernel and kernel module samples per application and implies “library.” “thread” gives separation for each thread and task. “cpu” does the separation for each processor. “all” implies all of these options. “none” turns off separation.

- **--event**
  This option defines the events to be measured as well as the interval (in number of events) to sample the Program Counter. The `--list-events` option gives a complete list of the available events. The commonly used events are CYCLES, which measures PPU, and SPU_CYCLES which measures the SPU.

At this point, the OProfile environment is ready to initiate the sampling. We can proceed with the commands shown in Example 5-18.

Example 5-18  Initiating Profiling

```
opcontrol --start # Fires profiling
app # start application ‘app’ (replace with the desired one)
```

Step 4: Stopping
As soon as the application returns, stop the sampling and dump the results as shown in Example 5-19.

Example 5-19  Stopping and collecting results

```
opcontrol --stop
opcontrol --dump
```
**Step 5: Reports**

After sampling data is collected, OProfile offers the `opreport` utility to format and generate profiles. The `opreport` tool allows the output to be created based on symbols and files names, as well as references the source code, for example, for the number of times that a function performed. Example 5-20 lists the commonly used options.

*Example 5-20  Commonly used options for opreport*

```bash
opreport -X -g -l -d -o output.opm
```

Each of the options is explained as follows:

- `-X`  Output is generated in XML format. Is required when used in conjunction with the VPA tool (see 5.6.6, “Visual Performance Analyzer” on page 400).
- `-g`  Maps each symbol to its corresponding source file and line.
- `-l`  Organizes sampling information per symbol.
- `-d`  For each symbol, shows per-instruction details.
- `-o`  Specifies the output file name.

### 5.6.4 Performance Debugging Tool

The PDT provides a means for tracing to record significant events during program execution and maintaining the sequential order of events. The PDT provides the ability to trace events of interest, in real time, and record relevant data from the SPEs and PPE.

The PDT achieves this objective by instrumenting the code that implements key functions of the events on the SPEs and PPE and collecting the trace records. This instrumentation requires additional communication between the SPEs and PPE as trace records are collected in the PPE memory. The traced records can then be viewed and analyzed by using additional SDK tools.

**Operation**

Tracing is enabled at the application level (user space). After the application is enabled, the tracing facility trace data is gathered every time the application runs.

Prior to each application run, the user can configure the PDT to trace events of interest. The user can also use the PDT API to dynamically control the tracing.

During the application run, the PPE and SPE trace records are gathered in a memory-mapped (mmap) file in the PPE memory. These records are written into the file system when appropriate. The event-records order is maintained in this
The SPEs use efficient DMA transfers to write the trace records into the mmap file. The trace records are written in the trace file using a format that is set by an external definition (using an XML file). The PDTR (see “PDTR” on page 392) and Trace Analyzer (see “Trace Analyzer” on page 409) tools, which use PDT traces as input, use the same format definition for visualization and analysis. Figure 5-26 illustrates the tracing architecture.

Considerations

Tracing 16 SPEs using one central PPE might lead to a heavy load on the PPE and the bus, and therefore, might influence the application performance. The PDT is designed to reduce the tracing execution load and provide a means for throttling the tracing activity on the PPE and each SPE. In addition, the SPE tracing code size is minimized, so that it fits into the small SPE local storage.

Events tracing is enabled by instrumenting selected functions of the following SDK libraries:

- On the PPE: DaCS, ALF, libspe2, and libsync
- On the SPE: DaCS, ALF, libsync, the spu_mfcio header file, and the overlay manager

Performance events are captured by the SDK functions that are already instrumented for tracing. These functions include SPEs activation, DMA transfers, synchronization, signaling, user-defined events, and so on. Statically
linked applications should be compiled and linked with the trace-enabled libraries. Applications that use shared libraries are not required to be rebuilt.

**Overall process**
The PDT tracing facility is designed to minimize the effort that is needed to enable the tracing facility for a given application. The process includes compiling, linking with trace libraries, setting environment variables, (optionally) adjusting the trace configuration file, and running the application. (In most cases on the SPU, code must be compiled since it is statically linked.)

**Step 1: Compilation**
The compilation part involves the specification of a few tracing flags and the tracing libraries. There two procedures, one for the PPE and one for SPE.

For the SPE, follow this procedure:

1. Add the following compilation flags. CFLAGS is the variable in the SDK makefile.
   ```
   -Dmain=_pdt_main -Dexit=_pdt_exit -DMFCIO_TRACE -DLIBSYNC_TRACE
   ```
2. Add the trace headers to the beginning of the include path. INCLUDE is the variable in the SDK makefile.
   ```
   -I/usr/spu/include/trace
   ```
3. Add the instrumented libraries in /usr/spu/lib/trace (for example, libtrace.a) to the linker process. LDFLAGS is the variable in the SDK makefile for the library path, and IMPORTS is the variable in the SDK makefile for the library.
   ```
   -L/usr/spu/lib/trace -ltrace
   ```
4. (Optional) To enable the correlation between events and the source code, for the analysis tools, rebuild the application by using the linking relocation flags. LDFLAGS is the variable in the SDK makefile.
   ```
   -Wl,q
   ```

For the PPE, follow this procedure:

1. (Optional, if using libsync) Add the following compilation flag. The CFLAGS variable is in the SDK makefile.
   ```
   -DLIBSYNC_TRACE
   ```
2. Add the trace headers to the beginning of the include path. The INCLUDE variable is in the SDK makefile.
   ```
   -I/usr/include/trace
   ```
3. Add the instrumented libraries (for example, libtrace.a) in /usr/lib/trace (or /usr/lib64/trace for 64-bit applications) to the linker process. The LDFLAGS variable is in the SDK makefile for the library path, and the IMPORTS variable is in the SDK makefile for the library. Enter either of the following commands depending on whether you have 64-bit applications:

- L/usr/lib/trace -ltrace
- L/usr/lib64/trace -ltrace

4. (Optional) To enable the correlation between events and the source code, for the analysis tools, rebuild the application by using the linking relocation flags. The LDFLAGS variable is in the SDK makefile:

- Wl,q

**Step 2: Preparing the run environment**

After the build process is complete, the PDT requires the environment variables, listed in Table 5-6, to be set prior to running the application.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>LD_LIBRARY_PATH</td>
<td>The full path to the traced library location is /usr/lib/trace (or -L/usr/lib64/trace for 64-bit applications).</td>
</tr>
<tr>
<td>PDT_KERNEL_MODULE</td>
<td>This is the PDT kernel module installation path. It should be /usr/lib/modules/pdt.ko.</td>
</tr>
<tr>
<td>PDT_CONFIG_FILE</td>
<td>This is the full path to the PDT configuration file for the application run. The PDT package contains a pdt_cbe_configuration.xml file in the /usr/share/pdt/config directory that can be used “as is” or copied and modified for each application run.</td>
</tr>
<tr>
<td>PDT_TRACE_OUTPUT</td>
<td>(Optional) This is the full path to the PDT output directory, which must exist prior to the application running.</td>
</tr>
<tr>
<td>PDT_OUTPUT_PREFIX</td>
<td>(Optional) This variable is used to add a prefix to the PDT output files names.</td>
</tr>
</tbody>
</table>

**Step 2a: (Optional, but recommended) Preparing configuration files**

A configuration XML file is used to configure the PDT. The PDT tracing facility that is built into the application at run time reads the configuration file that is defined by the PDT_CONFIG_FILE environment variable. The /usr/share/pdt/config directory contains a reference configuration file (pdt_cbe_configuration.xml). This file should be copied and then specifically modified for the requirements of each application.
The /usr/share/pdt/config directory also contains reference configuration files for applications that are using the DaCS and ALF libraries: pdt_dacs_config_cell.xml for DaCS and pdt_alf_config_cell.xml for ALF. In addition, a pdt_libsync_config.xml reference file is provided for applications that use the libsync library.

The first line of the configuration file contains the application name. This name is used as a prefix for the PDT output files. To correlate the output name with a specific run, the name can be changed before each run. The PDT output directory is also defined in the output_dir attribute. This location is used if the PDT_TRACE_OUTPUT environment variable is not defined.

The first section of the file, <groups>, defines the groups of events for the run. The events of each group are defined in other definition files, which are also in XML format, and are included in the configuration file. These files reside in the /usr/share/pdt/config directory. They are provided with the instrumented library and should not be modified by the programmer.

Each file contains a list of events with the definition of the trace-record data for each event. Some of the events define an interval (with StartTime and EndTime), and some are single events in which the StartTime is 0 and the EndTime is set to the event time.

The names of the trace-record fields are the same as the names defined by the API functions. There are two types of records: one for the PPE and one for the SPE. Each of these record types has a different header that is defined in a separate file: pdt_ppe_event_header.xml for the PPE and pdt_spe_event_header.xml for the SPE.

The SDK provides instrumentation for the following libraries (events are defined in the XML files):

- GENERAL (pdt_general.xml)
  These are the general trace events such as trace start and trace stop. Tracing of these events is always active.

- LIBSPE2 (pdt_libspe2.xml)
  These are the libspe2 events.

- SPU_MFCIO (pdt_mfcio.xml)
  These are the spu_mfcio events that are defined in the spu_mfcio.h header file.

- LIBSYNC (pdt_libsync.xml)
  These are the mutex events that are part of the libsync library.
DACS (pdt_dacs.xml, pdt_dacs_perf.xml, and pdt_dacs_spu.xml)
These are the DaCS events, separated into three groups of events.

ALF (pdt_alf.xml, pdt_alf_perf.xml, and pdt_alf_spu.xml)
These are the ALF events, separated into three groups of events.

The second section of the file contains the tracing control definitions for each type of processor. The PDT is made ready for the hybrid environment so that each processor has a host, <host>. On each processor, several groups of events can be activated in the group control, <groupControl>. Each group is divided into subgroups, and each subgroup, <subgroup>, has a set of events. Each group, subgroup, and event has an active attribute that can be either true or false. This attribute affects tracing as follows:

- If a group is active, all of its events are traced.
- If a group is not active, and the subgroup is active, all of its subgroup’s events are traced.
- If a group and subgroup are not active, and an event is active, that event is traced.

We recommended that you enable tracing only for those events that are of interest. We also recommend that you to turn off tracing of non-stalling events, since the relative overhead is higher. Depending on the number of processors that are involved, programs might produce events at a high rate. If this scenario occurs, the number of traced events might also be high.

**Step 3: Running the application**

After both the environment variables and configuration files are set, simply execute the application. The PDT produces trace files in a directory that is defined by the environment variable PDT_TRACE_OUTPUT. If this environment variable is not defined, the output location is taken from the definition provided by the output_dir attribute in the PDT configuration file. If neither is defined, the current path is used. The output directory must exist prior to the application run, and the user must have a write access to this directory. The PDT creates the files shown in Table 5-7 on page 392 in that output directory at each run.
Table 5-7  Output files

<table>
<thead>
<tr>
<th>File</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>&lt;prefix&gt;-&lt;app_name&gt;-yyyyymmddhhmmss.pex</code></td>
<td>Meta file of the trace.</td>
</tr>
<tr>
<td><code>&lt;prefix&gt;-&lt;app_name&gt;-yyyyymmddhhmmss.maps</code></td>
<td>Maps file from /proc/<code>&lt;pid&gt;/</code> for the address-to-name resolution performed by the PDTR tool (or <code>pdtr</code> command).</td>
</tr>
<tr>
<td><code>&lt;prefix&gt;-&lt;app_name&gt;-yyyyymmddhhmmss.&lt;N&gt;.trace</code></td>
<td>Trace file. An application might produce multiple trace files where <code>&lt;N&gt;</code> is the index.</td>
</tr>
</tbody>
</table>

**Output file variables:**

- The `<prefix>` variable is provided by the optional PDT_OUTPUT_PREFIX environment variable.
- The `<app_name>` variable is a string provided in the PDT configuration file `application_name` attribute.
- The `yyyyymmddhhmmss` variable is the date and time when the application started (trace_init() time).
- The `<N>` variable is the serial number of the trace file. The maximum size of each trace file is 32 MB.

**PDTR**

The PDTR command-line tool (`pdtr` command) provides both viewing and post processing of PDT traces on the target (client) machine. The alternative is to use the graphical Trace Analyzer, part of VPA (see 5.6.6, “Visual Performance Analyzer” on page 400).

To use this tool, you must instrument your application by building with the PDT. After the instrumented application has run and created the trace output files, you can run the `pdtr` command to show the trace output.

For example, consider the following PDT trace file set:

```
app-20071115094957.1.trace
app-20071115094957.maps
app-20071115094957.pex
```

In this case, run the `pdtr` command as follows:

```
pdtr [options] app-20071115094957
```
The `pdtr` command produces the output file `app-2007115094957.pep`.

The PDTR tool produces various summary output reports with lock statistics, DMA statistics, mailbox usage statistics, and overall event profiles. The tool can also produce sequential reports with time-stamped events and parameters per line.

Example 5-21, Example 5-22 on page 394, and Example 5-23 on page 394 show the reports that are produced by PDTR. See the PDTR man page for additional output examples and usage details.

**Example 5-21  General Summary Report**

General Summary Report

```
1.107017 seconds in trace

Total trace events: 3975

<table>
<thead>
<tr>
<th>Count</th>
<th>EvID</th>
<th>Event</th>
<th>min</th>
<th>avg</th>
<th>max</th>
<th>evmin,evmax</th>
</tr>
</thead>
<tbody>
<tr>
<td>672</td>
<td>1202</td>
<td>SPE_MFC_READ_TAG_STATUS</td>
<td>139.7ns</td>
<td>271.2ns</td>
<td>977.8ns</td>
<td>26, 3241</td>
</tr>
<tr>
<td>613</td>
<td>0206</td>
<td>_DACS_HOST_MUTEX_LOCK</td>
<td>349.2ns</td>
<td>1.6us</td>
<td>20.3us</td>
<td>432, 2068</td>
</tr>
<tr>
<td>613</td>
<td>0406</td>
<td>_DACS_HOST_MUTEX_UNLOCK</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>336</td>
<td>0402</td>
<td>SPE_MFC_GETF</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>240</td>
<td>1406</td>
<td>_DACS_SPE_MUTEX_UNLOCK</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>239</td>
<td>1206</td>
<td>_DACS_SPE_MUTEX_LOCK</td>
<td>279.4ns</td>
<td>6.6us</td>
<td>178.7us</td>
<td>773, 3152</td>
</tr>
<tr>
<td>224</td>
<td>0102</td>
<td>SPE_MFC_PUTF</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>99</td>
<td>0200</td>
<td>HEART_BEAT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>96</td>
<td>0302</td>
<td>SPE_MFC_GET</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>64</td>
<td>1702</td>
<td>SPE_READ_IN_MBOX</td>
<td>139.7ns</td>
<td>3.7us</td>
<td>25.0us</td>
<td>21, 191</td>
</tr>
<tr>
<td>16</td>
<td>0002</td>
<td>SPE_MFC_PUT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>2007</td>
<td>_DACS_MBOX_READ_ENTRY</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>2107</td>
<td>_DACS_MBOX_READ_EXIT_INTERVAL</td>
<td>11.7us</td>
<td>15.2us</td>
<td>25.9us</td>
<td>557, 192</td>
</tr>
<tr>
<td>16</td>
<td>0107</td>
<td>_DACS_RUNTIME_INIT_ENTRY</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>2204</td>
<td>_DACS_MBOX_WRITE_ENTRY</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>0207</td>
<td>_DACS_RUNTIME_INIT_EXIT_INTERVAL</td>
<td>6.6us</td>
<td>7.4us</td>
<td>8.2us</td>
<td>29, 2030</td>
</tr>
<tr>
<td>16</td>
<td>2304</td>
<td>_DACS_MBOX_WRITE_EXIT_INTERVAL</td>
<td>4.1us</td>
<td>6.0us</td>
<td>11.3us</td>
<td>2011, 193</td>
</tr>
<tr>
<td>16</td>
<td>0601</td>
<td>SPE_PROGRAM_LOAD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>0700</td>
<td>SPE_TRACE_START</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>0800</td>
<td>SPE_TRACE_END</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>2A04</td>
<td>_DACS_MUTEX_SHARE_ENTRY</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

...
Example 5-22  Sequential trace output

--- Trace File(s) -----------------------------------------------
Event type size: 0
Metafile version: 1

--- Trace File(s) -----------------------------------------------
0  0.000000  0.000ms  PPE_HEART_BEAT  PPU 00000001  TB:0000000000000000
1  0.005025  5.025ms  PPE_SPE_CREATE_GROUP  PPU F7FC73A0  TB:000000000001190F  *** Unprocessed event ***
2  0.015968  10.943ms  PPE_HEART_BEAT  PPU 00000001  TB:0000000000037D13
3  0.031958  15.990ms  PPE_HEART_BEAT  PPU 00000001  TB:000000000006FB69
4  0.047957  15.999ms  PPE_HEART_BEAT  PPU 00000001  TB:00000000000A7A3B
5  0.053738  5.781ms  PPE_SPE_CREATE_THREAD  PPU F7FC73A0  TB:00000000000BBD90
6  0.053768  0.030ms  PPE_SPE_WRITE_IN_MBOX  PPU F7FC73A0  TB:00000000000BBF3F  *** Unprocessed event ***
7  0.000000  0.000us  SPE_ENTRY  SPU 1001F348  Decr:00000001
8  163  11.384us  SPE_MFC_WRITE_TAG_MASK  SPU 1001F348  Decr:000000A4  *** Unprocessed event ***
9  170  0.489us  SPE_MFC_GET  SPU 1001F348  Decr:000000AB  Size: 0x80 (128), Tag: 0x4 (4)
10 176  0.419us  SPE_MFC_WRITE_TAG_UPDATE  SPU 1001F348  Decr:000000B1  *** Unprocessed event ***
11 183  0.489us  SPE_MFC_READ_TAG_STATUS_ENTRY  SPU 1001F348  Decr:000000B8
12 184  0.070us  SPE_MFC_READ_TAG_STATUS_EXIT  SPU 1001F348  Decr:000000B9  >>> delta tics:1 ( 0.070us)
rec:24 {DMA done[tag=4,0x4] rec:22  0.978us 130.9MB/s}
20  0  0.000us  SPE_ENTRY  SPU 1001F348  Decr:00000001
21 163  11.384us  SPE_MFC_WRITE_TAG_MASK  SPU 1001F348  Decr:000000A4  *** Unprocessed event ***
22 170  0.489us  SPE_MFC_GET  SPU 1001F348  Decr:000000AB  Size: 0x80 (128), Tag: 0x4 (4)
23 176  0.419us  SPE_MFC_WRITE_TAG_UPDATE  SPU 1001F348  Decr:000000B1  *** Unprocessed event ***
24 183  0.489us  SPE_MFC_READ_TAG_STATUS_ENTRY  SPU 1001F348  Decr:000000B8
25 184  0.070us  SPE_MFC_READ_TAG_STATUS_EXIT  SPU 1001F348  Decr:000000B9  >>> delta tics:1 ( 0.070us)
rec:32 {DMA done[tag=4,0x4] rec:30  0.978us 130.9MB/s}
33 4523  0.210us  SPE_MUTEX_LOCK_EXIT  SPU 1001F348  Lock:1001E280  Decr:000011AC  >>> delta tics:3 ( 0.210us)
34  0  0.000us  SPE_ENTRY  SPU 1001F9D8  Decr:00000001
35  96  6.705us  SPE_MFC_WRITE_TAG_MASK  SPU 1001F9D8  Decr:00000061  *** Unprocessed event ***
36 103  0.489us  SPE_MFC_GET  SPU 1001F9D8  Decr:00000068  Size: 0x80 (128), Tag: 0x4 (4)
37 109  0.419us  SPE_MFC_WRITE_TAG_UPDATE  SPU 1001F9D8  Decr:0000006E  *** Unprocessed event ***
38 116  0.489us  SPE_MFC_READ_TAG_STATUS_ENTRY  SPU 1001F9D8  Decr:00000075
39 117  0.070us  SPE_MFC_READ_TAG_STATUS_EXIT  SPU 1001F9D8  Decr:00000076  >>> delta tics:1 ( 0.070us)
rec:38 {DMA done[tag=4,0x4] rec:36  0.978us 130.9MB/s}

Example 5-23  Lock report

==================================================================================================================================
<table>
<thead>
<tr>
<th>Accesses</th>
<th>Hits</th>
<th>Misses</th>
<th>Hit hold time (us)</th>
<th>Miss wait time (us)</th>
<th>Account</th>
</tr>
</thead>
<tbody>
<tr>
<td>%Total</td>
<td>Count</td>
<td>%Account</td>
<td>Count</td>
<td>%Account</td>
<td>min, avg, max</td>
</tr>
<tr>
<td>---------</td>
<td>-----</td>
<td>-------</td>
<td>-------------------</td>
<td>-------------------</td>
<td>---------</td>
</tr>
</tbody>
</table>
* 600 (100.0) 3 (0.5) 597 (99.5) 100.8, 184.6, 402.4 13.3, 264.4, 568.0  shr_lock (0x10012180)
2 (66.7) 298 (49.9) 100.8, 101.1, 101.5 181.8, 249.7, 383.6  main (0x68c)(lspe=1)
1 (33.3) 199 (33.3) 200.7, 201.3, 202.5 13.3, 315.2, 568.0  main (0x68c)(lspe=2)
0 (0.0) 100 (16.8) 0.0, 0.0, 0.0 205.0, 206.8, 278.5  main (0x68c)(lspe=3)

Implicitly initialized locks (used before/without mutex_init)

See the PDTR man page for additional output examples and usage details.

5.6.5  FDPR-Pro

The Post-link Optimization for Linux on Power tool (FDPR-Pro or fdprpro) is a performance tuning utility that reduces execution time and real memory utilization of user space application programs. It optimizes the executable image of a program by collecting information about the behavior of the program under a workload. It then creates a new version of that program optimized for that
workload. The new program typically runs faster and uses less real memory than the original program.

**Operation**
The post-link optimizer builds an optimized executable program in three distinct phases:

1. Instrumentation phase, where the optimizer creates an instrumented executable program and an empty template profile file
2. Training phase, where the instrumented program is executed with a representative workload, and as it runs, it updates the profile file
3. Optimization phase, where the optimizer generates the optimized executable program file

You can control the behavior of the optimizer with options specified on the command line.

**Considerations**
The FDPR-Pro tool applies advanced optimization techniques to a program. Some aggressive optimizations might produce programs that do not behave as expected. You should test the resulting optimized program with the same test suite that is used to test the original program. You cannot re-optimize an optimized program by passing it as input to FDPR-Pro.

An instrumented executable, created in the instrumentation phase and run in the training phase, typically runs several times slower than the original program. This slowdown is caused by the increased execution time required by the instrumentation. Select a lighter workload to reduce training time to a reasonable value, while still fully exercising the desired code areas.
Overall process
The typical FDPR-Pro process encompasses instrumentation, training, and optimization, as illustrated in Figure 5-27.

**Figure 5-27** FDPR-Pro process

**Step 1: Preparing the input files**
The input to the `fdprpro` command must be an executable or a shared library (for PPE files) produced by the Linux linker. `fdprpro` supports 32-bit or 64-bit programs compiled by the GCC or XLC compilers.

Build the executable program with relocation information. To do this, call the linker with the `--emit-relocs` (or `-q`) option. Alternatively, pass the `-Wl,--emit-relocs` (or `-Wl,-q`) options to the GCC or XLC compiler.
If you are using the SDK makefiles structure (with make.footer), set the variables shown in Example 5-24 (depending on the compiler) in the makefile.

Example 5-24  make.footer variables

<table>
<thead>
<tr>
<th>Variable</th>
<th>Compiler</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDFLAGS_xlc</td>
<td>XCL</td>
<td>For XLC</td>
</tr>
<tr>
<td>LDFLAGS_gcc</td>
<td>GCC</td>
<td>For GCC</td>
</tr>
</tbody>
</table>

Step 2: Instrumentation phase

PPE and SPE executables are instrumented differently. For the PPE executable, the `fdprpro` command creates an instrumented file and a profile file. The profile file is later populated with profile information, while the instrumented program runs with a specified workload. In contrast, for the SPE executable, the profile (with extension .mprof) is created only when the instrumented program runs.

FDPR-Pro processes (instruments or optimizes) the PPE program and the embedded SPE images. Processing the SPE images is done by extracting them to external files, processing each of them, and then encapsulating them back into the PPE executable. Two modes are available in order to fully process the PPE/SPE combined file: integrated mode and standalone mode.

- Integrated mode

  The integrated mode of operation does not expose the details of SPE processing. This interface is convenient for performing full PPE/SPE processing, but flexibility is reduced. To completely process a PPE/SPE file, run the `fdprpro` command with the `-cell` (or `--cell-supervisor`) command-line option, as shown in the following example:

  ```
  fdprpro -cell -a instr myapp -o myapp.instr
  ```

- Standalone mode

  As opposed to integrated mode, where the same optimization options are used when processing the PPE file and when processing each of the SPE files, full flexibility is available in standalone mode. With full flexibility, you can specify the explicit commands that are needed to extract the SPE files, process them, and then encapsulate and process the PPE file. The following list shows the details of this mode.

  a. Extract the SPE files.

     SPE images are extracted from the input program and written as executable files in the specified directory:

     ```
     fdprpro -a extract -spedir somedir myapp
     ```
b. Process the SPE files.

The SPE images are processed one by one. You should place all of the output files in a distinct directory.

\[ \text{fdprpro -a (instr|opt) somewdir/spe_i [-f prof_i] [opts ...] outdir/spe_i} \]

\textbf{spe}_i: Replace the variable \(<spe_i>\) with the name of the SPU file obtained in the extraction process. The profile and optimization options must be specified only in the optimization phase (see “Step 4: Optimization phase” on page 399).

c. Encapsulate and process the PPE file.

The SPE files are encapsulated as a part of the PPE processing. The \(-\text{spedir}\) option specifies the output SPE directory.

\[ \text{fdprpro -a (instr|opt) --encapsulate -spedir outdir [ opts ...]} \]
\[ \text{myapp -o myapp.instr} \]

\textbf{SPE instrumentation}

When the optimizer processes PPE executables, it generates a profile file and an instrumented file. The profile file is filled with counts while the instrumented file runs. In contrast, when the optimizer processes SPE executables, the profile is generated when the instrumented executable runs. Running a PPE or SPE instrumented executable typically generates a number of profiles, one for each SPE image whose thread is executed. This type of profile accumulates the counts of all threads that execute the corresponding image. The SPE instrumented executable generates an SPE profile named \(<\text{spename}>.\text{mprof}\) in the output directory, where \(<\text{spename}\>) represents the name of the SPE thread.

The resulting instrumented file is 5% to 20% larger than the original file. Because of the limited local storage size of the CBEA, instrumentation might cause SPE memory overflow. If this happens, \texttt{fdprpro} issues an error message and exits. To avoid this problem, the user can use the \--ignore-function-list file or \texttt{-ifl} file option. The file that is referenced by the file parameter contains the names of the functions that should not be instrumented and optimized. This results in a reduced instrumented file size. Specify the same \texttt{-ifl} option in both the instrumentation and optimization phases.

\textbf{Step 3: Training phase}

The training phase consists of running the instrumented application with a representative workload, one that fully exercises the desired code areas. While the program runs with the workload, the PPE profile, by default with the .nprof extension, is populated with profile information. Simultaneously, SPE profiles, one for each executed SPE thread, are generated in the current directory.
If an old profile exists before instrumentation starts, `fdprpro` accumulates new data into it. In this way, you can combine the profiles of multiple workloads. If you do not want to combine profiles, remove the old SPE profiles (the .mprof files) and replace the PPE profile (by default .nprof file) with its original copy, before starting the instrumented program.

The instrumented PPE program requires a shared library named libfsprinst32.so for ELF32 programs, or libfdprinst64.so for ELF64 programs. These libraries are placed in the library search path directory during installation.

The default directory for the profile file is the directory that contains the instrumented program. To specify a different directory, set the environment variable FDPR_PROF_DIR to the directory that contains the profile file.

**Step 4: Optimization phase**

The optimization phase takes all profiling information generated during the test phase in order to optimize the application. Example 5-25 shows the typical optimization that occurs.

*Example 5-25  Typical optimization*

```bash
fdprpro -a opt -f myapp.nprof [ opts ... ] myapp -o myapp.fdpr
```

The same considerations shown in the instrumentation phase, with regard to *integrated mode* and *standalone mode*, are still valid during the optimization phase. The notable exceptions are the action command being performed (here `-a opt`) and the options (here related to the optimization phase). The routine involves extracting, processing, and encapsulating.

**Optimization options**

If you invoke `fdprpro` with the basic optimization flag `-O`, it performs code reordering optimization. It also optimizes branch prediction, branch folding, code alignment, and removal of redundant NOOP instructions.

To specify higher levels of optimization, pass one of the flags `-O2`, `-O3`, or `-O4` to the optimizer. Higher optimization levels perform more aggressive function inlining, data flow analysis optimizations, data reordering, and code restructuring, such as loop unrolling. These high-level optimization flags work well for most applications. You can achieve optimal performance by selecting and testing specific optimizations for your program.
5.6.6 Visual Performance Analyzer

The VPA is an Eclipse-based tool set and currently includes six plug-in applications that work cooperatively:

- Profile Analyzer
- Code Analyzer
- Pipeline Analyzer
- Counter Analyzer
- Trace Analyzer
- Control Flow Analyzer (experimental)

VPA aims to provide a platform independent, easy-to-use integrated set of graphical application performance analysis tools, leveraging existing platform specific non-GUI performance analysis tools to collect a comprehensive set of data. In doing so, VPA creates a consistent set of integrated tools to provide a platform independent drill-down performance analysis experience. Figure 5-28 illustrates the architecture of this tool set.

Figure 5-28 VPA architecture
However VPA does not supply performance data collection tools. Instead, it relies on platform-specific tools, such as OProfile and PDT, to collect the performance data as illustrated in Figure 5-29.

In general, each visualization tool acts complementary to the other:

**Profile Analyzer**
Is a system profile analysis tool. This plug-in obtains profile information from various platform specific tools, and provides analysis views for the user to identify performance bottlenecks.

**Pipeline Analyzer**
Gets pipeline information of Power processors, and provides two analysis views: scroll mode and resource mode.

**Code Analyzer**
Reads XCOFF (AIX binary file format) files or ELF files running Linux on Power Systems Servers, and displays program structure with block information. With related profile information, it can provide analysis views on the hottest program block as well as optimization suggestions.
Counter Analyzer
Reads counter data files generated by AIX `hpmcount` or `hpmstat` and provides multiple views to help users identify and eliminate performance bottlenecks by examining the hardware performance counter values, computed performance metrics, and CPI breakdown models.

Trace Analyzer
Reads in traces generated by the Performance Debugging Tool for the Cell/B.E. system. Displays time-based graphical visualization of the program execution as well as a list of trace contents and the event details for selection.

Control Flow Analyzer
Reads the call trace data file and displays the execution flow graph and call tree to help the user analyze when and where one method of invocation happens and how long it runs.

The VPA is better described by its own individual tools. In the case of the Cell/B.E. processor, we have the following available relevant tools:

- OProfile
- CPC
- PDT
- FDPR-Pro

Considering these Cell/B.E. performance tools, we focus on each respective visualization tool, namely the Profile Analyzer, Counter Analyzer, Trace Analyzer and Code Analyzer.

**Tool usage:** The following sections provide a brief overview of each Cell/B.E. relevant VPA tools. For more detailed coverage of tool usage, see Chapter 6, “The performance tools” on page 417.

Profile Analyzer
With the Profile Analyzer tool, you can navigate through a system profile, looking for performance bottlenecks. This tool provides a powerful set of graphical and text-based views for users to narrow down performance problems to a particular process, thread, module, symbol, offset, instruction, or source line. It supports profiles that are generated by the Cell/B.E. OProfile.

**Overall process**
Profile Analyzer works with properly formatted data, gathered by OProfile. First you run The the desired application with OProfile and format its data. Then you load the information in the Profile Analyzer and explore its visualization options.
**Step 1: Collecting the profile data**

As outlined in 5.6.3, “OProfile” on page 382, initiate the profile session as usual, adding the selected events to be measured. As a required step for VPA, configure the session to properly separate the samples with the following command:

```
opcontrol --separate=all
```

As soon as the measuring is ended, prepare the output data for VPA with the `opreport` tool as follows:

```
opreport -X -g -d -l myapp.opm
```

**Explanation of options:** Consult 5.6.3, “OProfile” on page 382, for an explanation of the options.

**Step 2: Loading the data**

Start the VPA tool. From the menu bar, select **Tools → Profile Analyzer** (Figure 5-30).

![Visual Performance Analyzer](image)

*Figure 5-30   Selecting Profile Analyzer*
Since we already have profile data from the previous step, load the information by selecting File → Open File. In the Open File window (Figure 5-31), select the .opm file that was generated previously.

![Open File window](image)

*Figure 5-31  Open File window*

In VPA, a profile data file loading process can run as a background job. When VPA is loading a file, you can click a button to have the loading job to run in the background. While the loading job is running in the background, you can use Profile Analyzer to view already loaded profile data files or start another loading job at the same time.
The process hierarchy view (Figure 5-32) appears by default in the top center pane. It shows an expandable list of all processes within the current profile. You can expand a process to view its module, later thread, and so on. You can also view the profile in the form of a thread or module, for example. You can define the hierarchy view by right-clicking the profile and choosing Hierarchy Management.

Code Analyzer
Code Analyzer displays detailed information about basic blocks, functions, and assembly instructions of executable files and shared libraries. It is built on top of FDPR-Pro technology and allows the addition of FDPR-Pro and tprof profile information. Code Analyzer can show statistics to navigate the code, to display performance comment and grouping information about the executable, and to map back to source code.

Overall process
The Code Analyzer works with the artifacts that are generated from the FDPR-Pro session on your executable. Initially the original executable is loaded, followed by the .nprof files generated. After that, you should be able to visualize the information.
Step 1: Collecting profile data
Initially, we should run at least one profiling session (without optimization) with your application. As previously explained in 5.6.5, “FDPR-Pro” on page 394, the first step is to instrument the desired application followed by the actual training, that is profile information generation. The expected results are profile files for both PPU code (*.nprof) and SPU code (*.mprof). Example 5-26 shows as typical session.

Example 5-26   Typical FDPR-Pro session

rm *.mprof # remove old data
mkdir ./somespudir# create temp folder
fdprpro -a instr -cell -spedir somespudir myapp # instrument
myapp ... # run your app with a meaningful workload

Explanation of options: Refer to 5.6.5, “FDPR-Pro” on page 394, for an explanation on the options.

Step 2: Loading the data
Start the VPA tool. From the menu bar, select Tools → Code Analyzer (Figure 5-33).
Locate the original application binary add it to the Code Analyzer choosing **File → Code Analyzer → Analyze Executable.** In the Choose Executable File to Analyze window (Figure 5-34), select the desired file and click **Open.**

*Figure 5-34  Selecting the Open File*
The executable is loaded. Information tabs should appear for both PPU code and SPU code as shown in Figure 5-35.

Figure 5-35  Executable information

To enhance the views with profile information for the loaded executable, you can use either an instrumentation profile file or a sampling profile. For that, choose File → Code Analyzer → Add Profile Information for each of the executable tabs that are available in the center of the window. The profile information that is added must match the executable tab that is selected. For the ppu tabs, add the *.nprof profiles, and for the spu tab, add the respective *.nprof file. Figure 5-36 on page 409 shows the added information.
Figure 5-36  Added profile information

**Trace Analyzer**

Trace Analyzer visualizes Cell/B.E. traces that contain such information as DMA communication, locking and unlocking activities, and mailbox messages. Trace Analyzer shows this data organized by core along a common time line. Extra details are available for each kind of event, for example, lock identifier for lock operations and accessed address for DMA transfers.

The tool introduces the following concepts with which you must be familiar:

**Events**

Events are records that have no duration, for example, records describing non-stalling operations, such as releasing a lock. Events input on performance is normally insignificant, but it be important for understanding the application and tracking down sources of performance problems.

**Intervals**

Intervals are records that may have non-zero duration. They normally come from stalling operations, such as acquiring a lock. Intervals are often a significant performance factor. Identifying long stalls and their sources is an important task in performance debugging. A special case of an interval is a *live interval*, which starts when an SPE thread begins to execute and ends when the thread exits.
Overall process
The Trace Analyzer tool works with trace information that is generated by the PDT. The tool then processes the trace for analysis and visualization. Most importantly, this processing adds context parameters, such as estimated wall clock time and unique SPE thread IDs, to individual records.

Step 1: Collecting trace data
As shown in 5.6.4, “Performance Debugging Tool” on page 386, the PDT tool requires the following steps to produce the trace files needed by the Trace Analyzer:

1. Re-compile and link with instrumented libraries.
2. Set up the runtime environment variables.
3. Execute the application.

If everything is properly configured, three tracing related files are generated in the configured output directory: .pex, .maps, and .trace.

Step 2: Loading the trace files
Start the VPA tool. Select Tools → Trace Analyzer (Figure 5-37).

Figure 5-37   Selecting Trace Analyzer
Select **File → Open File** and locate the .pex file generated during the tracing session. After loading in the trace data, the Trace Analyzer Perspective displays the data in its views and editors. See Figure 5-38.

From the view shown in Figure 5-38, going from the top left clockwise, we see the following components:

- The navigator view
- Trace Editor

This editor shows the trace visualization by core, where data from each core is displayed in a separate row, and each trace record is represented by a rectangle. Time is represented on the horizontal axis, so that the location and size of a rectangle on the horizontal axis represent the corresponding event's time and duration. The color of the rectangle represents the type of event, as defined by the Color Map View.
Figure 5-39 shows an example of the details of the Trace Editor view.

- **Details View**
  
  This view show the details of the selected record (if any).

- **Color Map View**
  
  With this view, the user can view and modify color mapping for different kinds of events.

- **Trace Table View**
  
  This view shows all the events on the trace in the order of their occurrence.

**Counter Analyzer**

The Counter Analyzer tool is a common tool to analyze hardware performance counter data among many IBM System i™ platforms, including systems that run on Linux on a Cell/B.E. processor.

The Counter Analyzer tool accepts hardware performance counter data in the form of a cross-platform XML file format. The tool provides multiple views to help the user identify the data. The views can be divided into two categories.
The first category is the “table” views, which are two-dimension tables that display data. The data can be raw performance counter values, derived metrics, counter comparison results, and so on.

The second category is the “plot” views. In these views, data is represented by a different kind of plots. The data can also be raw performance counter values, derived metrics, comparison results, and so on. In addition to these “table” views and “plot” views, “utility” views help the user configure and customize the tool.

**Overall process**

In the case of the Cell/B.E. processor, the Counter Analyzer works with count data that is produced by the CPC tool, in .pmf XML format files. After loading the counter data of the .pmf file, the Counter Analyzer Perspective displays the data in its views and editors.

**Step 1: Collecting count data**

To obtain the required counter data, proceed as explained in 5.6.2, “The CPC tool” on page 376. Make sure that the output is generated in the XML format.

Typically, you use CPC as shown in Example 5-27.

*Example 5-27   Collecting count data*

cpc -e EVENT,... --sampling-mode -c CPUS -i 100000000 –X file.pmf \
-t TIME # for system wide mode

cpc -e EVENT,EVENT,... -i INTERVAL –X file.pmf some_workload # for  
workload mode

**Step 2: Loading the count data**

Start the VPA too. From the menu bar, select **Tools → Counter Analyzer** (Figure 5-40).

![Figure 5-40   Selecting Counter Analyzer](image)
Load the count data by selecting **File → Open File**. In the Open File window, locate the .pmf file that was generated by CPC.

After loading the counter data of the .pmf file, the Counter Analyzer Perspective (Figure 5-41) displays the data in its views and editors. Primary information, including details, metrics, and CPI breakdown, is displayed in Counter Editor. Resource statistics information about the file (if available) is shown in the tabular view Resource Statistics. The View Graph shows the details, metrics, and CPI breakdown in a graphic way.

![Figure 5-41 Counter Analyzer Perspective](image)

The Counter Analyzer organizes the information according to the following concepts:

- **Performance Monitoring**
  The Counter Performance monitor counter provides comprehensive reports of events that are critical to performance on systems. It gathers critical hardware events, such as the number of misses on all cache levels, the number of floating point instructions executed, and the number of instruction loads that cause TLB misses.
 Metrics

The metric information is calculated with a user-defined formula and event count from a performance monitor counter. It is used to provide such performance information as the processor utilization rate, at a million instructions per second. This helps the algorithm designer or programmer identify and eliminate performance bottlenecks.

 CPI Breakdown Model

Cycles per instruction (CPI) is the measurement for analyzing the performance of a workload. CPI is defined as the number of processor clocked cycles that are needed to complete an instruction. It is calculated as shown in the following equation:

CPI = Total Cycles / Number of Instructions Completed

A high CPI value usually implies under utilization of machine resources.

For more information, consult the IBM Visual Performance Analyzer User Guide Version 6.1 manual at the following Web address:


Alternatively, visit the Visual Performance Analyzer page in alphaWorks at the following address:

http://www.alphaworks.ibm.com/tech/vpa
The performance tools

In this chapter, we explore a practical “hands-on” example of using the performance tools. We explain how to collect proper information and how to access relevant visualization features. This chapter includes the following topics:

- 6.1, “Analysis of the FFT16M application” on page 418
- 6.2, “Preparing and building for profiling” on page 418
- 6.3, “Creating and working with the profile data” on page 423
- 6.4, “Creating and working with trace data” on page 437
6.1 Analysis of the FFT16M application

In the section, we present a full example on how to explore the Cell/B.E. performance tools and, especially, how to visualize the results. The chosen target sample application for analysis is the FFT16M application in the IBM Software Developer Kit 3.0 demonstration bundle in /opt/cell/sdk/src/demos/FFT16M.

This manually tuned application performs a 4-way SIMD single-precision complex Fast Fourier Transform (FFT) on an array of size 16,777,216 elements. The following command options are available:

```
fft <ncycles> <printflag> [<log2_spus> <numa_flag> <largepage_flag>]
```

6.2 Preparing and building for profiling

For flexibility, we set up a “sandbox” styled project tree structure, which allows us to have more flexibility while modifying and generating the files.

6.2.1 Step 1: Copying the application from SDK tree

To work in a “sandbox” tree, we create our own copy of the project, on an accessible location, such as your home directory:

```
cp -R /opt/cell/sdk/demos/FFT16M ~/
```

6.2.2 Step 2: Preparing the makefile

Go to your recently created project structure and locate the following makefiles:

- ~/FFT16M/Makefile
- ~/FFT16M/ppu/Makefile
- ~/FFT16M/ppu/Makefile

As explained in the following sections, we make a few modifications to the makefiles to prevent them from installing executables back to the SDK tree. We also introduce the required compilation flags for profiling data.

Notes:

- No further makefile modifications, beyond these, are required.
- There are specific changes depending on whether you use the GCC or XLC compiler.
Modifying the ~/FFT16M/ppu/Makefile

In Example 6-1, we comment out the install directives.

Example 6-1   Changing ~/FFT16M/ppu/Makefile for gcc

```plaintext
# ###########################################################################
# # Target
# ###########################################################################
# PROGRAM_ppu= fft
#
# ###########################################################################
# # Objects
# ###########################################################################
# IMPORTS = ../spu/fft_spu.a -lspe2 -lpthread -lm -lnuma
#
# INSTALL_DIR= $(EXP_SDKBIN)/demos
# INSTALL_FILES= $(PROGRAM_ppu)
# LDFLAGS_gcc    = -Wl,-q
# CFLAGS_gcc = -g
#
# ###########################################################################
# # buildutils/make.footer
# ###########################################################################
#
ifdef CELL_TOP
   include $(CELL_TOP)/buildutils/make.footer
else
   include ../../../buildutils/make.footer
endif
```

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In Example 6-2, we introduce the \texttt{-g} and \texttt{-Wl,-q} compilation flags to preserve the relocation and line number information in the final integrated executable.

\textbf{Example 6-2  Changing \texttt{-/FFT16M/ppu/Makefile} for gcc}

%**************************************************
# Target
%**************************************************
#
PROGRAM\_ppu= fft

%**************************************************
# Objects
%**************************************************
#
PPU\_COMPILER = xlc

IMPORTS = ../spu/fft\_spu.a -lspe2 -lpthread -lm -lnuma

#INSTALL\_DIR= $(EXP\_SDKBIN)/demos
#INSTALL\_FILES= $(PROGRAM\_ppu)
LDFLAGS\_xlc = -Wl,-q
CFLAGS\_xlc = -g

%**************************************************
# buildutils/make.footer
%**************************************************
#
ifdef CELL\_TOP
   include $(CELL\_TOP)/buildutils/make.footer
else
   include ../../../../buildutils/make.footer
endif
Modifying the ~/FFT16M/spu/Makefile

In Example 6-3 and Example 6-4 on page 422, we introduce the -g and -Wl,-q compilation flags in order to preserve the relocation and the line number information in the final integrated executable.

Example 6-3  Modifying ~/FFT16M/spu/Makefile for gcc

```plaintext
# Target

PROGRAMS_spu:= fft_spu
LIBRARY_embed:= fft_spu.a

# Local Defines

CFLAGS_gcc:= -g --param max-unroll-times=1 # needed to keep size of program down
LDFLAGS_gcc    = -Wl,-q -g
```

```plaintext
ifdef CELL_TOP
    include $(CELL_TOP)/buildutils/make.footer
else
    include ../../../buildutils/make.footer
endif
```
Example 6-4   Modifying ~/FFT16M/spu/Makefile for xlc

```
# Target
SPU_COMPILER = xlc
PROGRAMS_spu:= fft_spu
LIBRARY_embed:= fft_spu.a

CFLAGS_xlc:= -g -qnounroll -O5
LDFLAGS_xlc:= -O5 -qflag=e:e -Wl,-q -g
```

Before the actual build, set the default compiler accordingly by entering the following command:
```
/opt/cell/sdk/buildutils/cellsdk_select_compiler [gcc|xlc]
```

Now we are ready for the build and enter the following command:
```
cd ~/FFT16M ; CELL_TOP=/opt/cell/sdk make
```
6.3 Creating and working with the profile data

Assuming that you have properly set up the project tree and had a successful build, you collect and work with the profile data.

6.3.1 Step 1: Collecting data with CPC

Before collecting the application data, execute a small test to verify that the cell-perf-counter (CPC) tool is properly working. Type the following command to measure clock-cycles and branch instructions committed on both hardware threads for all processes on all processors for 5 seconds:

cpc --cpus all --time 5s --events C

You should immediately see counter statistics.

Given that CPC is properly working, we collect counter data for the FFT16M application. The following example counts PowerPC instructions committed in one event-set. It also counts L1 cache load misses in a second event-set and writes the output in the XML format (suitable for counter analyzer) to the file fft_cpc.pmf:

cd ~/FFT16M

cpc --events C,2100,2119 --cpus all ---xml fft_cpc.pmf ./ppu/fft 40 1

As the result, you should have the ~/FFT16M/fft_cpc.pmf file.

6.3.2 Step 2: Visualizing counter information using Counter Analyzer

The generated counter information can now be visualized by using the Counter Analyzer tool in VPA:

1. Open VPA.
2. Select Tools → Counter Analyzer.
3. Choose File → Open File.
4. In the Open File window, locate the fft_cpc.pmf file and select it.
Figure 6-1 shows the result of the collected counter information.

6.3.3 Step 3: Collecting data with OProfile

In the following steps, we generate appropriate profile information (suitable for Profile Analyzer) for both PowerPC Processor Unit (PPU) and Synergistic Processor Unit (SPU), from the FFT16M application:
1. Initialize the OProfile environment for SPU and run the fft workload to collect SPU average cycle events as shown in Example 6-5.

   **Example 6-5  Initializing and running OProfile for SPU profiling**

   ```
   # As root
   opcontrol --deinit
   opcontrol --start-daemon
   opcontrol --init
   opcontrol --reset
   opcontrol --separate=all --event=SPU_CYCLES:100000
   opcontrol --start
   # As regular user
   fft 20 1
   # As root
   opcontrol --stop
   opcontrol --dump
   ```

   To generate the report, enter the following command:

   `opreport -X -g -l -d -o fft.spu.opm`

2. Repeat the steps for PPU as shown in Example 6-6.

   **Example 6-6  Initializing and running OProfile for PPU profiling**

   ```
   # As root
   opcontrol --deinit
   opcontrol --start-daemon
   opcontrol --init
   opcontrol --reset
   opcontrol --separate=all --event=CYCLES:100000
   opcontrol --start
   # As regular user
   fft 20 1
   # As root
   opcontrol --stop
   opcontrol --dump
   ```

   To generate the report, enter the following command:

   `opreport -X -g -l -d -o fft.ppu.opm`
6.3.4 Step 4: Examining profile information with Profile Analyzer

Next, load the generated profile information with Profile Analyzer:

1. Open VPA.
2. Select **Tools** → **Profile Analyzer**.
3. Choose **File** → **Open File**.
4. In the Open File window, locate the `fft.spu.opm` file and select it.

Figure 6-2 shows the window that opens.

![fft.spu.opm in Profile Analyzer](image-url)
5. Examine the disassembly information.

a. Select the `fft_spu` entry in the Modules section (center of the Profile Analyzer window in Figure 6-3) and double-click `main` under the Symbol/Functions view (right side of Figure 6-3). The result is displayed in the Disassembly view in the lower right pane of the window.

![Figure 6-3 Disassembly view for fft_spu.opm](image)
b. The tool might prompt you for that particular symbol’s source code. If you have the code, you can switch to the **Source Code** tab (Figure 6-4) in the lower right pane of the window.

![Figure 6-4   Source view for fft_spu.opm](image)

If desired, you can repeat this procedure to analyze the fft_ppu.opm profile results.

### 6.3.5 Step 5: Gathering profile information with FDPR-Pro

The principal function of the FDPR-Pro tool is optimization. In addition, by using this tool, you can investigate the application performance, while mapping back to the source code, when used in combination with the Code Analyzer.

We use the FDPR-Pro instrumentation to collect the profiling data:

1. Clean up the old profile information and create a temporary working directory for FDPR-Pro:
   ```
   cd ~/FFT16M/ppu ; rm -f *.mprof *.nprof ; mkdir sputmp
   ```
2. Instrument the fft executable by entering the following command:
   ```
   fdprpro fft -cell -spedir sputmp -a instr
   ```
Example 6-7 shows the output of the command.

Example 6-7  Sample output from FDPR-Pro instrumentation

FDPR-Pro Version 5.4.0.16 for Linux (CELL)
fdprpro ./fft -cell -spedir sputmp -a instr
> spe_extraction -> ./fft ... 
... 
> processing_spe_file -> sputmp/fft_spu ... 
... 
> reading_exe ... 
> adjusting_exe ... 
... 
> analyzing ... 
> building_program_infrastructure ... 
@Warning: Relocations based on section .data -- section may not be reordered
> building_profiling_cfg ... 
> spe_encapsulation -> sputmp/out ... 
>> processing_spe -> sputmp/out/fft_spu ... 
> instrumentation ... 
>> throw&_catch_fixer ... 
>> adding_universal_stubs ... 
>> running_markers_and_instrumenters ... 
>> linker_stub_fixer ... 
>> dynamic_entries_table_sections_bus_fixer ... 
>> writing_profile_template -> fft.nprof ... 
> symbol_fixer ... 
> updating_executable ... 
> writing_executable -> fft.instr ... 
bye.

3. Run the generated instrumented profile:

./fft.instr 20 1

There should be two relevant generated files:

- ~/FFT16M/ppu/fft.nprof #, which contains PPU profile information
- ~/FFT16M/ppu/fft_spu.mprof #, which contains SPU profile information
6.3.6 Step 6: Analyzing profile data with Code Analyzer

The Code Analyzer tool imports information from the FDPR-Pro and creates a visualization for it, which you can use by following these steps:

1. With VPA open, select **Tools → Code Analyzer → Analyze Executable**.
2. In the window that opens, locate the original fft executable file.

In the center of the window (Figure 6-5), two editor tabs are displayed: one for PPU and one for SPU.

![Figure 6-5  PPU and SPU editors in Code Analyzer](image)
3. Associate the PPU profile information:
   a. Select the **PPU** editor tab view.
   b. From the menu bar, select **File → Code Analyzer → Add Profile Info** (Figure 6-6).
   c. Locate the **fft.nprof** file.

![Figure 6-6   Adding profile info](image)

4. Repeat the same procedure in step 3 for the SPU part. This time, click the **SPU** editor tab and locate the **fft_spu.mprof** file.
Immediately after loading the profile information, you see the colors of the instructions on both editor tabs (Figure 6-7). Red indicates highly frequently executed instructions.

![Code Analyzer showing rates of execution](image)

*Figure 6-7  Code Analyzer showing rates of execution*
Code Analyzer offers a variety of ways to use the code:

- In addition to the instructions, associate the source code.
  
  a. Select a symbol in the program tree, right-click and select **Open Source Code**.

  b. Locate the proper source code.

As a result, the Source Code tab is displayed at the center of the window (Figure 6-8), where you can see rates of execution per line of source code.

![Figure 6-8  Code Analyzer source code tab](image-url)
Calculate dispatch grouping boundaries for both fft PPE and fft SPU tabs. Select each tab and click the **Collect display information about dispatch groups** button (Figure 6-9).

![Collect & display information about dispatch groups](image)

You can simultaneously click the **Collect hazard info** button to collect comments about performance bottlenecks, above source lines that apply. See Figure 6-10.

![Hazard Info commented source code](image)
Display pipeline population for each dispatch group. Click the **Dispatch Info** tab in the lower right pane (the Instruction Properties tab) and click the **Link with table** button. See Figure 6-11.

**Figure 6-11** Dispatch Info tab with “Link with Table” option
The Latency Info tab, in the lower right pane (Figure 6-12), displays latencies for each selected instruction.

![Figure 6-12 Latency Info view](image)

- The Code Analyzer also offers the possibility of inspecting SPU Timing information, at the pipeline level, with detailed stages of the Cell pipeline population. Select the fft SPU editor tab, locate the desired symbol in the program tree, right-click, and select Show SPU-Timing as shown in Figure 6-13.

![Figure 6-13 Selecting SPU-Timing information](image)
6.4 Creating and working with trace data

The PDT tool produces tracing data, which can be viewed and analyzed in the Trace Analyzer tool. To properly collect trace data, we must recompile the fft application according to the required PDT procedures.
6.4.1 Step 1: Collecting trace data with PDT

To collect trace data with PDT:

1. Prepare the SPU makefile according to the PDT requirements as shown in Example 6-8 and Example 6-9 on page 439, depending on the compiler of your choice.

Example 6-8 Modifying ~/FFT16M/spu/Makefile for gcc compiler

```bash
# Target
PROGRAMS_spu:= fft_spu
LIBRARY_embed:= fft_spu.a

# Local Defines
CFLAGS_gcc:= -g --param max-unroll-times=1 -Wall -Dmain=_pdt_main
-Dexit=_pdt_exit -DMFCIO_TRACE -DLIBSYNC_TRACE
LDFLAGS_gcc    = -Wl,-q -g -L/usr/spu/lib/trace
INCLUDE                = -I/usr/spu/include/trace
IMPORTS                = -ltrace

# buildutils/make.footer
```
Example 6-9  Modifying ~/FFT16M/spu/Makefile for xlc compiler

#########################################################################
#                           #
#       Target             #
#########################################################################
#
SPU_COMPILER = xlc
PROGRAMS_spu:= fft_spu
LIBRARY_embed:= fft_spu.a

#########################################################################
#                           #
#       Local Defines      #
#########################################################################
#
CFLAGS_xlc:= -g -qnounroll -O5
CPP_FLAGS_xlc := -I/usr/spu/include/trace -Dmain=_pdt_main
               -Dexit=_pdt_exit -DMFCIO_TRACE -DLIBSYNC_TRACE
LDFLAGS_xlc:= -O5 -qflag=e:e -Wl,-q -g -L/usr/spu/lib/trace -ltrace

#########################################################################
#                           #
#       buildutils/make.footer                            #
#########################################################################
#
ifdef CELL_TOP
   include $(CELL_TOP)/buildutils/make.footer
else
   include ../../../../../buildutils/make.footer
endif
2. Rebuild the fft application:
   
   cd ~/FFT16M ; CELL_TOP=/opt/cell/sdk make

3. To focus on stalls, which we strongly recommend, set up a configuration file with only the relevant stalls (mailboxes and read tag status for SPE):
   
   a. Copy the default XML to where the FFT is run, so that we can modify it:
      
      cp /usr/share/pdt/config/pdt_cbe_configuration.xml ~/FFT16M
   
   b. Open the copied file for editing. On the first line, change the application name value to “fft”.
   
   c. Search for <configuration name="SPE">. Below that line is the MFCIO group tag. Set it to active="false".
   
   d. Delete the SPE_MFC group. This should be sufficient to trace only the stalls in the SPE.

4. Prepare the environment by setting the following variables:
   
   export LD_LIBRARY_PATH=/usr/lib/trace
   export PDT_KERNEL_MODULE=/usr/lib/modules/pdt.ko
   export PDT_CONFIG_FILE=~/FFT16M/pdt_cbe_configuration.xml

5. Run the fft application at least three times for better sampling:
   
   cd ~/FFT16M/ppu ; ./fft 1 1 4 1 0
   
   You should have the three trace files, .pex, .map and .trace, that are available right after the execution.

Notes:

- The default PDT_CONFIG_FILE for the SDK establishes the trace files prefix as “test.” If you have not modified the file, look for the trace files with “test” as the prefix.

- Remember to unset the LD_LIBRARY_PATH environment variable, before running the original (non-PDT) binary later.
6.4.2 Step 2: Importing the PDT data into Trace Analyzer

The Trace Analyzer allows the visualization of the application's stages of execution. It works with data generated from the PDT tool. More specifically, it reads information that is available in the generated .pex file. To visualize the data on the Trace Analyzer:

1. With VPA open, select **Tools → Trace Analyzer**.
2. Select **File → Open File**.
3. In the Open File window, locate the .pex file that was generated in the previous steps.

The window shown in Figure 6-15 on page 441 is displayed.

![Trace Analyzer window](image)

Figure 6-15 Trace Analyzer window

This window corresponds to the FFT16M application that was run with 16 SPEs and no huge pages. As we can observe, a less intensive blue has been selected
for the MFC_IO group. We now see the difference between the borders and the internals of the interval. Additionally we use the color map to change the color of SPE_READ_IN_MBOX to be red rather than the default blue for its group. You see a huge stall in the middle. This is where the benchmark driver verifies the result of the test run to make sure the benchmark computes correctly. The timed run is the thin blue strip after the stall.

Next we zoom into this area, which is all that interests us in this benchmark. As we can observe in Figure 6-16, the mailboxes (red bars) break the execution into six stages. The different stages have different behaviors. For example, the third and sixth stages are much longer than the rest and have a lot of massive stalls.
By using Trace Analyzer, we can select a stall to obtain further details by clicking the stall (the yellow highlight circled in the SPE10 row in Figure 6-16). The selection marker rulers on the left and top show the location of the selected item and can be used to return to it if you scroll away. The data collected by the PDT for the selected stall is shown in the Record Details tab in the lower right corner of Figure 6-16. The stall is huge at almost 12K ticks. We check the Cell/B.E. performance tips for the cause of the stall and determine that translation look-aside buffers (TLB) misses are a possible culprit and huge pages are a possible fix.

By looking at this trace visualization sample, we can discover a significant amount of information regarding potential application problems. It is possible to observe how well balanced the application is by looking at the execution and start/stop time for each SPU. Since the visualization is broken down by type, which are the cause of stalls in the code, we can identify synchronization problems.
Chapter 7. Programming in distributed environments

In this chapter, we provide an overview of a few of the distributed programming techniques that are available on the Cell Broadband Engine (Cell/B.E.) processor. We introduce the Hybrid Programming Model, as well as its libraries in IBM Software Developer Kit (SDK) 3.0, and Dynamic Application Virtualization (DAV).

Specifically, we discuss the following topics:

- 7.1, “Hybrid programming models in SDK 3.0” on page 446
- 7.2, “Hybrid Data Communication and Synchronization” on page 449
- 7.3, “Hybrid ALF” on page 463
- 7.4, “Dynamic Application Virtualization” on page 475
7.1 Hybrid programming models in SDK 3.0

The Cell Broadband Engine Architecture (CBEA) is one answer for the problems that the computer industry faces in regard to performance degradation on traditional single-threaded-styled solutions. The power, frequency, and memory wall problems lead to the use of multi-core solutions and the exploitation of memory hierarchies enforcing the data locality.

Figure 7-1 [27 on page 625] shows the slow down in single thread performance growth.

![Image: SPECINT: Slowdown in single thread performance growth]

Figure 7-1  Single thread performance

Although the Cell/B.E. processor has been performing in an outstanding manner for certain types of applications, we must consider other requirements in this picture. For example, we must consider power balance, existing application integration, and larger cluster scenarios, where network latency has a great influence on the performance.
The Hybrid Model System architecture (Figure 7-2) proposal is a combination of characteristics from traditional superscalar multicore solutions and Cell/B.E. accelerated features. The idea is to use traditional, general purpose superscalar clusters of machines as “processing masters” (hosts) to handle large data partitioning and I/O bound computations (for example, message passing). Meanwhile the clusters off-load well-characterized, computational-intensive functions to computing kernels running on Cell/B.E. accelerator nodes.

Figure 7-2   Hybrid Model System architecture

This solution enables finer grain control over the applications’ parallelism and a more flexible offering, because it easily accommodates Multiple Process Multiple Data (MPMD) tasks with Single Process Multiple Data (SPMD) tasks.

Motivations
There is a well known balance between generality and performance. While the Cell/B.E. processor is not a general all-purpose solution, it maintains strong general purpose capabilities, especially with the presence of the Power Processor Element (PPE). See Figure 7-3.

Figure 7-3   Balance between special and general purpose solutions
To understand the motivations behind the Hybrid architecture, we must understand a few high performance scenario requirements:

- When moving from single-threaded to multi-core solutions, although there is a significant performance boost (throughput), there might be a power (energy) demand increase.
- Legacy solutions based on homogenous architectures exist.
- Some applications need finer grain parallel computation control, because different components have different computational profiles.

The Hybrid Model System architecture addresses these requirements in the following ways:

- Increasing the throughput performance with the addition of the accelerators (Cell/B.E. system), in a more energy efficient way
  
  Since the Cell/B.E. processor has outstanding performance for computation-specific parallel tasks, given the same energy footprint, the whole system performs better when kept as a homogeneous solution (Figure 7-4).

- Accommodating existing solutions, which includes general purpose units
- Achieving finer grain parallelism, because it is capable of mixing MPMD and SPMD at different levels (host and accelerator)
7.2 Hybrid Data Communication and Synchronization

The Hybrid version of the Data Communication and Synchronization (DaCS) library provides a set of services that eases the development of applications and application frameworks in a heterogeneous multi-tiered system (for example a 64 bit x86 system (x86_64) and one or more Cell/B.E. systems). The DaCS services are implemented as a set of application programming interfaces (APIs) that provide an architecturally neutral layer for application developers on a variety of multi-core systems.

One of the key abstractions that further differentiates DaCS from other programming frameworks is a hierarchical topology of processing elements, each referred to as a DaCS Element (DE). Within the hierarchy, each DE can serve one or both of the following roles:

- A general purpose processing element, acting as a supervisor, control, or master processor
  This type of element usually runs a full operating system and manages jobs running on other DEs. This element is referred to as a host element (HE).

- A general or special purpose processing element running tasks assigned by an HE, which is referred to as an accelerator element (AE)

Figure 7-5   Hybrid DaCS system
7.2.1 DaCS for Hybrid implementation

The DaCS for Hybrid (DaCSH) architecture (Figure 7-6) is an implementation of the DaCS API specification, which supports the connection of an HE on an x86_64 system to one or more AEs on Cell/B.E. systems. In SDK 3.0, DaCSH only supports the use of sockets to connect the HE with the AEs. DaCSH provides access to the PPE, allowing a PPE program to be started and stopped and allowing data transfer between the x86_64 system and the PPE. The SPEs can only be used by the program running on the PPE.

![DaCS for Hybrid architecture](image)

A PPE program that works with the SPEs can also be a DaCS program. In this case, the program uses DaCS for Cell (DaCSC, see the Data Communication and Synchronization Library for Cell Broadband Engine Programmer's Guide and API Reference). The PPE acts as an AE for DaCSH (communicating with the x86_64 system) and as an HE for DaCSC (communicating with the SPEs). The DaCS API on the PPE is supported by a combined library. When the PPE is used with both DaCSH and DaCSC, the library automatically uses the parameters that

---

are passed to the API to determine if the PPE is an AE communicating with its HE (DaCSH) or an HE communicating with its AEs (DaCSC).

To manage the interactions between the HE and the AEs, DaCSH starts a service on each of them. On the host system, the service is the host DaCS daemon (hdacsd). On the accelerator, the service is the accelerator DaCS daemon (adacsd). These services are shared between all DaCSH processes for an operating system image.

For example, consider an x86_64 system that has multiple cores that each run a host application using DaCSH. In this case, only a single instance of the hdacsd service is needed to manage the interactions of each of the host applications with their AEs via DaCSH. Similarly, on the accelerator, if the Cell/B.E. system is on a blade server (which has two Cell/B.E. processors), a single instance of the adacsd service is needed to manage both of Cell/B.E. systems acting as AEs, even if they are used by different HEs.

When a host application starts using DaCSH, it connects to the hdacsd service. This service manages the system topology from a DaCS perspective (managing reservations) and starts the accelerator application on the AE. Only process management requests use the hdacsd and adacsd services. All other interactions between the host and accelerator application flow via a direct socket connection.

**Services**

The DaCS services can be divided into the following categories:

- **Resource reservation**
  The resource reservation services allow an HE to reserve AEs below itself in the hierarchy. The APIs abstract the specifics of the reservation system (operating system, middleware, and so on) to allocate resources for an HE. When reserved, the AEs can be used by the HE to execute tasks for accelerated applications.

- **Process management**
  The process management services provide the means for an HE to execute and manage accelerated applications on AEs, including remote process launch and remote error notification among others. The hdacsd provides services to the HE applications. The adacsd provides services to the hdacsd and HE application, including the launching of the AE applications on the accelerator for the HE applications.
- **Group management**
  The group management services provide the means to designate dynamic groups of processes for participation in collective operations. In SDK 3.0, this is limited to process execution synchronization (barrier).

- **Remote memory**
  The remote memory services provide the means to create, share, transfer data to, and transfer data from a remote memory segment. The data transfers are performed by using a one-sided put or get remote direct memory access (rDMA) model. These services also provide the ability to scatter/gather lists of data and provide optional enforcement of ordering for the data transfers.

- **Message passing**
  The message passing services provide the means for passing messages asynchronously, using a two-sided send/receive model. Messages are passed point-to-point from one process to another.

- **Mailboxes**
  The mailbox services provide a simple interface for synchronous transfer of small (32-bit) messages from one process to another.

- **Process synchronization**
  The process synchronization services provide the means to coordinate or synchronize process execution. In SDK 3.0, this is limited to the barrier synchronization primitive.

- **Data synchronization**
  The data synchronization services provide the means to synchronize and serialize data access. These include management of wait identifiers for synchronizing data transfers, as well as mutex primitives for data serialization.

- **Error Handling**
  The error handling services enable the user to register error handlers and gather error information.

### 7.2.2 Programming considerations

The DaCS library API services are provided as functions in the C language. The protocols and constants that are required are made available to the compiler by including the DaCS header file `dacs.h` as follows:

```c
#include <dacs.h>
```

In general, the return value from these functions is an error code. Data is returned within parameters that are passed to the functions.
Process management model
When working with the host and accelerators, there must be a way to uniquely identify the participants that are communicating. From an architectural perspective, each accelerator can have multiple processes simultaneously running. Therefore, it is not enough to only identify the accelerator. Instead the unit of execution on the accelerator (the DaCS process) must be identified by using its DaCS element ID (DE ID) and its process ID (Pid). The DE ID is retrieved when the accelerator is reserved (by using dacs_reserve_children()) and the Pid when the process is started (by using dacs_de_start()).

Since the parent is not reserved, and no process is started on it, two constants are provided to identify the parent: DACS_DE_PARENT and DACS_PID_PARENT. Similarly, to identify the calling process itself, the constants DACS_DE_SELF and DACS_PID_SELF are provided.

Resource sharing model
The APIs that support the locking primitives, remote memory access, and groups follow a consistent pattern of creation, sharing, usage, and destruction:

► Creation
An object is created that is shared with other DEs, for example with dacs_remote_mem_create().

► Sharing
The object created is then shared by linked share and accept calls. The creator shares the item, for example, with dacs_remote_mem_share(). The DE with which it is shared accepts it, which in this example, is with dacs_remote_mem_accept(). These calls must be paired. When one is invoked, it waits for the other to occur. This is done for each DE that the share is action with.

► Usage
Usage might require closure (such as in the case of groups), or the object might immediately be available for use. For example, remote memory can immediately be used for put and get.

► Destruction
The DEs that have accepted an item can release the item when they are done with it, for example, by calling dacs_remote_mem_release(). The release does not block, but notifies the creator that it is no longer being used and cleans up any local storage. The creator does a destroy (in this case dacs_remote_mem_destroy()) that waits for all of the DEs it has shared with to release the item and then destroys the shared item.
API environment
To make these services accessible to the runtime code, each process must create a DaCS environment. This is done by calling the special initialization service dacs_runtime_init(). When this service returns, the environment is set up so that all other DaCS function calls can be invoked. When the DaCS environment is no longer required, the process must call dacs_runtime_exit() to free all resources that are used by the environment.

Important: DaCS for Hybrid and DaCS for the Cell/B.E. system share the same API set, although they are two different implementations. For more information about DaCS API, refer to 4.7.1, “Data Communication and Synchronization” on page 291.

7.2.3 Building and running a Hybrid DaCS application

Three versions of the DaCS libraries are provided with the DaCS packages:

- Optimized libraries
  The optimized libraries have minimal error checking and are intended for production use.

- Debug libraries
  The debug libraries have much more error checking than the optimized libraries and are intended to be used during application development.

- Traced libraries
  The traced libraries are the optimized libraries with performance and debug trace hooks in them. These are intended to be used to debug functional and performance problems that might occur. The traced libraries use the interfaces that are provided by the Performance Debug Tool (PDT) and require that this tool be installed.

Both static and shared libraries are provided for the x86_64 and PPU. The desired library is selected by linking to the chosen library in the appropriate path. The static library is named libdacs.a, and the shared library is named libdacs.so.
**DaCS configuration**

The hdacsd and adacsd are both configured by using their respective /etc/dacsd.conf files. The default versions of these files are automatically installed with each of the daemons. These default files contain comments on the parameters and values that are currently supported.

When one of these files is changed, the changes do not take affect until the respective daemon is restarted as described previously.

**DaCS topology**

The topology configuration file /etc/dacs_topology.config is only used by the host daemon service. Ensure that you back up this file before you change it. Changes do not take effect until the daemon is restarted.

The hdacsd might stop if there is a configuration error in the dacs_topology.config file. Check the log file specified by the dacsd.conf file (default is /var/log/hdacsd.log) for configuration errors.

The topology configuration file identifies the hosts and accelerators and their relationship to one another. The host can contain more than one processor core, for example a multicore x86_64 blade. The host can be attached to one or more accelerators, for example a Cell/B.E. blade. By using the topology configuration file, you can specify a number of configurations for this hardware. For example, it can be configured so that each core is assigned one Cell/B.E. accelerator or it might be configured so that each core can reserve any (or all) of the Cell/B.E. accelerators.

The default topology configuration file, shown in Example 7-1, is for a host that has four cores and is attached to a single Cell/B.E. blade.

**Example 7-1  Default topology file**

```
<DaCS_Topology
 version="1.0">
 <hardware>
  <de tag="OB1" type="DACS_DE_SYSTEMX" ip="192.168.1.100">
   <de tag="OC1" type="DACS_DE_SYSTEMX_CORE"></de>
   <de tag="OC2" type="DACS_DE_SYSTEMX_CORE"></de>
   <de tag="OC3" type="DACS_DE_SYSTEMX_CORE"></de>
   <de tag="OC4" type="DACS_DE_SYSTEMX_CORE"></de>
  </de>
  <de tag="CB1" type="DACS_DE_CELLBLADE" ip="192.168.1.101">
   <de tag="CBE11" type="DACS_DE_CBE"></de>
   <de tag="CBE12" type="DACS_DE_CBE"></de>
  </de>
 </hardware>
</DaCS_Topology>
```
The `<hardware>` section identifies the host system with its four cores (OC1-OC4) and the Cell/B.E. BladeCenter (CB1) with its two Cell/B.E. systems (CBE11 and CBE12).

The `<topology>` section identifies what each core (host) can use as an accelerator. In this example, each core can reserve and use either the entire Cell/B.E. BladeCenter (CB1) or one or more of the Cell/B.E. systems on the BladeCenter.

The ability to use the Cell/B.E. system is implicit in the `<canreserve>` element. This element has an attribute that defaults to `false`. When it is set to true, only the Cell/B.E. BladeCenter can be reserved. If the fourth `<canreserve>` element was changed to `<canreserve he="OC4" ae="CB1" only="TRUE"/></canreserve>`, then OC4 can only reserve the Cell/B.E. BladeCenter. The usage can be made more restrictive by being more specific in the `<canreserve>` element. If the fourth `<canreserve>` element is changed to `<canreserve he="OC4" ae="CBE12"/></canreserve>`, then OC4 can only reserve CBE12 and cannot reserve the Cell/B.E. BladeCenter.

Modify the topology configuration file to match your hardware configuration. Make a copy of the configuration file before changing it. At a minimum, update the IP addresses of the ip attributes to match the interfaces between the host and the accelerator.

**DaCS daemons**

The daemons can be stopped and started by using the shell `service` command in the `sbin` directory. For example, to stop the host daemon, type the following command:

```
/sbin/service hdacsd stop
```

To restart the host daemon, type the following command:

```
/sbin/service hdacsd start
```

The adacsd can be restarted in a like manner. See the man page for `service` for more details about the service command.
Running an application
A hybrid DaCS application on the host (x86_64) must have CPU affinity to start, which can be done on the command line. The following command line example shows how to set affinity of the shell to the first processor:

```
taskset -p 0x00000001 $$
```

The bit mask, starting with 0 from right to left, is an index to the CPU affinity setting. Bit 0 is on or off for CPU 0, bit 1 for CPU 1, and bit number x is CPU number x. $$ means the current process gets the affinity setting. The following command returns the mask setting as an integer:

```
taskset -p$$
```

You can use the -c option to make `taskset` more usable. For example, the following command sets the processor CPU affinity to CPU 3 for the current process:

```
taskset -pc 3 $$
```

The -pc parameter sets by process and CPU number. The following command returns the current processor setting for affinity for the current process:

```
taskset -pc $$
```

According to the man page for `taskset`, a user must have CAP_SYS_NICE permission to change CPU affinity. See the man page for `taskset` for more details.

To launch a DaCS application, use a `taskset` call as shown in the following example, where the application program is MyDaCSApp and is passed an argument of arg1:

```
taskset 0x00000001 MyDaCSApp arg1
```
7.2.4 Step-by-step example

In this section, we create a simple Hybrid DaCS Hello World application, from building the application to deploying it.

Step 1: Verifying the configuration

To properly build and run this example, we must verify the following requirements on the configuration:

- You need one x86_64 blade server and one QS21 Cell/B.E. blade, both configured with the SDK 3.0.
- Verify that you have properly configured DaCS topology files in your x86_64 node.
- Make sure hdacsd is started on the x86_64 and adacsd is started on the QS21 blade.

Step 2: Creating the build structure

Although the build process occur on the host (x86_64) machine, create the directory structure shown in Example 7-2, under the root of your user home directory, on both the host and accelerator machines.

Example 7-2  Directory structure

```
hdacshello
hdacshello/ppu
hdacshello/ppu/spu
```

We also need a makefile in each of the created folders (only on the host machine), which we create as shown in Example 7-3, Example 7-4 on page 459, and Example 7-5 on page 459.

Example 7-3  hdacshello/Makefile

```
DIRS := ppu
INCLUDE := -I/opt/cell/sdk/prototype/usr/include
IMPORTS := /opt/cell/sdk/prototype/usr/lib64/libdacs_hybrid.so
CC_OPT_LEVEL := -g
PROGRAM := hdacshello
include $(CELL_TOP)/buildutils/make.footer
```
Example 7-4  hdacshello/ppu/Makefile

DIRS := spu
INCLUDE := -I/opt/cell/sdk/sysroot/usr/include
IMPORTS :=
/opt/cell/sysroot/opt/cell/sdk/prototype/usr/lib64/libdacs_hybrid.so
spu/hdacshello_spu
LDFLAGS += -lstdc++
CC_OPT_LEVEL = -g
PROGRAM_ppu64 := hdacshello_ppu
include $(CELL_TOP)/buildutils/make.footer

Example 7-5  hdacshello/ppu/spu/Makefile

INCLUDE := -I/opt/cell/sdk/sysroot/usr/include
IMPORTS := /opt/cell/sysroot/usr/spu/lib/libdacs.a
CC_OPT_LEVEL := -g
PROGRAM_spu := hdacshello_spu
include $(CELL_TOP)/buildutils/make.footer

Step 3: Creating the source files
Create the source files as shown in Example 7-6, Example 7-7 on page 460, and Example 7-8 on page 462 for each part of the application (on the host machine).

Example 7-6  hdacshello/hdacshello.c

#include <dacs.h>
#include <stdlib.h>
#include <string.h>
#include <stdio.h>
de_id_t cbe[2];
dacs_process_id_t pid;

int main(int argc __attribute__((unused)), char* argv[] __attribute__((unused)))
{
    DACS_ERR_T dacs_rc;
    dacs_rc = dacs_runtime_init(NULL,NULL);
    printf("HOST: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);
    uint32_t num_cbe = 1;
```c
    dacs_rc = dacs_reserve_children(DACS_DE_CBE,&num_cbe,cbe);
    printf("HOST: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);
    printf("HOST: %d : num children = %d, cbe = %08x\n",__LINE__,num_cbe,cbe[0]);
    fflush(stdout);
    char const * argp[] = {0};
    char const * envp[] = {0};
    char program[1024];
    getcwd(program,sizeof(program));
    strcat(program,"/ppu/hdacshello_ppu");

    dacs_rc =
    dacs_de_start(cbe[0],program,argp,envp,DACS_PROC_REMOTE_FILE,&pid);
    printf("HOST: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    int32_t status = 0;
    dacs_rc = dacs_de_wait(cbe[0],pid,&status);
    printf("HOST: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    dacs_rc = dacs_release_de_list(num_cbe,cbe);
    printf("HOST: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    dacs_rc = dacs_runtime_exit();
    printf("HOST: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    return 0;
}
```

Example 7-7   hdacshello/ppu/hdacshello_ppu.c

```c
#include <dacs.h>
#include <libspe2.h>
#include <malloc.h>
#include <inttypes.h>
#include <string.h>
#include <stdio.h>
#include <stdlib.h>
```
extern spe_program_handle_t hdacshello_spu;

de_id_t spe[2];
dacs_process_id_t pid;

int main(int argc __attribute__((unused)), char* argv[] __attribute__((unused)))
{
    DACS_ERR_T dacs_rc;
    dacs_rc = dacs_runtime_init(NULL,NULL);
    printf("PPU: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    uint32_t num_spe = 1;
    dacs_rc = dacs_reserve_children(DACS_DE_SPE,&num_spe,spe);
    printf("PPU: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);
    printf("PPU: %d : num children = %d, spe = %08x\n",__LINE__,num_spe,spe[0]);
    fflush(stdout);

    char const * argp[] = {0};
    char const * envp[] = {0};
    void * program;
    program = &hdacshello_spu;
    DACS_PROC_CREATION_FLAG_T flags = DACS_PROC_EMBEDDED;
    dacs_rc = dacs_de_start(spe[0],program,argp,envp,flags,&pid);
    printf("PPU: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    int32_t status = 0;
    dacs_rc = dacs_de_wait(spe[0],pid,&status);
    printf("PPU: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    dacs_rc = dacs_release_de_list(num_spe,spe);
    printf("PPU: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    dacs_rc = dacs_runtime_exit();
    printf("PPU: %d : rc = %s\n",__LINE__,dacs_strerror(dacs_rc));
    fflush(stdout);

    return 0;
}
Example 7-8  hdacshello/ppu/spu/hdacshello_spu.c

```c
#include <dacs.h>
#include <stdio.h>
#include <stdlib.h>

int main(int argc __attribute__((unused)), char* argv[] __attribute__((unused)))
{
    DACS_ERR_T dacs_rc;

    dacs_rc = dacs_runtime_init(NULL, NULL);
    printf("SPU: %d : rc = %s\n", __LINE__, dacs_strerror(dacs_rc));
    fflush(stdout);

    printf("Hello !\n"); fflush(stdout);

    dacs_rc = dacs_runtime_exit();
    printf("SPU: %d : rc = %s\n", __LINE__, dacs_strerror(dacs_rc));
    fflush(stdout);

    return 0;
}
```

Step 4: Building and deploying the files
Change to the topmost folder (hdacshello) and build the application as shown in Example 7-9.

Example 7-9  Building the application

```
cd ~/hdacshello
CELL_TOP=/opt/cell/sdk make
```

If everything proceeded as expected, the following binaries are available:

- hdacshello/hdacshello
- hdacshello/ppu/hdacshello_ppu
- hdacshello/ppu/spu/hdacshello_spu

Ensure that you have permission to execute each of binaries, for example, by using the following command:
```
chmod a+x ~hdacshello/hdacshello  # repeat on the other executables
```
Next, deploy the CBE binary hdacshello/ppu/hdacshello_ppu to the matching location on the accelerator (QS21 Cell Blade) machine. You can use `scp`, as shown in the following example:

```
scp ~/hdacshello/ppu/hdacshello_ppu user@qs21:~/hdacshello/ppu
```

**Step 5: Running the application**
Since Hybrid DaCS requires variables and commands to be run, create a helper script as shown in Example 7-10.

**Example 7-10  hdacshello/run.sh**

```bash
# Set the environment
export
LD_LIBRARY_PATH=/opt/cell/sdk/prototype/usr/lib64:$LD_LIBRARY_PATH
export
DACS_START_ENV_LIST="LD_LIBRARY_PATH=/opt/cell/sdk/prototype/usr/lib64:
_LD_LIBRARY_PATH"
# Set the shell's cpu affinity
taskset -pc 1 $$

# Launch the target program
~/hdacshello/hdacshello
```

Make sure that all daemons are properly running on the host and the accelerator, and execute the helper script:

```
~/hdacshello/run.sh
```

### 7.3 Hybrid ALF

There are two implementations of ALF:

- **ALF Cell implementation**
  The ALF Cell implementation executes on the Cell/B.E. PPU host and schedules tasks and work blocks on Cell/B.E. SPUs.

- **ALF Hybrid implementation**
  The ALF Hybrid implementation executes on an x86_64 host and schedules tasks and work blocks on an associated set of Cell/B.E. SPUs through a communications mechanism, which in this case uses the DaCS library.
7.3.1 ALF for Hybrid-x86 implementation

ALF for Hybrid-x86 is an implementation of the ALF API specification in a system configuration with an Opteron x86_64 system connected to one or more Cell/B.E. processors. In this implementation, the Opteron system serves as the host, the SPEs on the Cell/B.E. BladeCenter servers act as accelerators, and the PPEs on the Cell/B.E. processors act as facilitators only. From the ALF application programmer’s perspective, the application interaction, as defined by the ALF API, is between the Hybrid-x86 host and the SPE accelerators.

This implementation of the ALF API uses the DaCS library as the process management and data transport layer. Refer to 7.2.1, “DaCS for Hybrid implementation” on page 450, for more information about how to set up DaCS in this environment.

Figure 7-7 shows the hybrid ALF stack.

To manage the interaction between the ALF host run time on the Opteron system and the ALF accelerator run time on the SPE, this implementation starts a PPE process (ALF PPE daemon) for each ALF run time. The PPE program is provided as part of the standard ALF runtime library. Figure 7-8 on page 465 shows the hybrid ALF flow.
7.3.2 Programming considerations

As occurs with DaCS and Hybrid DaCS, ALF and Hybrid ALF are two different implementations of the same API set. For more information about the ALF programming model, refer to 4.7.2, “Accelerated Library Framework” on page 298.

7.3.3 Building and running a Hybrid ALF application

Three versions of the ALF for Hybrid-x86 libraries are provided with the SDK:

- Optimized
  The optimized library has minimal error checking on the SPEs and is intended for production use.

- Error-check enabled
  The error-check enabled version has a lot more error checking on the SPEs and is intended to be used for application development.

- Traced
  The traced version are the optimized libraries with performance and debug trace hooks in them. These version are intended for debugging functional and performance problems that are associated with ALF.

Additionally, both static and shared libraries are provided for the ALF host libraries. The ALF SPE runtime library is only provided as static libraries.
An ALF for Hybrid-x86 application must be built as two separate binaries as follows:

- The first binary is for the ALF host application, for which you must do the following actions:
  a. Compile the x86_64 host application with the
     -D_ALF_PLATFORM_HYBRID define variable, and specify the
     /opt/cell/sdk/prototype/usr/include directory.
  b. Link the x86_64 host application with the ALF x86_64 host runtime library,
     alf_hybrid, found in the /opt/cell/sdk/prototype/usr/lib64 directory and the
     DaCS x86_64 host runtime library, dacs_hybrid, also found in the
     /opt/cell/sdk/prototype/usr/lib64 directory.

- The second binary is for the ALF SPE accelerator computational kernel. You
  must perform the following actions:
  a. Compile the application’s SPE code with the
     -D_ALF_PLATFORM_HYBRID define variable, and specify the
     /opt/cell/sysroot/usr/spu/include and the
     /opt/cell/sysroot/opt/cell/sdk/prototype/usr/spu/include directories.
  b. Link the application’s SPE code with the ALF SPE accelerator runtime
     library, alf_hybrid, in the /opt/cell/sysroot/opt/cell/sdk/prototype/usr/spu/lib
     directory.
  c. Use the `ppu-embedspu` utility to embed the SPU binary into a PPE ELF
     image. The resulting PPE ELF object needs to be linked as a PPE shared
     library.

### 7.3.4 Running a Hybrid ALF application

**Library access:** Ensure that the dynamic libraries libalf_hybrid and
libdacs_hybrid are accessible. You can set this in `LD_LIBRARY_PATH` as
shown in the following example:

```bash
export LD_LIBRARY_PATH=/opt/cell/sdk/prototype/usr/lib64
```

To run an ALF application:

1. Build the ALF for Hybrid-x86 application. Build the host application as an
   executable, `my_appl`, and the accelerator computational kernel as a PPE
   shared library, `my_appl.so`. 
2. Copy the PPE shared library with the embedded SPE binaries from the host where it was built to a selected directory on the Cell/B.E. processor where it is to be executed, for example:

   scp my_appl.so <CBE>:/tmp/my_directory

3. Set the environment variable ALF_LIBRARY_PATH to the directory shown in step 2 on the Cell/B.E processor, for example:

   export ALF_LIBRARY_PATH=/tmp/my_directory

4. Set the CPU affinity on the Hybrid-x86 host, for example:

   taskset -p 0x00000001 $$

5. Run the x86_64 host application on the host environment, for example:

   ./my_appl

### 7.3.5 Step-by-step example

In this section, we show how to create and deploy a simple Hybrid ALF Hello World application.

**Step 1: Creating the build structure**

The build process occurs on the host (x86_64) machine. Create the directory structure shown in Example 7-11, under the root of your user home directory.

```
Example 7-11  Directory structure

halfhello
halfshello/host
halfhello/spu
```

We also need a makefile in each of the created folders. See Example 7-12, Example 7-13, and Example 7-14 on page 468.

```
Example 7-12  halfhello/Makefile

DIRS := spu host
include $(CELL_TOP)/buildutils/make.footer
```

```
Example 7-13  halfhello/spu/Makefile

INCLUDE := -I/opt/cell/sysroot/opt/cell/sdk/prototype/usr/spu/include
IMPORTS := -lalf_hybrid
LDFLAGS := -L/opt/cell/sysroot/opt/cell/sdk/prototype/usr/spu/lib
CPPFLAGS := -D_ALF_PLATFORM_HYBRID_
OBJS_alf_hello_world_sp := main_spu.o
```
PROGRAMS_spu := alf_hello_world_spu
SHARED_LIBRARY_embed64 := alf_hello_world_hybrid_sp64.so
include $(CELL_TOP)/buildutils/make.footer

Example 7-14  halfhello/host/Makefile

TARGET_PROCESSOR := host
INCLUDE := -I/opt/cell/sdk/prototype/usr/include
IMPORTS := -lalf_hybrid -lpthread -ldl -ldacs_hybrid -l numa -l stdc++ -lrt
LDFLAGS := -L/opt/cell/sdk/prototype/usr/lib64
CPPFLAGS := -D_ALF_PLATFORM_HYBRID_
PROGRAM := alf_hello_world_hybrid_host64
include $(CELL_TOP)/buildutils/make.footer

Step 2: Creating the source files

Create the source files for each part of the application as shown in Example 7-15 and Example 7-16 on page 473.

Example 7-15  Host source code (~/halfhello/host/main.c)

```c
#include <stdio.h>
#include <alf.h>
#include <unistd.h>
#include <stdlib.h>
#include <string.h>
#define DEBUG
#ifdef DEBUG
#define debug_print(fmt, arg...) printf(fmt,##arg)
#else
#define debug_print(fmt, arg...) { }
#endif
#define IMAGE_PATH_BUF_SIZE 1024    // Max length of path to PPU image
char spu_image_path[IMAGE_PATH_BUF_SIZE];   // Used to hold the complete path to SPU image
char library_name[IMAGE_PATH_BUF_SIZE];    // Used to hold the name of spu library
char spu_image_name[] = "alf_hello_world_sp64";
char kernel_name[] = "comp_kernel";
char input_dtl_name[] = "input_prep";
```
char output_dtl_name[] = "output_prep";

int main()
{
    int ret;
    alf_handle_t handle;
    alf_task_desc_handle_t task_desc_handle;
    alf_task_handle_t task_handle;
    alf_wb_handle_t wb_handle;
    void *config_parms = NULL;

    sprintf(library_name, "alf_hello_world_hybrid_spu64.so");

dbg_print("Before alf_init\n");

    if ((ret = alf_init(config_parms, &handle)) < 0) {
        fprintf(stderr, "Error: alf_init failed, ret=%d\n", ret);
        return 1;
    }

    dbg_print("Before alf_num_instances_set\n");
    if ((ret = alf_num_instances_set(handle, 1)) < 0) {
        fprintf(stderr, "Error: alf_num_instances_set failed, ret=%d\n", ret);
        return 1;
    } else if (ret > 0) {
        dbg_print("alf_num_instances_set returned number of SPUs=%d\n", ret);
    }

    dbg_print("Before alf_task_desc_create\n");
    if ((ret = alf_task_desc_create(handle, 0, &task_desc_handle)) < 0) {
        fprintf(stderr, "Error: alf_task_desc_create failed, ret=%d\n", ret);
        return 1;
    } else if (ret > 0) {
        dbg_print("alf_task_desc_create returned number of SPUs=%d\n", ret);
    }

    dbg_print("Before alf_task_desc_set_int32\n");
    if ((ret = alf_task_desc_set_int32(task_desc_handle, ALF_TASK_DESC_MAX_STACK_SIZE, 4096)) < 0) {
        fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);

return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_WB_PARM_CTX_BUF_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_WB_IN_BUF_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_WB_OUT_BUF_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_WB_INOUT_BUF_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_TSK_CTX_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_TSK_CTX_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int32\n");
if ((ret = alf_task_desc_set_int32(task_desc_handle,
ALF_TASK_DESC_TSK_CTX_SIZE, 0)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int32 failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_set_int64\n");
if ((ret = alf_task_desc_set_int64(task_desc_handle, ALF_TASK_DESC_ACCEL_IMAGE_REF_L, (unsigned long long)spu_image_name)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int64 failed, ret=%d\n", ret);
    return 1;
}

dump_print("Before alf_task_desc_set_int64\n");
if ((ret = alf_task_desc_set_int64(task_desc_handle, ALF_TASK_DESC_ACCEL_LIBRARY_REF_L, (unsigned long long)library_name)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int64 failed, ret=%d\n", ret);
    return 1;
}

dump_print("Before alf_task_desc_set_int64\n");
if ((ret = alf_task_desc_set_int64(task_desc_handle, ALF_TASK_DESC_ACCEL_KERNEL_REF_L, (unsigned long long)kernel_name)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int64 failed, ret=%d\n", ret);
    return 1;
}

dump_print("Before alf_task_desc_set_int64\n");
if ((ret = alf_task_desc_set_int64(task_desc_handle, ALF_TASK_DESC_ACCEL_INPUT_DTL_REF_L, (unsigned long long)input_dtl_name)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int64 failed, ret=%d\n", ret);
    return 1;
}

dump_print("Before alf_task_desc_set_int64\n");
if ((ret = alf_task_desc_set_int64(task_desc_handle, ALF_TASK_DESC_ACCEL_OUTPUT_DTL_REF_L, (unsigned long long)output_dtl_name)) < 0) {
    fprintf(stderr, "Error: alf_task_desc_set_int64 failed, ret=%d\n", ret);
    return 1;
}

dump_print("Before alf_task_create\n");
if ((ret = alf_task_create(task_desc_handle, NULL, 1, 0, 0, &task_handle)) < 0) {
    fprintf(stderr, "Error: alf_task_create failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_desc_destroy\n");
if ((ret = alf_task_desc_destroy(task_desc_handle)) < 0) {
    fprintf(stderr, "Error: alf_exit alf_task_desc_destroy, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_wb_create\n");
if ((ret = alf_wb_create(task_handle, ALF_WB_SINGLE, 1, &wb_handle)) < 0) {
    fprintf(stderr, "Error: alf_wb_create failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_wb_enqueue\n");
if ((ret = alf_wb_enqueue(wb_handle)) < 0) {
    fprintf(stderr, "Error: alf_wb_enqueue failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_finalize\n");
if ((ret = alf_task_finalize(task_handle)) < 0) {
    fprintf(stderr, "Error: alf_task_finalize failed, ret=%d\n", ret);
    return 1;
}

d debug_print("Before alf_task_wait\n");
if ((ret = alf_task_wait(task_handle, -1)) < 0) {
    fprintf(stderr, "Error: alf_task_wait failed, ret=%d\n", ret);
    return 1;
} else if (ret > 0) {
    debug_print("alf_task_wait returned number of work blocks=%d\n", ret);
}

d debug_print("In main: alf_task_wait done.\n");
d debug_print("Before alf_exit\n");
if ((ret = alf_exit(handle, ALF_EXIT_POLICY_FORCE, 0)) < 0) {
    fprintf(stderr, "Error: alf_exit failed, ret=%d\n", ret);
}
return 1;
}

des_gic_print("Execution completed successfully, exiting.\n");
return 0;
}

Example 7-16  SPU source code (~$/halfhello/host/main_spu.c)
#include <stdio.h>
#include <alf_accel.h>
int debug = 1;                  // set to 0 to turn-off debug

int comp_kernel(void *p_task_context, void *p_parm_context,
    void *p_input_buffer, void *p_output_buffer,
    void *p_inout_buffer,
    unsigned int current_count, unsigned int
    total_count)
{
    if (debug)
        printf
            ("Entering alf_accel_comp_kernel, p_task_context=%p, 
            p_parm_context=%p, p_input_buffer=%p, p_output_buffer=%p, 
            p_inout_buffer=%p, current_count=%d, total_count=%d\n", 
            p_task_context, p_parm_context, p_input_buffer,
            p_output_buffer, p_inout_buffer, current_count, total_count);
    printf("Hello World!\n");
    if (debug)
        printf("Exiting alf_accel_comp_kernel\n");
    return 0;
}

int input_prep(void *p_task_context, void *p_parm_context, void *p_dtl,
    unsigned int current_count, unsigned int
    total_count)
{
    if (debug)
        printf
            ("Entering alf_accel_input_list_prepare, p_task_context=%p, 
            p_parm_context=%p, p_dtl=%p, current_count=%d, total_count=%d\n", 
            p_task_context, p_parm_context, p_dtl, current_count,
            total_count);
if (debug)
    printf("Exiting alf_accel_input_list_prepare\n");
return 0;
}

int output_prep(void *p_task_context, void *p_parm_context, void *
    *p_dtl, unsigned int current_count, unsigned int total_count)
{
    if (debug)
        printf
            ("Entering alf_accel_output_list_prepare, p_task_context=%p, p_parm_context=%p, p_dtl=%p, current_count=%d, total_count=%d\n", p_task_context, p_parm_context, p_dtl, current_count, total_count);
    if (debug)
        printf("Exiting alf_accel_output_list_prepare\n");
    return 0;
}

ALF_ACCEL_EXPORT_API_LIST_BEGIN
    ALF_ACCEL_EXPORT_API("", comp_kernel);
    ALF_ACCEL_EXPORT_API("", input_prep);
    ALF_ACCEL_EXPORT_API("", output_prep);
ALF_ACCEL_EXPORT_API_LIST_END

---

**Step 3: Building and deploying the files**

Change to the topmost folder and build the application as shown in Example 7-17.

*Example 7-17  Building and deploying the files*

```bash
cd ~/halfhello
CELL_TOP=/opt/cell/sdk make
```

If everything proceeded as expected, the following binaries are available:

- halfhello/alf_hello_world_hybrid_host64
- halfhello/spu/alf_hello_world_hybrid_spu64.so

Next, deploy the ALF SPU shared library halfhello/spu/alf_hello_world_hybrid_spu64.so to the matching location on the accelerator (QS21 Cell Blade) machine. You can use `scp` as shown in the following example:

```bash
scp ~/halfhello/spu/alf_hello_world_hybrid_spu64.so user@qs21:/tmp
```
Step 4: Running the application
To properly run the application, execute the sequence shown in Example 7-18.

Example 7-18  Running the application

```bash
# Set the environment
export LD_LIBRARY_PATH=/opt/cell/sdk/prototype/usr/lib64
export ALF_LIBRARY_PATH=/tmp
taskset -pc 1 $$
~/halfhello/alf_hello_world_hybrid_host64
```

Before running the application, make sure all Hybrid DaCS daemons are properly running on the host and the accelerator.

7.4 Dynamic Application Virtualization

IBM Dynamic Application Virtualization (DAV) is a technology offering that is available from the IBM alphaWorks Web site at the following address:


DAV implements the function offload programming model. By using DAV, applications that run under Microsoft Windows can transparently tap on the compute power of the Cell/B.E. processor. The originality lies in the fact that the application that we want to accelerate does not require any source code changes. Of course, the offloaded functions must be written to exploit the accelerator, which is the Cell/B.E. processor in this case, but the main application remains unaware. It benefits from an increased level of performance.

The technology was developed initially for the financial services sector but can be used in other areas. The ideas apply wherever an application exhibits computational kernels with a high computational intensity (ratio of computation over data transfers) and cannot be rewritten by using other Cell/B.E. frameworks. The reason it cannot be rewritten is possibly because we do not have its source code or because we do not want to port the whole application to the Cell/B.E. environment.

DAV opens up a whole new world of opportunities to exploit the CBEA for applications that did not initially target the Cell/B.E. processor. It also offers increased flexibility. For example, an application can have its front-end run on a mobile computer or desktop with GUI features and have the hard-core, number crunching part run on specialized hardware such as the Cell/B.E. technology-based blade servers.
From the accelerator perspective, DAV is a way to gain ground into more application areas without needing to port the necessary middleware to run the full application. This is true for every type of acceleration model. The accelerator platform, which is the Cell/B.E. processor, does not need to support many database clients, external file systems, job schedulers, or grid middleware. This is handled at the client level. This separation of work lets the accelerator have a light operating system and middleware layer because it only provides raw computing power.

7.4.1 DAV target applications

In its current implementation, DAV can be used to speed up Microsoft Windows applications written in Visual C/C++ or Visual Basic (Excel spreadsheets). DAV supports accelerator platforms (the server) running Linux. They can be any x86 or ppc64 server, including Cell/B.E. blade servers. DAV refers to the Windows part of the application as the *client side* and the accelerator part as the *server side*.

The best candidate functions for offloading to a Cell/B.E. platform are the ones that show a high ratio of computation over communication. The increased levels of performance that are obtained by running on the Cell/B.E. processor should not be offset by the communication overhead between the client and the server. The transport mechanism that is currently used by DAV is TCP/IP sockets. Other options can be explored in the future to lower the latency and increase the network bandwidth. Clearly, any advance on this front will increase, for a given application, the number of functions that can be accelerated.

Workloads that have a strong affinity for parallelization are optimal. A good example is option pricing by using the Monte Carlo method, as described in Chapter 8, “Case study: Monte Carlo simulation” on page 499.

7.4.2 DAV architecture

The DAV cleverly uses dynamic link libraries (DLLs). In fact, only functions that reside in a DLL can be offloaded to the server. DAV fakes the Microsoft Windows DLL that contains the client code with a new one that communicates with the server to implement the offloaded functions. Enabling an application with DAV means that you must take the following steps:

1. On the client side, identify the functions to be offloaded, including their C prototypes, the data they need on input, and the output they produce. These functions must reside in a DLL.
2. On the server side, write an implementation of the exact same functions, exploiting all the Cell/B.E. strengths. The implementation can use any Cell/B.E. programming techniques. The functions must be made into 32-bit DLLs (the equivalent of shared libraries in Linux.)

3. Back on the client side, fake the application with a DAV DLL, called the stub library, that replaces the original DLL. This DLL satisfies the application but interfaces with the DAV infrastructure to ship data back and forth between the client and the server.

The whole process is shown in Figure 7-9. It shows the unchanged application linked with the stub library on the client side (left) and the implementation of the accelerated library on the server side (right).

![Figure 7-9 DAV process flow](image)

1. Deploy library to remote machine.
2. Install IBM DAV locally and remotely.
3. Use IBM DAV Tooling to generate stub library.
4. Change application linkage from library to stub library.
5. That’s it! No code changes required.
7.4.3 Running a DAV-enabled application

To run the application, we must ensure that the DAV run time on the client side can contact the server. On the server side, we must ensure that the DAV server is running, waiting to be called by the client DAV infrastructure. In this section, we show how to start the DAV service on the DAV server and set the various DAV runtime configuration parameters.

The client application is then run as usual, but it is faster, thanks to the acceleration provided by the Cell/B.E. processor.

7.4.4 System requirements

To install the DAV components, you need Microsoft Windows XP SP2 on the client side and Red Hat Enterprise Linux 5 (RHEL5) or Fedora7 on the Cell/B.E. side. After downloading the DAV software, you receive three files:

- DAVClientInstall.exe
  Run this file to install the client part of the DAV infrastructure. This file must be installed on any machine where a DAV application will run.

- dav-server.ppc64.rpm
  Install this RPM file on every server that is to become a DAV accelerator node.

- DAVToolingInstall.exe
  Run this file to install the DAV Tooling part, the one that creates the stub DLLs from the C prototypes of the functions to be offloaded. This executable file must be installed only on the machine where the stub DLLs are to be created. This file requires the DAV client package to be installed as well.

A C compiler is required to create the DAV stub library. Although a Cygwin environment with gcc might work, we used Microsoft Visual C++® 2005 Express Edition, which is likely to be more common among Microsoft developers. This is a trial version of Microsoft Visual C++ 2005. The installation is described in the DAV user guide available at the following Web address:


The DAV client package comes with a few samples in the C:\Program Files\IBM\DAV\sample directory. We describe the use of DAV following one of the examples.
7.4.5 A Visual C++ application example

In this section, we show the steps to enable a simple Visual C++ application by using DAV. The application initializes two arrays and calls two functions to perform computations on the arrays. Example 7-19 shows the C source code for the main program.

Example 7-19 The main() function

```c
#include "Calculate.h"
int main(int argc, char * argv[]) {  
    #define N 100
    int i;
    double in[N],out[N];

    for(i=0;i<N;i++)
        in[i]=1.*i;

    printf("calculate_Array sent %f\n",calculate_Array(in,out,N));
    printf("calculate_Array2 sent %f\n",calculate_Array2(in,out,N));

    return 0;
}
```

The two functions `calculate_Array` and `calculate_Array2` are shown in Example 7-19 on page 479. They are the ones that we wish to offload to the accelerator. They take the `in` array as input and compute the `out` array and the `ret` result as shown in Example 7-20.

Example 7-20 The computational functions

```c
#include "Calculate.h"
double calculate_Array(double *in,double *out,int size) {
    int i;
    double ret=0.;
    for(i=0;i<size;i++) {
        out[i]=2.*in[i];
        ret+=in[i];
    }
    return ret;
}

double calculate_Array2(double in[],double out[],int size) {
```
int i;
double ret=0.;
for(i=0;i<size;i++) {
    out[i]=0.5*in[i];
    ret+=out[i];
}
return ret;

The function prototypes are defined in a header file. This file is important because this is the input for the whole DAV process. In real situations, the C source code for the main program, the functions, and the header might not be available. You should find a way to create a prototype for each function that you want to offload. This is the only source file that is absolutely required to get DAV to do what it needs to do. You might have to ask the original writer of the functions or do reverse engineering to determine the parameters that are passed to the function.

Example 7-21 shows our header file for this example.

Example 7-21  The Calculate.h header file.

// Calculate.h

#if defined(__cplusplus)
extern "C" {
#endif

double calculate_Array(double *in,double *out,int size);
double calculate_Array2(double in[],double out[],int size);

#if defined(__cplusplus)
}
#endif

The following steps are required to enable DAV acceleration for this application:
1. Build the original application, which creates an executable file and a DLL file for the functions.
2. Run the DAV Tooling component by using the header file for the functions. This creates the stub DLL that replaces the original DLL.
3. Instruct the original application to use the stub DLL.
4. On the Cell/B.E. processor, compile the functions and put them in a shared library.
5. Start the DAV server on the Cell/B.E. processor.
6. Change the DAV parameters on the client machine, so that it points to the right accelerator node.
7. Run the application with DAV acceleration.

**Building the original application**

Figure 7-10 shows how the original application is built. The `.c` and `.h` files are the source code of the compute functions.
We used Visual C++ 2005 Express Edition for this. Figure 7-11 shows the two projects: CalculateApp, which is the main program, and Calculate, which are the functions.

```c
// Calcul.cpp : Defines the entry point for the application.

#include "stdafx.h"

#include "Calculate.h"

#if defined(_cplusplus)
extern "C"
#endif

int main(int argc, _TCHAR* argv[])
{
    #define N 100
    int i;
    double in[N],out[N];

    for(i=0; i<N; i++)
        in[i]=1.*i;

    printf("calculate_Array returned %f\n", calc
    printf("calculate_Array2 returned %f\n", cal
```
Applying the DAV Tooling component

Next, we apply the DAV Tooling, which produces the stub DLLs and the source code for the server-side data marshalling, as shown in Figure 7-12. This step only requires the .h file.

Figure 7-12   The DAV Tooling step
The DAV Tooling component comes bundled with Eclipse 3.3. Your header file should be ready before starting the tooling. After starting DAV Tooling, follow these steps:

1. Open a new project as shown Figure 7-13 and choose the IBM DAV Tooling wizard.

*Figure 7-13   Opening a new project*
2. In the Header File Location window (Figure 7-14), complete the following steps:
   a. Choose a name for the project and open the header file. We use the one supplied in the IBM DAV client package in `C:\Program Files\IBM\DAV\sample\Library\Library.h`.
   b. Click the **Check Syntax** button to check the syntax of the header file. The Finish button does not be active until you check the syntax.
   c. Click the **Finish** button.

![Figure 7-14  Choosing the header file](image-url)
3. In the DAV Tooling window (Figure 7-15), for each function in the list, double-click the function prototype. Complete the semantic information, describing for each argument, the type of data behind it. This information is stored in DAV to generate the necessary code to send data between the client and the server.

![Figure 7-15 Creating the semantic information for the functions](image)

4. After every function is completely described, click the **Generate Stub Code** button (Figure 7-16) to generate the stub DLL.

![Figure 7-16 Generate Stub Code button to create the stub DLL](image)

We experienced a slight problem with the Visual C++ 2005 Express Edition here. The free version lacks libraries that the linker tries to bring in when creating the stub DLL. These libraries (odbccp32.lib and odbc32.lib) are not needed. Therefore, we changed the DAV script C:\Program Files\IBM\IBM DAV Tooling\eclipse\plugins\com.ibm.oai.appweb.tooling_1.0.1\SDK\compilers\vc, so that it would not try to link them.
This step creates many important files for the client and the server side. The files are in the C:Documents\settings\username\workspace\projectname directory. The following files are in that directory:

- **Client directory**
  - `<libraryname>_oai.dll`
  - `<libraryname>_oai.lib`
  - `<libraryname>_stub.dll`
  - `<libraryname>_stub.lib`

- **Server directory**
  - `<libraryname>_oai.cpp`
  - `<libraryname>_oai.h`
  - `makefile`
  - `changes_to_server_config_file`

The client files are needed to run the DAV-enabled application. They must be available in the search path for shared libraries. The _stub tagged files contain the fake functions. The _oai tagged files contain the client side code for the input and output data marshalling between the client and the server. We copied the four files to the C:\Program Files\IBM\DAV\bin directory. We point the executable file to this path later, so that it can find the DLLs when it needs to load them.

The server files (tagged _oai) contain the server side code for the input and output data marshalling between the client and the server. These files must be copied over and compiled on the server node. The makefile is provided to do so. The changes_to_server_config_file file contains the instructions to make the changes to the DAV configuration files on the server to serve our now offloaded functions.
Building the server side shared libraries

Next, we must implement the offloaded functions on the Cell/B.E. server. In our case, we compiled them by using the gcc compiler. In an actual case, we implement the functions by using the compute power of the SPE. The process is shown in Figure 7-17.

As on the client side, we build the lib<libraryname>_oai.so library that contains the data marshalling library. We also build the lib<libraryname>.so library that contains the actual computational functions. The lib<libraryname>.so library is
built by using the Calculate.cpp and Calculate.h files listed in Example 7-20 on page 479 and Example 7-21 on page 480 using a command similar to the following example:

```
$ gcc -shared -fPIC -o libLibrary.so Calculate.cpp
```

We then have two libraries called `libLibrary.so` and `libLibrary_oai.so` that we decided to put in `/usr/dav/services/Library`. This path can be changed to another location provided that you also change the server config file accordingly.

### Configuring the server

To configure the server to receive requests from the client, we begin by setting a system wide IBM_DAV_PATH environment variable to point to the location where DAV has been installed. The default is `/usr/dav`. We added a `dav.sh` file under `/etc/profile.d`.

Next, we change the server DAV config file in `$IBM_DAV_PATH/IBM_DAV.conf`. We incorporate the changes that were suggested in the `changes_to_server_config_file` file when we did the tooling step. These changes provide the name by which the service will be called and path information regarding where to find the shared libraries for the application and the data marshalling libraries. We can also adjust the logging settings and the port number to which the DAV server will be listening. The exact same port number must be specified in the client config file. Example 7-22 shows the contents of the server file.

**Example 7-22  Contents of the IBM_DAV.conf file**

```
#Logging level - 2 = normal
dav.log.level=2

#Log file location - change as desired
#ensure that the location specified has write access for the user running dav
dav.log.directory=/var/log
dav.log.filename=ibm_dav.log

dav.listen.port=12456
dav.listen.max=20

#Services
dav.server.service.restarts=1
dav.server.service.restart.interval=3

dav.server.service.root=services
```
# Relative path specified resulting in path of "$IBM_DAV_PATH/services"
# If this entry is missing it defaults to "$IBM_DAV_PATH/bin"

# Service Entries
# These entries will be automatically generated by IBM DAV Tooling in a
# "changes_to_server_config_file.txt" file.
# The entries should be inserted here

# This is the sample library and its library path resulting in a
# location of "$IBM_DAV_PATH/services/Library"
dav.server.service.Library=Library get_Library_oai Library_oai
dav.server.service.Library.root=Library

Finally, we start the DAV server (Example 7-23).

Example 7-23   Starting the DAV server

    # $IBM_DAV_PATH/bin/davStart -t Library

    Library is the name of the service in this example. The Cell/B.E. server is ready
to receive requests from the client application. We now return to the client side.

**Linking the client application with the stub library**
Before running the accelerated application, we must relink the application with
the stub DLL. This is done by using Visual C++ 2005, changing the linker
parameters in the project properties dialog, and putting the C:\Program
Files\IBM\DAV\bin directory at the top of the list of searched paths.

**Setting the client side parameters**
We must tell the DAV client side where to get its acceleration from. We do this by
editing the IBM_DAV.conf file on the client. The file is located under C:\Program
Files\IBM\DAV. Example 7-24 shows the contents of this file. Our Cell/B.E. blade
address is listed there together with the port number where the DAV server is
listening to.

Example 7-24   The client IBM_DAV.conf file

    # Logging level - 2 = normal
dav.log.level=2

    # Log file location - change as desired
dav.log.directory=c:\
dav.log.filename=ibm_dav.log
Connections to servers
#If sample1 is not available, sample2 will be tried
dav.servers=qdc221
qdc221.dav.server.ip=9.3.84.221
qdc221.dav.server.port=12456

The server is ready. The application has been instructed to load its computational DLL from the path where the stud DLL has been stored. The DAV client knows which accelerator it can work with. We are ready to run the application by using Cell/B.E. acceleration.

Running the application
We now run the application, which, although we have not changed it at all, will benefit from Cell/B.E. acceleration. Running the application entails the following process flow, which is also illustrated in Figure 7-18 on page 492:

1. The application calls a function that now lives in the stub library.
2. Control is passed to the data marshalling library that sends input data to the Cell/B.E. server.
3. The data is shipped to the Cell/B.E. server.
4. The data is received by the data marshalling library on the server side, which in turn, calls the compute functions.
5. The results are handed back to the data marshalling library.
6. The output data is transferred back to the client side.
7. The data movement library handles the output data received.
8. The control is passed back to the stub compute library, which resumes the main application.
Figure 7-18  Flow of the Cell/B.E. accelerated application

Figure 7-19 on page 493 shows the application starting.
After starting, the application immediately calls the `CalculateArray` function, which comes from the stub library. The stub library calls the server side. The system on which we ran the application has the Zone Alarms Checkpoint Integrity Flex firewall installed. This software traps all socket connections for security purposes. This is a simple way to see that our application is now going to the network for its computational functions as shown in Figure 7-20.
Although you do not want this to happen all the time, this software shows that we are going to the network. When we set the firewall to allow the connection to 9.3.84.221:12456, our Cell/B.E. server listening on port 12456, the server side of DAV loads the library that contains our computation functions, gets them to do their work, and returns the output data.

Upon returning from the computational routines, the application continues its work as shown in Figure 7-21.

![Command Prompt - CalculateApp.exe](image1)

**Figure 7-21** Results returned from the DAV server

If we look at the server side, we see that transactions are logged to the DAV log file as shown in Figure 7-22. We are calling from 9.41.223.243.

![root@qdc221 ~]# tail -f /var/log/ibm_dav.log](image2)

**Figure 7-22** All transactions are logged
Looking at the processes running on the server, we see that two processes are running on the host, the davStart daemon and the davService process forked to handle our request as shown in Figure 7-23.

![Figure 7-23 The DAV processes running on the server](image)

We also see from the maps file that our shared libraries have been loaded into the DAV server process as shown in Figure 7-24. See the seventh to tenth lines of output.

![Figure 7-24 The maps file for the davService process](image)

In essence, this is how IBM DAV works. Every application, provided that it gets its computational functions from a DLL, can be accelerated on a Cell/B.E. system by using DAV.
7.4.6 Visual Basic example: An Excel 2007 spreadsheet

IBM DAV is a helpful tool for bringing Cell/B.E. acceleration to Microsoft Windows applications that run on Intel processors. It requires no source code changes to the application and provides the quickest path to the Cell/B.E. system because it limits the porting effort to the number crunching functions only. Still, some usage considerations are important.

IBM DAV is best used for accelerating highly computational functions that require little input and output. This is a general statement that is exacerbated by the fact that the data transfer is currently performed with TCP/IP, which has high latency. Keep in mind that no state data can be kept on the accelerator side between two invocations. Also, all the data needed by the accelerated functions must be passed as arguments for the DAV run time to ship all the required information to the accelerator. The functions should be self-contained.

The davServer process that runs on the Cell/B.E. system is a 32-bit Linux program. Therefore, we are limited to 32-bit shared libraries on the server side.
Application re-engineering

In this part, we focus on specific application re-engineering topics as explained in the following chapters:

- Chapter 8, “Case study: Monte Carlo simulation” on page 499
- Chapter 9, “Case study: Implementing a Fast Fourier Transform algorithm” on page 521
Case study: Monte Carlo simulation

Monte Carlo simulation is a popular computational technique that is used in the Financial Services Sector and in other engineering and scientific areas, such as computational physics. In the Financial Services Sector, for example, this technique is used to calculate stock prices, interest rates, exchange rates, commodity prices, and risk management that requires estimating losses with certain probability over a time period.

The Monte Carlo simulation technique is also used to calculate option pricing (option value). An option is an agreement between a buyer and a seller. For example, the buyer of a European call option buys the right to buy a financial instrument for a preset price (strike price) at expiration (maturity). Similarly, the buyer of a European put option buys the right to sell a financial instrument for a preset price at expiration. The buyer of an option is not obligated to exercise the option. For example, if the market price of the underlying asset is below the strike price on the expiration date, then the buyer of a call option can decide not to exercise that option.

In this chapter, we show how to implement Monte Carlo simulation on the Cell Broadband Engine (Cell/B.E.) system to calculate option pricing based on the Black-Scholes model by providing sample codes. We include techniques to improve the performance and provide performance data. Also, since such mathematical functions as log, exp, sine and cosine are used extensively in
option pricing, we discuss the use of the following IBM Software Developer Kit (SDK) 3.0 libraries by providing examples and makefile:

- **SIMD Math**
  
  This library consists of single-instruction, multiple-data (SIMD) versions (short vector versions) of the traditional libm math functions on the Cell/B.E. server.

- **Mathematical Acceleration Subsystem (MASS)**
  
  This library provides both SIMD and vector versions of mathematical intrinsic functions, which are tuned for optimum performance on the Cell/B.E. server. MASS treats exceptional values differently and can produce slightly different results, compared to SIMD math.

### 8.1 Monte Carlo simulation for option pricing

Option pricing involves the calculation of option payoff values that depend on the price of the underlying asset. The Black-Scholes model is based on the assumption that the price of an asset follows the geometric Brownian Motion, which is described by a Stochastic Differential Equation (SDE). An Euler scheme is used to discretize the SDE. The resulting equation can be solved by using Monte Carlo simulation [12 on page 624 or 23 on page 625].

Example 8-1 shows the basic steps for the Monte Carlo simulation to calculate the price \( S \) of an asset (stock) at different time steps \( 0 = t_0 < t_1 < t_2 < \ldots t_M = T \), where \( T \) is time to expiration (maturity). The current stock price at time \( t_0, S(t_0) = S_0 \), the interest rate \( (r) \), and the volatility \( (v) \) are known. In this example, \( N \) is the number of cycles and \( M \) is the number of time points in each cycle.

**Example 8-1   Pseudo code for Monte Carlo cycles**

```plaintext
for i=0, 1, 2, .., N-1
{
    for j=0, 1, 2, .. M-1
    {
        get a standard normal random number \( X_{ij} \)
        \( dt_j = t_j - t_{j-1} \)
        \( S^i(t_j) = S^i(t_{j-1}) \times \exp((r-0.5v^2) \times dt_j + v \times \sqrt{dt_j} \times X_{ij}) \)
    }
}
```
Example 8-2 shows pseudo code for calculating European option call and put values. In this example, \( S \) is the spot price, and \( K \) is the strike price.

**Example 8-2  European option pricing**

\[
\begin{align*}
C_i &= \text{MAX}(0,S_{[M-1]} - K); \quad i=0,1, ..., N-1 \\
P_i &= \text{MAX}(K-S_{[M-1]},0); \quad i=0,1, ..., N-1
\end{align*}
\]

Average current call value = \( \exp(-rT)*(C_0+C_1+...+C_{N-1})/N \)

Average current put value = \( \exp(-rT)*(P_0+P_1+...+P_{N-1})/N \)

Example 8-3 shows pseudo code to calculate Asian option call and put values. Again, \( S \) is the spot price, and \( K \) is the strike price.

**Example 8-3  Asian option pricing**

\[
\begin{align*}
B_i &= \left( S_i(t_0) + S_i(t_1) + ... + S_i(t_{M-1}) \right)/M; \quad i=0,1,2, ..., N-1 \\
C_i &= \text{MAX}(0,B_i-K); \quad i=0,1,2, ..., N-1 \\
P_i &= \text{MAX}(B_i-K, 0); \quad i=0,1,2, ..., N-1
\end{align*}
\]

Average current call value = \( \exp(-rT)*(C_0+C_1+...+C_{N-1})/N \)

Average current put value = \( \exp(-rT)*(P_0+P_1+...+P_{N-1})/N \)

Different types of options are traded in the market. For example, an American call or put option gives the buyer the right to sell or buy the underlying asset at strike price on or before the expiration date [12 on page 624].

The main computational steps for option values are in Example 8-1 on page 500. In addition, the most time consuming part is getting the standard Gaussian (normal) random variables. These random numbers have the following probability density function with mean 0 and standard deviation 1:

\[
f(x) = \frac{1}{2\pi} \exp\left(-\frac{1}{2}x^2\right), \quad -\infty < x < \infty
\]
8.2 Methods to generate Gaussian (normal) random variables

In this section, we discuss some of the available algorithms to generate Gaussian random numbers. The general procedure entails the following steps:

1. Generate a 32-bit random unsigned integer \( x \).
2. Convert \( x \) to a float or double uniform random variable \( y \) in (0,1).
3. Transform \( y \) to a standard normal random variable \( z \).

Mersenne Twister [11 on page 624] is a popular method to generate random numbers that are 32-bit unsigned integers. This method has a long period, good distribution property, and efficient use of memory.

Other methods are equally as good as the Mersenne Twister method. SDK 3.0 provides the Mersenne Twister method as one of the random number generators [21 on page 624]. This method takes an unsigned integer as a seed and generates a sequence of random numbers. Then it changes these unsigned 32-bit integers to uniform random variables in (0, 1). Finally it transforms the uniform random numbers to standard normal random numbers for which the Box-Muller method or its variant polar method can be used. The Box-Muller method and the polar method require two uniform random numbers and return two normal random numbers [22 on page 625].

Example 8-4 shows sample code to generate two standard normal single-precision random numbers. The code can be easily changed to generate two standard normal double-precision random numbers.

Example 8-4  Code to generate standard normal random numbers

```c
//generate uniform random numbers
float rand_unif()
{
    float c1= 0.5f, c2 = 0.2328306e-9f;
    return (c1 + (signed) rand_MT() * c2);
}

//Box-Muller method
void box_muller_normal(float *z)
{
    float pi=3.14159f;
    float t1,t2;
```
\begin{verbatim}
  t1 = sqrtf(-2.0f * logf(rand_unif()));
  t2 = 2.0f*pi*rand_unif();
  z[0] = t1 * cos(t2);
  z[1] = t1 * sin(t2);
}

// polar method
void pollar_normal(float *z)
{
  float t1, t2, y;

  do {
    t1 = 2.0f * rand_unif() - 1.0f;
    t2 = 2.0f * rand_unif() - 1.0f;
    y = t1 * t1 + t2 * t2;
  } while ( y >= 1.0f );

  y = sqrtf((-2.0f * logf(y)) / y);
  z[0] = t1 * y;
  z[1] = t2 * y;
}
\end{verbatim}

8.3 Parallel and vector implementation of the Monte Carlo algorithm on the Cell/B.E. architecture

In this section, we show how to parallelize and vectorize the Monte Carlo simulation algorithm for option pricing on Cell. In addition, we discuss the use of the SIMD Math, MASS SIMD, and MASS Vector libraries.
8.3.1 Parallelizing the simulation

Figure 8-1 shows the steps for parallelizing the Monte Carlo simulation.

Each of the following steps correspond to those shown in Figure 8-1:

1. Input data
   Read input values such as the number of Monte Carlo simulations, spot_price, strike_price, interest_rate, volatility, time_to_maturity, and num_of_time_steps.

2. Work partitioning
   Decide how many SPEs to use based on the number of simulations that are required. Partition the work and prepare the data needed for each SPE.

3. Set up SPE threads
   Create SPE contexts and execution threads, loads the SPE program, and start the execution of SPE threads by using libspe2 functions.

4. Get data
   Each SPE gets initial seed values and data for Monte Carlo simulation by using direct memory access (DMA).
5. Gaussian random numbers
   Each SPE generates unsigned integer random numbers and converts them to
   standard Gaussian random numbers by using Box-Muller transformation.

6. Monte Carlo cycles
   Each SPE performs its portion of Monte Carlo cycles and computes an
   average option value as result.

7. Send the result
   Each SPE sends its result to the Power Processor Element (PPE) by using
   DMA.

8. Collect results
   Check if the work is done by the SPEs and collect the results from the SPEs.

9. Final value
   Calculate the average of the results that are collected and compute the final
   option value.

Some of the steps are explained further in the discussion that follows. Because
the computational steps for getting European option call and put values and
Asian option values are similar, we restrict our discussion to European option call
value.

In general, the number of Monte Carlo simulations, \( N \), is large (hundreds of
thousands or millions) since the rate of convergence for the Monte Carlo
simulation method is \( 1/\sqrt{N} \). Further, the Monte Carlo cycles (simulations) are
independent. Therefore, we can divide the number of cycles, \( N \) in Example 8-1 on
page 500, by the number of available SPEs and distribute the workload among
the SPEs in step 2 on page 504 (Figure 8-1 on page 504). Further, in each
Synergistic Processor Element (SPE), four Monte Carlo cycles can be done
simultaneously for single precision (two cycles for double precision) by using
Cell/B.E. SIMD instructions. Therefore, the number of Monte Carlo cycles that
are allocated for each SPE must be a multiple of four for single precision and two
for double precision.

As noted previously, in Example 8-1 on page 500, the main computational part
generates standard normal random numbers. Therefore, it requires careful
implementation to reduce the overall computing time. For computational
efficiency and because of limited local storage available on SPEs, avoid
precomputing all random numbers and storing them. Instead, generate the
random numbers during the Monte Carlo cycles in each SPE.

However, this poses a major challenge in the sense that we cannot simply
implement the serial random number generators, such as Mersenne Twister, that
generates a sequence of random numbers based on a single seed value as
input. At a minimum, generating random numbers on SPEs in parallel requires different seed values as input.

Using different seeds on different SPEs is not enough since the generated random numbers can be correlated. That is, the random numbers are not independent. Therefore, the quality of Monte Carlo simulations is not good, which leads to inaccurate results. These remarks apply to all parallel machines, not specific to the Cell/B.E. system.

One way to avoid these problems is to use parallel random number generators such as Dynamic Creator [10 on page 624]. Dynamic Creator is based on the Mersenne Twister algorithm, which depends on a set of parameters, called Mersenne Twister parameters, to generate a sequence of random numbers for a given seed. With the Dynamic Creator algorithm, we can be certain that the Mersenne Twister parameters are different on different SPEs so that the generated sequences are independent of each other, resulting in high quality random numbers overall. Also, Dynamic Creator provides the capability to precompute these parameters, which can be done on a PPE and saved in an array, but only once for a given period.

Following the logical steps given in step 2 on page 504 of Figure 8-1 on page 504, on PPE, we store the values, such as the number of simulations \((J=N/\text{number}\_\text{of}\_\text{SPUs})\), interest rate \((r)\), volatility \((v)\), number of time steps, Mersenne Twister parameters, and the initial seeds in the control structure, defined in Example 8-5 on page 506, to transfer the data to SPEs. Based on these input values, in step 6 on page 505 (Figure 8-1), the average call value is calculated as follows:

\[
C_k = \frac{(C^0 + C^1 + \ldots + C^{J-1})}{J}; \quad k = 1, 2, \ldots \text{number}\_\text{of}\_\text{SPUs},
\]

Here \(C^i\) is defined as shown in Example 8-2 on page 501. As indicated in step 9 on page 505 (Figure 8-1), these results are combined to compute the present call value:

\[
\exp(-rT)*((C_1 + C_2 + \ldots + C_k)/\text{number}\_\text{of}\_\text{SPUs})
\]

In Example 8-5, we define the control structure that will be used to share data between PPU and SPUs.

**Example 8-5  Control structure to share data between the PPU and SPUs**

```c
typedef struct _control_st {
    unsigned int seedvs[4]; /* array of seeds */
    unsigned int dcvala[4]; /* MT parameters */
    unsigned int dcvalb[4]; /* MT parameters */
    unsigned int dcvalc[4]; /* MT parameters */
    int num_simulations; /* number of MC simulations */
} _control_st;
```
In Example 8-6, we provide a sample code for the main program on SPUs to calculate the European option call value.

**Example 8-6  SPU main program for the Monte Carlo simulation**

```c
#include <spu_mfcio.h>

/* control structure */
control_st cb __attribute__ ((aligned (128)));

int main(unsigned long long speid, unsigned long long parm)
{
    /*
     * DMA control structure cb into local store.
     */
    spu_writech(MFC_WrTagMask, 1 << 0);
    spu_mfcdma32((void *)&cb, (unsigned int)parm,
                  sizeof(cb), 0, MFC_GET_CMD);
    (void)spu_mfcstat(2);

    //Get input values for Monte Carlo simulation.
    my_num_simulations = cb.num_simulations;
    s = cb.spot_price;
    x = cb.strike_price;
    r = cb.interest_rate;
    sigma = cb.volatility;
    T = cb.time_to_maturity;
    nt = cb.num_time_steps;

    //get seed
```
seed = ((vector unsigned int){cb.seedvs[0], cb.seedvs[1],
cb.seedvs[2], cb.seedvs[3]});

//
//Get Mersenne Twister parameters that are different on SPUs
A     = ((vector unsigned int){cb.dcvala[0], cb.dcvala[1],
cb.dcvala[2], cb.dcvala[3]});
maskB = ((vector unsigned int){cb.dcvalb[0], cb.dcvalb[1],
cb.dcvalb[2], cb.dcvalb[3]});
maskC = ((vector unsigned int){cb.dcvalc[0], cb.dcvalc[1],
cb.dcvalc[2], cb.dcvalc[3]});

//Initialize the random number generator
rand_dc_set(seed, A, maskB, maskC);

// compute European option average call value -- Monte Carlo simulation
monte_carlo_eur_option_call(s, x, r, sigma, T, nt,
    ,my_sim_size, &value);

// send the value to PPU
spu_writetch(MFC_WrTagMask, 1 << 0);
    spu_mfcdma32((void *)(&value),
        (unsigned int)(cb.valp), sizeof(float),0,MFC_PUT_CMD);

// wait for the DMA to complete
    (void)spu_mfcstat(2);

    return 0;
}

Attention: In control_st, padding, pad[28], is done so that its size is 128 bytes. Otherwise, the DMA commands in the previous program return a bus error message because the minimum length for a DMA is 128 bytes.

8.3.2 Sample code for a European option on the SPU

In this section, we provide sample code to compute an European option call value on SPUs. Since the Monte Carlo cycles are independent, four cycles can be done simultaneously by vectorizing the outer loop in Example 8-1 on page 500. Example 8-7 uses the SPU vector intrinsics and SIMD Math functions.
Example 8-7  SPU vector code for European option call value

```c
#include <spu_intrinsics.h>
#include <simdmath/expf4.h>
#include <simdmath/sqrtf4.h>
//
#include <sum_across_float4.h>
//
/*
s    spot price
x    strike (exercise) price,
r    interest rate
sigma volatility
t_m   time to maturity
nt    number of time steps
*/
void monte_carlo_eur_option_call(float s, float x, float r,
float sigma, float t_m, int nt, int nsim, float *avg)
{
    vector float c, v0, q, zv;
    vector float dtv, rv, vs, p, y, rvt, sigmav, xv;
    vector float sum, u, sv, sinitv, sqdtv;
    int i, j, tot_sim;
    //
    v0    = spu_splats(0.0f);
    c     = spu_splats(-0.5f);
    dtv   = spu_splats((t_m/(float)nt));
    sqdtv = _sqrtf4(dtv);
    sigmav = spu_splats(sigma);
    sinitv = spu_splats(s);
    xv     = spu_splats(x);
    rv     = spu_splats(r);
    vs     = spu_mul(sigmav, sigmav);
    p      = spu_mul(sigmav, sqdtv);
    y      = spu_madd(vs, c, rv);
    rvt    = spu_mul(y, dtv);

    tot_sim = ( (nsim+3)&~3 ) >> 2;
    sum    = spu_splats(0.0f);

    for (i=0; i < tot_sim; i++)
    {
        sv    = sinitv;
        for (j=0; j < nt; j++)
        {
```
rand_normal(zv);
    u = spu_madd(p, zv, rvt);
    sv = spu_mul(sv, _expf4(u));
}
    q = spu_sub(sv, xv);
    sv = _fmaxf4(q, v0);

    sum = spu_add(sum, sv);

}    

*avg = _sum_across_float4(sum) / ((float)tot_sim*4.0f);

In this example, in the step that computes tot_sim, we first make the number of simulations a multiple of four before dividing it by four. In addition to using SIMDmath functions expf4 and sqrtf4, we used the function _fmaxf4 to get the component-wise maximum of two vectors and SDK library function _sum_across_float4 to compute the sum of the vector components.

8.4 Generating Gaussian random numbers on SPUs

In this section, we discuss some techniques to generate Gaussian (normal) random numbers on SPUs. The main parts in Example 8-8 on page 511 are the routine rand_normal for generating standard Gaussian random numbers and the rest of the instructions for option value calculations.

As noted in 8.2, “Methods to generate Gaussian (normal) random variables” on page 502, obtaining standard Gaussian random numbers computation:

1. Generating random unsigned integer numbers.
2. Computing standard normal random numbers, for example, using Box-Muller method or Polar method.

For step 1, we recommend using a parallel random number generator such as Dynamic Creator [10 on page 624]. The library [22 on page 625] consists of functions to generate different random numbers, but it does not include Dynamic Creator. The code in Dynamic Creator for generating random numbers requires only integer computations, which can be easily changed to vector instructions using SPU intrinsics.
For the vector version of Dynamic Creator, the components of a seed vector should be independent. Therefore, on 16 SPUs, 64 independent seed values, unsigned integers, are required. For example, one can use thread IDs as seed values.

For Dynamic Creator, as indicated in Example 8-7, only three Mersenne Twister parameters (A, maskB, and maskC) that are different on SPUs need to be set when the random number generator is initialized. The rest of the Mersenne Twister parameters do not change and can be inlined in the random number generator code.

For step 2, we show the sample SPU code in Example 8-8, which is a vector version of the Box-Muller method in Example 8-4 on page 502. In this example, we convert the vectors of random unsigned integers to vectors of floats, generate uniform random numbers, and use Box-Muller transformation to obtain vectors of standard normal random numbers.

Example 8-8  Single precision SPU code to generate Gaussian random numbers

```c
#include <spu_mfcio.h>
#include <simdmath/sqrtf4.h>
#include <simdmath/cosf4.h>
#include <simdmath/sinf4.h>
#include <simdmath/logf4.h>

void rand_normal_sp(vector float *z)
{
    vector float u1,u2,v1,v2,w1,p1;
    vector float c1 = ((vector float) { 0.5f,0.5f,0.5f,0.5f});
    vector float c2 = ((vector float) { 0.2328306e-9f, 0.2328306e-9f,
                                      0.2328306e-9f, 0.2328306e-9f});
    vector float c3 = ((vector float) { 6.28318530f, 6.28318530f,
                                      6.28318530f, 6.28318530f});
    vector float c4 = ((vector float) { -2.0f,-2.0f,-2.0f,-2.0f});
    vector unsigned int y1, y2;

    // get y1, y2 from random number generator.

    // convert to uniform random numbers
    v1 = spu_convtf( (vector signed int) y1, 0);
    v2 = spu_convtf( (vector signed int) y2, 0);
    u1 = spu_madd( v1, c2, c1);
    u2 = spu_madd( v2, c2, c1);
} 
```
// Box-Muller transformation
   w1 = _sqrtf4( spu_mul(c4, _logf4(u1)) );
   p1 = spu_mul(c3, u2);
   z[0] = spu_mul(w1, _cosf4(p1) );
   z[1] = spu_mul(w1, _sinf4(p1) );

In Example 8-8, the C statement (vector signed int) y1 converts the
components of y1 that are unsigned integers to signed integers. The instruction
spu_convtf converts each component of the resulting vector signed int to a
floating-point value and divides it by 2^{scale}, where scale=0.

For double precision, the generated 32-bit unsigned integer random numbers are
converted to floating-point values and extended to double-precision values. A
double-precision vector has only two elements because each element is 64-bits
long. In Example 8-8, the vector y1 has four elements that are 32-bit unsigned
integer random numbers. Therefore, we can compute the required two
double-precision vectors of uniform random numbers for Box-Muller
transformation by shuffling the components of y1. Doing this avoids the
computation of y2 in Example 8-8. Example 8-9 explains this idea.

Example 8-9  Double-precision SPU code to generate Gaussian random numbers

```c
#include <spu_mfcio.h>
#include <simdmath/sqrtd2.h>
#include <simdmath/logd2.h>
#include <simdmath/cosd2.h>
#include <simdmath/sind2.h>

//

void rand_normal_dp(vector double *z)
{
   vector double u1,u2,v1,v2,w1,p1;
   vector double c1 = { 0.5, 0.5};
   vector double c2 = { 0.2328306e-9, 0.2328306e-9};
   vector double c3 = { 6.28318530, 6.28318530};
   vector double c4 = { -2.0, -2.0};

   vector unsigned char pattern={4, 5, 6, 7, 0, 1, 2, 3, 12, 13, 14, 15,
8, 9, 10, 11};
   vector unsigned int y1, y2;

   // get y1 from random number generator.
```
\[ y_2 = \text{spu\_shuffle}(y_1, y_1, \text{pattern}); \]

// convert to uniform random numbers
\[ v_1 = \text{spu\_extend}(\text{spu\_convtf}(\text{vector signed int} y_1, 0)); \]
\[ v_2 = \text{spu\_extend}(\text{spu\_convtf}(\text{vector signed int} y_2, 0)); \]
\[ u_1 = \text{spu\_madd}(v_1, c_2, c_1); \]
\[ u_2 = \text{spu\_madd}(v_2, c_2, c_1); \]

// Box-Muller transformation
\[ w_1 = \text{_sqrtd2}(\text{spu\_mul}(c_4, \text{logd2}(u_1))); \]
\[ p_1 = \text{spu\_mul}(c_3, u_2); \]
\[ z[0] = \text{spu\_mul}(w_1, \text{cosd2}(p_1)); \]
\[ z[1] = \text{spu\_mul}(w_1, \text{sind2}(p_1)); \]

Next we discuss ways to tune the code on the SPUs. To improve register utilization and instruction scheduling, the outer loop in Example 8-7 on page 509 can be unrolled. Further, a vector version of the Box-Muller method and Polar method computes two vectors of standard normal random numbers out of two vectors of unsigned integer random numbers. Therefore, we can use all generated vectors of normal random numbers by unrolling the outer loop, for example, to a depth of two or four.

In Example 8-8 on page 511 and Example 8-9 on page 512, we used mathematical intrinsic functions, such as \text{exp}, \text{sqrt}, \text{cos} and \text{sin}, from SIMD math library, which takes advantage of SPU SIMD (vector) instructions and provides significant performance gain over the standard \text{libm} math library [20 on page 624]. MASS, which available in SDK 3.0, provides mathematical intrinsic functions that are tuned for optimum performance on PPU and SPU. The MASS libraries provide better performance than SIMD math libraries for most of the intrinsic functions. In some cases, the results for MASS functions might not be as accurate as the corresponding functions in the SIMD math libraries. In addition, MASS can handle the edges differently.

**More information:** For comparison of the accuracy of results between MASS functions and SIMD math functions, see the Mathematical Acceleration Subsystem publications [21 on page 624].

We found that, with our implementation of European option pricing, performance can be improved by using MASS. Also the accuracy of the computed result of using MASS is about the same as that of using SIMD math. The current version of MASS in SDK 3.0 provides only single-precision math intrinsic functions.
Use of MASS inline functions instead of SIMD math functions is easy. For example, the MASS functions require only changing the SIMD math include statements in Example 8-7 on page 509 and in Example 8-8 on page 511 by the corresponding include statements given in Example 8-10 and in Example 8-11, respectively.

Example 8-10  Include statements to include MASS intrinsic functions on SPUs

```c
#include <mass/expf4.h>
#include <mass/sqrtf4.h>
```

Example 8-11  Include statements to include MASS intrinsic functions on SPUs

```c
#include <mass/sqrtf4.h>
#include <mass/cosf4.h>
#include <mass/sinf4.h>
#include <mass/logf4.h>
```

Alternatively, in Example 8-10 and in Example 8-11, you can use MASS SIMD library functions instead of inlining them. To this, you must change the include statements and the function names as shown in Example 8-12, which is a modified version of Example 8-8 on page 511.

Example 8-12  SPU code to generate Gaussian random numbers using the MASS SIMD library

```c
#include <spu_mfcio.h>
#include <mass_simd.h>

//

void rand_normal_sp(vector float *z)
{
    vector float u1,u2,v1,v2,w1,p1;
    vector float c1 = ((vector float) { 0.5f,0.5f,0.5f,0.5f});
    vector float c2 = ((vector float) { 0.2328306e-9f, 0.2328306e-9f,
                                       0.2328306e-9f, 0.2328306e-9f});
    vector float c3 = ((vector float) { 6.28318530f, 6.28318530f,
                                        6.28318530f, 6.28318530f});
    vector float c4 = ((vector float) { -2.0f,-2.0f,-2.0f,-2.0f});
    vector unsigned int y1, y2;

    // get y1, y2 from random number generator.
```
//convert to uniform random numbers
    v1 = spu_convtf( (vector signed int) y1, 0);
    v2 = spu_convtf( (vector signed int) y2, 0);
    u1 = spu_madd( v1, c2, c1);
    u2 = spu_madd( v2, c2, c1);

// Box-Muller transformation
    w1 = sqrtf4( spu_mul(c4, logf4(u1)) );
    p1 = spu_mul(c3, u2);
    z[0] = spu_mul(w1, cosf4(p1));
    z[1] = spu_mul(w1, sinf4(p1));
}

Further, you must add the libmass_simd.a MASS SIMD library at the link step. Example 8-13 shows the makefile to use the library.

Example 8-13  Makefile to use the MASS SIMD library

CELL_TOP = /opt/cell/sdk/
SDKLIB = /opt/cell/sdk/prototype/sysroot/usr/lib

# Choose xlc over gcc because it gives slightly better performance
SPU_COMPILER = xlc
#

PROGRAMS_spu    := mceuro_spu

INCLUDE = -I/usr/spu/include

OBJS  = mceuro_spu.o

LIBRARY_embed   := mceuro_spu.a

# use default optimization because higher levels do not improve
CC_OPT_LEVEL          := -O3

IMPORTS = /usr/spu/lib/libmass_simd.a

###########################################################################
#                         make.footer                                       
###########################################################################
To create a double-precision version of Example 8-13 on page 515, see the methods used in Example 8-9 on page 512.

Recall that the number of Monte Carlo cycles for European option pricing is typically very large, in the hundreds of thousands or millions. In such cases, the call overhead for math functions can degrade the performance. To avoid call overhead and improve the performance, you can use the MASS Vector library [21 on page 624] to provide the math functions to calculate the results for an array of input values with a single call. To link to MASS vector library, you must replace libmass_simd.a with libmassv.a in IMPORTS in the makefile as shown in Example 8-13 on page 515.

Example 8-14 shows how to restructure the code to use the MASS vector functions.

```
Example 8-14  Code to use MASS vector functions

#include <spu_mfcio.h>
#include <massv.h>

//define buffer length
#define BL 32
#define BL2 64

void rand_normal_sp(vector float *z) {
    vector float c1 = ((vector float) { 0.5f,0.5f,0.5f,0.5f});
    vector float c2 = ((vector float) { 0.2328306e-9f, 0.2328306e-9f,
        0.2328306e-9f, 0.2328306e-9f});
    vector float c3 = ((vector float) { 6.28318530f, 6.28318530f,
        6.28318530f, 6.28318530f});
    vector float c4 = ((vector float) { -2.0f,-2.0f,-2.0f,-2.0f});
    vector unsigned int y[BL2] __attribute__((aligned (16)));
    vector float u1[BL] __attribute__((aligned (16)));
    vector float u2[BL] __attribute__((aligned (16)));
    vector float w[BL] __attribute__((aligned (16)));
    vector float v1,v2;
    int i, size=4*BL;
```
// get random numbers in the array y
// convert to uniform random numbers

for (i=0, j=0; i < BL; i++, j+=2)
{
    v1 = spu_convtf( (vector signed int) y[j], 0);
    v2 = spu_convtf( (vector signed int) y[j+1], 0);
    u1[i] = spu_madd( v1, c2, c1);
    u2[i] = spu_madd( v2, c2, c1);
}

// MASS
vslog ((float *) u1, (float *) u1, &size);

for (i=0; i < BL; i++)
{
    u1[i] = spu_mul(c4,u1[i]);
    u2[i] = spu_mul(c3,u2[i]);
}

vssqrt ( (float *)u1, (float *)u1, &size);
vsincos ( (float *)w, (float *)u2, (float *)u2, &size);

for (i=0, j=0; i < BL; i++, j+=2)
{
    z[j] = spu_mul(u1[i], u2[i]);
    z[j+1] = spu_mul(u1[i], w[i]);
}

In Example 8-14 on page 516, pointers to vector floats, such as u1 and u2, are cast to pointers of floats in the MASS vector function calls since the array arguments in the MASS vector functions are defined as pointers to floats. For details, see the prototypes for the SPU MASS vector functions in /usr/spu/include/massv.h.
8.5 Improving the performance

In this section, we discuss ideas to improve the performance and provide performance data.

**More information:** General guidelines for improving performance of an application on SPUs are provided in “Maximizing the power of the Cell Broadband Engine processor: 25 tips to optimal application performance developerWorks” [13 on page 624].

With our European option pricing code implementation, we found that XLC gives better performance than gcc. Further, besides using MASS, we used the following techniques to improve performance:

- Unrolled the outer loop in Example 8-7 to improve register utilization and instruction scheduling
  
  Note that, since a vector version of the Box-Muller method and Polar method computes two vectors of standard normal random numbers out of two vectors of unsigned integer random numbers, all generated vectors of normal random numbers can be used by unrolling the outer loop, for example, to a depth of eight or four.

- Inlined some of the routines to avoid call overhead

- Avoided branching *(if and else statements)* as much as possible within a loop

- Reduced the number of global variables to help compiler with register optimization

Moreover, we used the -O3 compiler optimization flag with XLC. We also tried higher level optimization flags such as -O5, which did not make a significant performance difference, compared to -O3.

Example 8-15 shows the input values for the Monte Carlo simulation.

*Example 8-15  Input values for Monte Carlo simulation*

```c
num_simulations = 200000000
    spot_price = 100.0
    strike_price = 50.0
    interest_rate = 0.10
    volatility = 0.40;
    time_to_maturity = 1
    num_time_steps = 1
```
Figure 8-2 shows the performance results in terms of millions of simulations per second (M/sec) for the tuned single precision code for European option on QS21 (clock speed 3.0 GHz).

Figure 8-2  Performance of the Monte Carlo simulation on QS21
Case study: Implementing a Fast Fourier Transform algorithm

In this chapter, we describe the code development process and stages for a Fast Fourier Transform (FFT) library that is included as a prototype project in the Software Developer Kit (SDK) 3 for the Cell Broadband Engine (Cell/B.E.) server. FFT algorithms are key in many problem domains including the seismic industry.

We provide a real-world example of the code development process for the Cell/B.E. processor running on an IBM BladeCenter QS21 blade server. The snippets of source code found in this chapter are included for illustration and do not constitute a complete FFT solution.

Specifically, this chapter includes the following topics:

- 9.1, “Motivation for an FFT algorithm” on page 522
- 9.2, “Development process” on page 522
- 9.3, “Development stages” on page 526
- 9.4, “Strategies for using SIMD” on page 529
9.1 Motivation for an FFT algorithm

FFTs are used in many applications that process raw data looking for a signal. There are many FFT algorithms ranging from relatively simple powers-of-two algorithms to powerful, but processor-intensive algorithms that are capable of working on arbitrary inputs. FFT algorithms are well suited to the Cell/B.E. processor because they are floating-point intensive and exhibit regular data access patterns.

The IBM SDK 3.0 for the Cell/B.E. processor contains a prototype FFT library that is written to explicitly exploit the features of the Cell/B.E. processor. This library uses several different implementations of an algorithm to solve a small class of FFT problems. The algorithm is based on a modified Cooley-Tukey type algorithm. All of the implementations use the same basic algorithm, but each implementation does something different to make the algorithm perform the best for a particular range of problem sizes.

The first step in any development process is to start with a good algorithm that maps well to the underlying architecture of the machine. The best compilers and hardware cannot hide the deficiencies that are imposed by a poor algorithm. It is necessary to start with a good algorithm.

When selecting the algorithm, we used the following considerations in addition to others:

- Require support for problem sizes that can be factored into powers of 2, 3, and 5. This eliminates the straight “power-of-two” algorithm or Prime Factor Algorithm (PFA).
- A single problem should fit within the memory of a Synergistic Processor Unit (SPU). This keeps the code simpler by eliminating the need for two or more SPUs to coordinate and work on a single problem.

9.2 Development process

There is no formal set of rules or process for developing or porting an existing application to the Cell/B.E. processor. The program team that wrote the FFT library developed an iterative development process mapped on top of a predefined set of logical stages. The iterative development process was useful during some or all of the development stages.
Figure 9-1 illustrates the stages and process used during implementation of the FFT code for Cell/B.E. processor. We describe this figure in more detail in the sections that follow.

9.2.1 Code

The “code” box in the Process box in Figure 9-1 represents the actual physical writing of code. This can include the implementation of a formal program specification or coding done without any documentation other than code directly from the programmer.

Experienced programmers might require fewer code iterations, while less experienced programmers normally take more iterations. There is no concept of duration other than it is typical to break code sessions into logical functional elements.

Coding can be done in any manner that is convenient to the programmer. The IBM Eclipse integrated development environment (IDE) for the Cell/B.E. processor in the SDK is a good choice for writing and debugging Cell/B.E. code.
Some of the team that implemented the FFT library used in IBM Eclipse IDE for the Cell/B.E. processor while others on the team used the classic Linux-based editors and debuggers or printf for C programmers.

9.2.2 Test

The “test” box inside the Process box in Figure 9-1 on page 523 represents the testing of code that has been compiled. For Cell/B.E. applications, testing is still a necessary and critical step in producing well performing applications.

The test process for the Cell/B.E. application undoubtedly requires testing for code performance. Testing is normally focused on code that runs on the Cell/B.E. SPU. The focus is more on making sure all code paths are processed and the accuracy of the output.

9.2.3 Verify

The “verify” box in the Process box in Figure 9-1 on page 523 represents an important aspect of many Cell/B.E. applications. The need for verification of code that runs on the SPU is of special importance. The Cell/B.E. Synergistic Processor Element (SPE) single-precision floating point is not the same implementation as found in the PowerPC Processor Unit (PPU) and other processors.

Code that is ported from other processor platforms presents a unique opportunity for the verification of Cell/B.E. programs by comparing output from the original application for accuracy. A binary comparison of single-precision floating point is performed to eliminate conversions to and from textual format.

The FFT coding team chose to write a separate verification program to verify output. The verification code uses a well-known open source FFT library that is supported on PowerPC processors. The results from the Cell/B.E. FFT code were compared at a binary level with the results from the open source FFT results. Example 9-2 on page 525 shows how the single-precision floating point was converted to displayable hexadecimal and then converted back to single-precision floating point. These functions are one way to compare SPE and PPE floats.
Example 9-1 shows a simple C function from the test program. It illustrates how two single-precision floating point values that represent a complex number are output in displayable hexadecimal form.

Example 9-1  Single-precision floats as displayable hexadecimal

typedef union {
    unsigned int i;
    float        f;
} Conv_t;

void printOutput( float f1, float f2 ) {
    if ( Output ) {
        Conv_t t1, t2;
        t1.f = f1;
        t2.f = f2;
        printf( "%08x %08x\n", t1.i, t2.i );
    } else {
        printf( "%f %f\n", f1, f2 );
    }
}

Example 9-2 shows a code snippet from the verification program, which reads displayable hexadecimal from stdin and converts it into a complex data type. The verify program reads multiple FFT results from stdin, which explains why variable p is a two-dimensional matrix.

Example 9-2  Reading complex numbers from a displayable hexadecimal format

typedef struct {
    float real;
    float imag;
} Complex;

Complex *t = team[i].srcAddr;
MT_FFTW_Complex *p = fftw[i].srcAddrFC;
unsigned int j;
for ( j=0; j < n; j++ ) {
    volatile union {
        float f;
        unsigned int i;
    } t1, t2;
    scanf( "%x %x\n", &t1.i, &t2.i );
#define INPUT_FMT "i=%d j=%d r=+%13.10f/%8.8X i=+%13.10f/%8.8X\n"
Chapter 9. Case study: Implementing a Fast Fourier Transform algorithm
(verbose ? fprintf(stdout, INPUT_FMT, i, j, t1.f, t1.i, t2.f, t2.i) : 0);
    t[j].real = p[j][0] = (double)t1.f;
    t[j].imag = p[j][1] = (double)t2.f;
} // for j  (number of elements in each fft)

The code shown in Example 9-2 on page 525 accurately reconstitutes a single-precision floating point value. This is only true for data that is generated by the same mathematical, floating-point representations (that is PowerPC to PowerPC). The FFT library verification program compares single-precision floating point values between PowerPC and SPE single-precision format. The format of the two floating point representations are similar and can normally be ignored for verification purposes.

9.3 Development stages

The development stages list in Figure 9-1 on page 523 were arrived at prior to the beginning of the code development and later revised. The purpose of introducing these stages is to provide a way to monitor progress of the project, which seems like a practical way to develop the solution.

In the following sections, we describe the different stages and include sample code to illustrate the evolution of some aspects of the code that are unique to the Cell/B.E. processor.

9.3.1 x86 implementation

The goal or deliverable from this stage is a functional FFT implementation that runs on x86 Linux hardware. Little effort was invested in producing the code that performed optimally on x86. The goal was to prove that the basic algorithm produced correct results and met our requirements for running in an SPU.

This version of the FFT code included an initial functional interface from a test program to the actual FFT code and a primitive method for displaying the results. The verification of FFT output was essential to ensure accuracy.

9.3.2 Port to PowerPC

The x86 FFT implementation was ported to PowerPC hardware by recompiling the x86 source code on an IBM QS20 blade. The effort to do this was almost trivial as can be expected.
The PowerPC version of the code performed much slower than the x86 version of the code. This performance is due to the difference in the relative power of the PPU portion of the Cell/B.E. processor compared to a fairly high-end dual-core x86 CPU on which the x86 code was developed. Even though this code was simple and single threaded, the x86 processor core that was used for development is a much more power processor core than the PPU. Fortunately, for the Cell/B.E. processor, the power of the chip lies in the eight SPUs, which we take advantage of. (The PPU is eventually left to manage the flow of work to the SPUs.)

9.3.3 Single SPU

The first real specific Cell/B.E. coding task was to take the PowerPC (PPU) code and run the compute kernel on a single SPU. This involved restructuring the code by adding calls to the SDK to load an SPU module. More code was added to implement a simple direct memory access (DMA) model for streaming data into the SPU and streaming it back to main store memory.

Example 9-3 shows a code snippet that demonstrates how a single SPU thread is started from the main FFT PPU code. The use of spe_context_run() is a synchronous API and blocks until the SPU program finishes execution. The multiple-SPU version of this code, which uses pThread support, is the preferred solution and is discussed in 9.3.5, “Multiple SPUs” on page 529.

Example 9-3  Running a single SPU

```c
#include <stdio.h>
#include <stdlib.h>
#include <libspe2.h>

spe_context_ptr_t ctx;
unsigned int entry = SPE_DEFAULT_ENTRY;

/* Create context */
if ((ctx = spe_context_create (0, NULL)) == NULL) {
    perror ("Failed creating context");
    exit (1);
}

/* Load program into context */
if (spe_program_load (ctx, &dma_spu)) {
    perror ("Failed loading program");
    exit (1);
}
```
/* Run context */
if (spe_context_run(ctx,&entry,0,buffer,(void *)128,NULL)< 0) {
    perror ("Failed running context");
    exit (1);
}
/* Destroy context */
if (spe_context_destroy (ctx) != 0) {
    perror("Failed destroying context");
    exit (1);
}

Note: The program in Example 9-3 on page 527 can be easily compiled and run as a spulet.

9.3.4 DMA optimization

The initial versions of the code were processor-intensive and slow because they did not use the SIMD features of the SPUs. The code was not well structured or optimized. As a result, the time spent doing DMA transfers was small relative to the time spent computing results. In this stage of the project, DMA optimization was not considered to be important. Early measurements showed that, without double buffering, DMA transfer time was only about two percent of the total time it took to solve a problem.

The time spent in the computation phase of the problem shrank as the code became more optimized by using various techniques, which are described later in this section. As a result, DMA transfer time grew larger relative to the time spent performing the computation phase. Eventually the team turned their attention from focusing on the computation phase to the growing DMA transfer time.

The programming team knew double buffering could be used to hide the time it takes to do DMA transfers if done correctly. At this phase of the project, the team was determined that DMA transfers were about 10 percent of the total time it took to solve a problem. Correctly implemented DMA transfers were added to the code at this phase to overlap with current ongoing computations. Given the optimization effort that had been put into the base algorithm, the ability to reduce our run time per problem by 10 percent was worth the effort in the end.

However, double buffering does not come for free. In this case, the double-buffering algorithm required three buffer areas, one buffer for incoming data and outgoing DMAs and two for the current computation. The use of a single buffer for both input and output DMAs was possible by observing that the data transfer time was significantly smaller than compute time. It was possible to
allocate the third buffer area when working on a small problem, but for the largest problems, memory was already constrained and another buffer area was not possible.

The solution was to have several implementations of the algorithm, where each specific implementation would use a different data transfer strategy:

- For large problems where an extra data buffer was not possible, double buffering would not be attempted.
- For smaller problems where an extra data buffer was possible, one extra data buffer would be allocated and used for background data transfers, both to and from SPU memory.

The library on the PPU that dispatched the problems to the SPUs now had to look at the problem size that was provided by the user and choose between two different implementations. The appropriate implementation was then loaded on an SPU, and problems that were suitable for that implementation were sent to that SPU.

With this solution, the code was flexible enough, so that small problems could take advantage of double buffering and large problems were still able to execute.

### 9.3.5 Multiple SPUs

The original project requirements specified that no single FFT problem would be larger than 10,000 points. This allowed for implementing a serial FFT algorithm where a single SPU could solve the largest sized FFT problem instead of a more complicated parallel algorithm using multiple SPUs in concert. (The SDK provides an example of a large parallel FFT algorithm in the sample codes.)

As a result, extending the code to run on multiple SPUs allowed for increasing the throughput, not the maximum problem size. All of the SPUs that get used will work independently, getting their work from a master work queue maintained in PPU memory.

### 9.4 Strategies for using SIMD

The SPUs are designed from the ground up as vector units. If a coding style does not use these vector registers, 75% of the potential performance is lost.

FFT implementations based on “power-of-two” algorithms are relatively easy to implement on a vector processor because of the regular data access patterns and boundary alignments of the data. A non “power-of-two” algorithm is
significantly more difficult to map to a vector architecture because of the data access patterns and boundary issues.

The FFT prototype library uses two different vectorization strategies:

- Striping multiple problems across a vector
- Synthesizing vectors by loop unrolling

### 9.4.1 Striping multiple problems across a vector

A relatively easy way to use the vector registers of the SPU is to treat each portion of the register independently from other portions of the register. For example, given four different floating point values from four different problems, all of the floating point values can reside in a single register at a time. This allows for computation on four different problems in a register at the same time. The code to do this a simple extension of the original scalar code. However, except now instead of loading one value from one problem into a register, the four values are loaded from four problems into a single vector register. One way to visualize this technique is to consider it as “striping” multiple problems across a single register.

If there are four FFT problems in memory and those problems are of the same length, then this is trivial to implement. All four problems are done in lockstep, and the code that performs the indexing, the twiddle calculations, and other code can be reused for all four problems.

The problem with this approach is that it requires multiple FFT problems to reside in SPU memory. For the smallest problem sizes (up to around 2500 points), this is feasible. However, for larger problem sizes, this technique cannot be leveraged because the memory requirements are too great for a single SPU.

### 9.4.2 Synthesizing vectors by loop unrolling

For the larger problems in the FFT prototype, “striping” multiple problems across a register is not possible because there is not enough space in SPU memory for multiple problems to reside. However, it is still necessary to use the vector features of the SPU for performance reasons.

The following solution resolves this problem:

- Unroll the inner computation loop by a factor of four to give four sets of scalar values to work with.
- Use shuffle operations to synthesize vector registers from the scalar values.
Perform the math using the vector registers.

Split the results in the vector registers apart and put them back into their proper places in memory.

The advantage of this solution is that it allows the use of the vector capabilities of the SPU with only one FFT problem resident in memory. Parallelism is achieved by performing loop unrolling on portions of the code.

A disadvantage of this solution is the need to constantly assemble and disassemble vector registers. Although there is a large net gain, a speedup of four times was not realized by using the vector registers. This is because the time spent assembling and disassembling the vector registers was fairly large compared to the time spent computing with those vector registers. (On a more intensive calculation, this technique would show a greater improvement.)

### 9.4.3 Measuring and tweaking performance

The primary tools for measuring performance of the FFT code are the simulator, the spu-timing tool, and a stopwatch. (An actual stopwatch was not used, but a simple stopwatch was implemented in the code). Since the FFT was developed, additional performance tools have become available, some of which are explained in this book.

The spu-timing tool allowed for analyzing the code generated by the compiler to look for inefficient sections of code. Understanding why the code was inefficient provided insight into where to focus efforts on generating better performing code. The simulator gave insight as to how the code would behave on real hardware. The dynamic profiling capability of the simulator let us verify our assumptions about branching behavior, DMA transfer time, and overall SPU utilization.

The stopwatch technique provided verification that the changes to the code were actually faster than the code that was replaced.

In the following sections, we discuss some of the performance lessons that we learned during this project.

**The SIMD Math library**

An FFT algorithm is an intensive user of the sinf() and cosf() function in the math library. These are expensive functions to call. Therefore, minimizing their use is critical.

An important feature of the SIMD Math library are the special vector versions of sinf() and cosf() that take four angles in a vector as input and return four sine or cosine values in a vector as output. These functions were used to dramatically
reduce the amount of time spent computing sines and cosines. The source code for the functions is available in the SDK, which allows you to adapt them for your specific needs. A merged sinf() and cosf() function was coded that generates eight outputs (four sines and four cosines) from a single vector of input angles. (This merged function was even faster than calling the two vector functions separately.) There are also inline versions of the functions available, which are used to improve branching performance.

**Improving performance with code inlining and branch hints**

In the code, we used two methods to improve performance as explained in the following sections.

**Code inlining**

Code inlining is an effective way to improve branching performance, because the fastest branch is the one that you might not have taken. During the development cycle, code is often written in functions or procedures to keep the code modular. During the performance tuning stage, these functions and procedure calls are reviewed for potential candidates that can be inlined. Besides reducing branching, inlining has a side benefit of giving the compiler more code to work with, allowing it to possibly use better the SPU pipelines.

A certain amount of judgement is required when inlining code. If the code is called from many locations, the compiled size of the code can grow to an unacceptable level. It is also possible to have so much code in a basic block that the compiler starts to spill registers to the stack, which leads to degraded performance.

**Branch hint directives**

Most of the branching in the FFT code is caused by loop structures. The compilers generate the correct branch hints for loop structures, so we do not worry about them. In the rare case where “if-then-else” logic is needed, an analysis of the code is performed to determine which path is the most used, and a branch hint is inserted manually. This improves the performance of the run time by about one percent, which is well worth the effort. Code that has a lot of “if-then-else” logic, where the most chosen path can be predicted, is likely to benefit more from branch hints than this code does.

**Effective use of the shuffle intrinsic**

In 9.4.1, “Striping multiple problems across a vector” on page 530, we discussed a technique where four FFT problems in memory were “striped” across vector registers. This allowed all four problems to be computed in lockstep by using code that was basically an extension of the original scalar code.
When the FFT problems first come into memory, they are in an interleaved format where the real portion of a complex number is followed immediately by the imaginary portion of the complex number. Loading a vector from memory results in two consecutive complex numbers from the same problem in a vector register. That then requires shuffle operations to split the reals from the imaginaries and more shuffle operations to end up with four problems striped across the vector registers.

It is most efficient to do all of the data re-arrangement after the problems are in SPU memory, but before computation begins. Example 9-4 shows a simple data structure that is used to represent the problems in memory and a loop that does the re-arrangement.

**Example 9-4 Rearranging complex numbers**

typedef union {
    Complex_t prob[4][MAX_PROB_SIZE_C2C_4];
    union {
        struct {
            float real[MAX_PROB_SIZE_C2C_4*4];
            float imag[MAX_PROB_SIZE_C2C_4*4];
        } sep;
        struct {
            vector float real[MAX_PROB_SIZE_C2C_4];
            vector float imag[MAX_PROB_SIZE_C2C_4];
        } vec;
    } u;
} Workarea_t __attribute__ ((aligned (128)));

Workarea_t wa[2];
// Separate into arrays of floats and arrays of reals
// Do it naively, one float at a time for now.
short int i;
for (i=0; i < worklist.problemSize; i++) {
    wa[0].u.sep.real[i*4+0] = wa[1].prob[0][i].real;
    wa[0].u.sep.real[i*4+1] = wa[1].prob[1][i].real;
    wa[0].u.sep.real[i*4+2] = wa[1].prob[2][i].real;
    wa[0].u.sep.real[i*4+3] = wa[1].prob[3][i].real;
    wa[0].u.sep.imag[i*4+0] = wa[1].prob[0][i].imag;
    wa[0].u.sep.imag[i*4+1] = wa[1].prob[1][i].imag;
    wa[0].u.sep.imag[i*4+2] = wa[1].prob[2][i].imag;
    wa[0].u.sep.imag[i*4+3] = wa[1].prob[3][i].imag;
}
The data structure provides access to memory in three ways:

- Four problems where the reals and imaginaries are interleaved
- One array of floats representing all of the reals, and one array of floats representing all of the imaginaries
- One vector array of floats representing all of the reals, and one vector array of floats representing all of the imaginaries

These arrays have one fourth as many elements as the previous arrays because they are vector arrays.

On entry to the code wa[1] contains four problems in interleaved format. This code copies all of the interleaved real and imaginary values from wa[1] to wa[0], where they appear as separate arrays of reals and imaginaries. At the end of the code, the view in wa[0] is of four problems striped across vectors in memory.

While this code works, it is hardly optimal:

- Only four bytes are moved at a time.
- It is scalar code. Therefore, the compiler must shuffle the floats into the preferred slots of the vector registers, even though the values are going to be written back to memory immediately.
- When writing each value to memory, the compiler must generate code to load a vector, merge the new data into the vector, and store the vector back into memory.

A review of the assembly code generated by the compiler and annotated by the spu_timing tool (Example 9-5) shows that the code is inefficient for this architecture. Notice the large number of stall cycles. The loop body takes 126 cycles to execute.

Example 9-5  spu_timing output for suboptimal code

```
.L211:
002992 1                                            2                   hbrp    # 2
002993 0                                             3456               shli    $17,$25,3
002994 0                                              4567              shli    $79,$25,4
002995 0                                               5678             shli    $35,$25,2
002996 0D                                               67              il      $61,4
002996 1D                                               6               lnop
002997 0D                                                78             a       $10,$17,$82
002997 1D 012                                            789            lqx     $39,$17,$82
002998 0D                                                 89             a      $78,$17,$30
002998 1D 0123                                           89            lqx     $36,$79,$80
002999 0D 0                                               9             ai      $6,$35,1
002999 1D 012                                              9            cwx     $38,$79,$80
003000 0D 01                                               ai $7,$35,2
```
The code in Example 9-6, while less intuitive, shows how the shuffle intrinsic can be used to dramatically speed up the rearranging of complex data.

**Example 9-6 Better performing complex number rearrangement code**

```c
// Shuffle patterns
vector unsigned char firstFloat  = (vector unsigned char){  0,  1,  2,  3, 16, 17, 18, 19,
0,  0,  0,  0,  0,  0,  0,  0 };
vector unsigned char secondFloat = (vector unsigned char){  4,  5,  6,  7, 20, 21, 22, 23,
0,  0,  0,  0,  0,  0,  0,  0 };
vector unsigned char thirdFloat  = (vector unsigned char){  8,  9, 10, 11, 24, 25, 26, 27,
0,  0,  0,  0,  0,  0,  0,  0 };
vector unsigned char fourthFloat = (vector unsigned char){ 12, 13, 14, 15, 28, 29, 30, 31,
0,  0,  0,  0,  0,  0,  0,  0 };
vector unsigned char firstDword  = (vector unsigned char){  0,  1,  2,  3,  4,  5,  6,  7,
16, 17, 18, 19, 20, 21, 22, 23 };;

vector float *base0 = (vector float *)&wa[1].prob[0];
vector float *base1 = (vector float *)&wa[1].prob[1];
vector float *base2 = (vector float *)&wa[1].prob[2];
vector float *base3 = (vector float *)&wa[1].prob[3];
short int i;
for ( i=0; i < worklist/problemSize; i=i+2 ) {

    // First, read a quadword from each problem
    vector float q0 = *base0;
    vector float q1 = *base1;
    vector float q2 = *base2;
    vector float q3 = *base3;

    vector float r0 = spu_shuffle(
            spu_shuffle( q0, q1, firstFloat ),
            spu_shuffle( q2, q3, firstFloat ) ,
            firstDword );
```

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This code executes much better on the SPU for the following reasons:

- All loads and stores to main memory are done by using full quadwords, which makes the best use of memory bandwidth.
- The compiler is working exclusively with vectors, avoiding the need to move values into “preferred slots.”

Example 9-7 on page 538 shows the code annotated with the spu_timing tool. This loop body is much shorter at only 23 cycles. It also has virtually no stall cycles. The body of the second loop is approximately five times faster than the body of the first loop. However, examination of the code shows that the second loop is executed one half as many times as the first loop. With the savings between the reduced cycles for the loop body and the reduced number of iterations, the second loop works out to be 10 times faster than the first loop. This result was verified with the simulator as well.
### Example 9-7  spu_timing output for more optimal code

```
.L211:
002992 0D  23       ai      $74,$16,1
002992 1D  234567  lqd     $72,0($15)
002993 0D  3456     shli    $59,$16,4
002993 1D  345678  lqx     $73,$15,$27
002994 0D  4567     shli    $56,$74,4
002994 1D  456789  lqx     $70,$15,$26
002995 0D  56       cgt     $55,$20,$22
002995 1D  0      56789  lqx     $71,$15,$25
002996 0   67       ori     $16,$22,0
002997 0   78       ai      $15,$15,16
002998 Od  89       ai      $22,$22,2
002999 1d 012     -9      shufb  $68,$72,$73,$23
003000 1 0123   shufb  $66,$72,$73,$21
003001 l 1234   shufb  $69,$70,$71,$23
003002 1 2345   shufb  $67,$70,$71,$21
003003 1 3456   shufb  $64,$72,$73,$18
003004 1 4567   shufb  $65,$70,$71,$18
003005 1 5678   shufb  $62,$72,$73,$19
003006 1 6789   shufb  $63,$70,$71,$19
003007 1 7890   shufb  $61,$68,$69,$17
003008 1 8901   shufb  $60,$66,$67,$17
003009 1 9012   shufb  $58,$64,$65,$17
003010 1 0123   shufb  $57,$62,$63,$17
003011 1 123456 stqx   $61,$59,$80
003012 1 234567 stqx   $60,$59,$24
003013 1 345678 stqx   $58,$56,$24
003014 1 456789 stqx   $57,$56,$80
003015 1 5678   brnz   $55,.L211
```

.L252:
Part 4

Systems

This part includes Chapter 10, “SDK 3.0 and BladeCenter QS21 system configuration” on page 541, which covers detailed system installation, configuration, and management topics.
Chapter 10. SDK 3.0 and BladeCenter QS21 system configuration

In this chapter, we discuss the installation of the IBM BladeCenter QS21 blade server and the IBM Software Developer Kit (SDK) 3.0 on a QS21 blade server. We also discuss firmware considerations, blade management considerations. In addition, we suggest a method for installing a distribution so that it uses a minimal amount of the resources of BladeCenter QS21.

Specifically, this chapter includes the following topics:

► 10.1, “BladeCenter QS21 characteristics” on page 542
► 10.2, “Installing the operating system” on page 543
► 10.3, “Installing SDK 3.0 on BladeCenter QS21” on page 560
► 10.4, “Firmware considerations” on page 565
► 10.5, “Options for managing multiple blades” on page 569
► 10.6, “Method for installing a minimized distribution” on page 593

The Cell Broadband Engine Software Development Kit 2.1 Installation Guide Version 2.1, which is available from IBM alphaWorks, explains the necessary steps to install the operating system on a QS21 blade and the additional steps to set up a diskless system.1 In this chapter, we consider this detail and address the topics that are complementary to this guide.

10.1 BladeCenter QS21 characteristics

The BladeCenter QS21 blade server has two 64-bit Cell Broadband Engine (Cell/B.E.) processors that are directly mounted onto the blade planar board in order to provide multiprocessing capability. The memory on a BladeCenter QS21 blade server consists of 18 XDR memory modules per Cell/B.E. chip, which creates 1 GB of memory per Cell/B.E. chip.

The BladeCenter QS21 is a single-wide blade that uses one BladeCenter H slot and can coexist with any other blade in the same chassis. To ensure compatibility with existing blades, the BladeCenter QS21 provides two midplane connectors that contain Gigabit Ethernet links, Universal Serial Bus (USB) ports, power, and a unit management bus. The local service processor supports environmental monitoring, front panel, chip initialization, and the Advanced Management Module Interface of the BladeCenter unit.

The blade includes support for an optional InfiniBand expansion card and an optional Serial Attached SCSI (SAS) card.

Additionally, the BladeCenter QS21 blade server has the following major components:

- Two Cell/B.E. processor chips (Cell/B.E.-0 and Cell/B.E.-1) operating at 3.2 GHz
- 2 GB extreme data rate (XDR) system memory with ECC, 1 GB per Cell/B.E. chip, and two Cell/B.E. companion chips, one per Cell/B.E. chip
- 2x8 Peripheral Component Interconnect Express (PCIe) as high-speed daughter cards (HSDC)
- 1 PCI-X as a daughter card
- Interface to optional DDR2 memory, for use as the I/O buffer
- Onboard Dual Channel Gb-Ethernet controller BCM5704S
- Onboard USB controller NEC uPD720101
- One BladeCenter PCI-X expansion card connector
- One BladeCenter High-Speed connector for 2 times x 8 PCIe buses
- One special additional I/O expansion connector for 2 times 16 PCIe buses
- Four dual inline memory module (DIMM) slots (two slots per Cell/B.E. companion chip) for optional I/O buffer DDR2 VLP DIMMs
- Integrated Renesas 2166 service processor (baseboard management controller (BMC) supporting Intelligent Platform Management Interface (IPMI) and Serial over LAN (SOL))
An important characteristic of the BladeCenter QS21 blade server is that it does not contain onboard hard disk or other storage. Storage on the BladeCenter QS21 blade server can be allocated through a network or a SAS-attached device. In the following section, we discuss the installation of the operating system, through a network storage, on the QS21 blade server.

**Support:** BladeCenter QS21 support is available only with the BladeCenter H Type 8852 Unit.

## 10.2 Installing the operating system

The BladeCenter QS21 blade server does not contain local storage. Storage can be provided through a SAS device or through another server in the same network. We briefly explain the steps to install an operating system through the network storage.

Refer to 10.2.4, “Example of installation from network storage” on page 550, in which we show how to apply these steps through the use of bash scripts.

### 10.2.1 Important considerations

To begin, consider the following requirements and information:

- The BladeCenter QS21 is accessible only through an SOL or serial interface. For a serial interface, a specific UART breakout cable is required.

- A POWER technology-based system with storage is required for initial the installation of Linux, which is necessary for a root file system.

**Alternative:** If you do not have a POWER technology-based system, you can execute your initial installation on a BladeCenter QS21 with USB storage attached.

- A Dynamic Host Configuration Protocol (DHCP) server is required to upload the zImage kernel to the BladeCenter QS21. This can only be done through Ethernet modules on the BladeCenter H chassis.

- While the media tray on the BC-H chassis works on the BladeCenter QS21, it is not a supported feature.

- For Red Hat Enterprise Linux (RHEL) 5.1, an individual kernel zImage must be created for each BladeCenter QS21.
For Fedora 7, the same kernel zImage can be applied across multiple blades and is accessible through the Barcelona Supercomputing Center at the following address:

http://www.bsc.es/

Each blade must have its own separate root file system over the Network File System (NFS) server. This restriction applies even if the root file system is read only.

**Shared read-only directories:** While each BladeCenter QS21 must have its own root file system, some directories can be shared as read only among multiple BladeCenter QS21 blade servers.

SWAP is not supported over NFS. Any root file system that is NFS mounted cannot have a SWAP space.

SELinux cannot be enabled on nfsroot clients.

SDK 3.0 supports both Fedora7 and RHEL 5.1, but it is officially supported only for RHEL 5.1

External Internet access is required for installing SDK 3.0 open source packages.

**Tip:** If your BladeCenter QS21 does not have external Internet access, you can download the SDK 3.0 open source components from the Barcelona Supercomputing Center Web site from another machine that has external Internet access and apply them to the BladeCenter QS21.

### 10.2.2 Managing and accessing the blade server

There are currently six options for managing and configuring the blade server, including the ability to access the blade’s console.
Advanced Management Module through the Web interface

The Advanced Management Module (AMM) is a management and configuration program for the BladeCenter system. Through its Web interface, the AMM allows configuration of the BladeCenter unit, including such components as the BladeCenter QS21. Systems status and an event log are also accessible to monitor errors that are related to the chassis or its connected blades.

Advanced Management Module through a command-line interface

In addition to the AMM being accessible through a Web browser, it is directly accessible through a command-line interface. Through this interface, you can issue commands to control the power and configuration of the blade server along with other components of the BladeCenter unit.

For more information and instructions about using the command-line interface, refer to the *IBM BladeCenter Management Module Command-Line Interface Reference Guide* (part number 42C4887) on the Web at the following address:


Serial over LAN

The SOL connection is one option for accessing the blade’s console. By accessing the blade’s console, you can view the progress of the firmware and access the Linux terminal. By default, the blade server sends output and receives over the SOL connection.

To establish an SOL connection, you must configure the SOL feature and start an SOL session. You must also ensure that the BladeCenter and AMM are configured properly to enable the SOL connection.

For further information and details about establishing an SOL connection, refer to the *IBM BladeCenter Serial over LAN Setup Guide* (part number 42C4885) on the Web at the following address:


Serial interface

In addition to using SOL to access the BladeCenter QS21 server's console, you can use a serial interface. This method requires the connection of a specific UART cable to the BladeCenter H chassis. This cable is not included with the BladeCenter H chassis, so you must access it separately.
Ensure to set the following parameters for serial connection on the terminal client:

- 115200 baud
- 8 data bits
- No parity
- One stop bit
- No flow control

By default, input is provided to the blade server through the SOL connection. Therefore, if you prefer input to be provided through a device connected to the serial port, ensure that you press any key on that device while the server boots.

**SMS utility program**

The System Management Services (SMS) utility program is another utility that can provide, in some cases, more information than what is accessible through the AMM.

To access the SMS utility program, you must have input to the blade’s console (accessible either through a serial interface or SOL) as it is starting. Early in its boot process, press F1 as shown in Example 10-1.

*Example 10-1 Initial BladeCenter QS21 startup*

QS21 Firmware Starting
Check ROM = OK
Build Date = Aug 15 2007 18:53:50
FW Version = "QB-1.9.1-0"

Press "F1" to enter Boot Configuration (SMS)

Initializing memory configuration...
MEMORY
  Modules = Elpida 512Mb, 3200 MHz
  XDRlibrary = v0.32, Bin A/C, RevB, DualDD
  Calibrate = Done
  Test = Done

SYSTEM INFORMATION
  Processor = Cell/B.E.(TM) DD3.2 @ 3200 MHz
  I/O Bridge = Cell Companion chip DD2.x
  Timebase = 26666 kHz (internal)
  SMP Size = 2 (4 threads)
  Boot-Date = 2007-10-26 23:52
  Memory = 2048MB (CPU0: 1024MB, CPU1: 1024MB)
The following configurations on the SMS utility cannot be implemented on the AMM:

- SAS configurations
- Choosing which firmware image to boot from, either TEMP or PERM
- Choosing static IP for network startup (otherwise, the default method is strictly DHCP)

For more information about firmware, refer to 10.4, “Firmware considerations” on page 565.

### 10.2.3 Installation from network storage

The BladeCenter QS21 blade server does not contain storage, and you cannot directly install Linux on a network device attached to the blade server. Therefore, you must create the initial installation on a disk. From this initial installation, you can establish a network boot up that can be used by the BladeCenter QS21.

Due to the POWER technology-based architecture for the Cell/B.E. system, the system to create the initial installation for storage must be on a 64-bit POWER technology-based system. After this installation, you copy the resulting root file system to an NFS server, make it network bootable so that it can be mounted via NFS, and adapt it to the specifics of the individual blade server.

#### Setting up the root file system

We begin by obtain a root file system that can be NFS mounted to the BladeCenter QS21. To set up the NFS:

1. Install RHEL 5.1 or Fedora 7 on a 64-bit POWER-technology based system.
2. Copy the root file system from the POWER-technology based installation to an NFS server.

**Multiple copies:** You can create multiple copies of this file system if you want to apply the same distribution on multiple BladeCenter QSS1 systems. For more information, refer to 10.2.4 “Example of installation from network storage” on page 550.

3. Edit the copied root file system to reflect the specific blade to which you want to mount the file system.
Obtaining a zImage kernel with the NFS root enabled

The RHEL 5.1 and Fedora 7 default kernels do not have options that allow NFS root to be enabled. Therefore, you are unable to boot an nfsroot system on these default kernels.

For these reasons, you must have a zImage kernel with an initial RAM disk (initrd) that supports booting from NFS and apply it to your BladeCenter QS21 through a Trivial File Transfer Protocol (TFTP) server.

To obtain a zImage kernel and apply it to the TFTP server:

1. Obtain or create your zImage file:
   - For Fedora 7, download the zImage file from the following address:
     http://www.bsc.es/projects/deepcomputing/linuxoncell/cellsimulator/sdk3.0/zImage.initrd-2.6.22-5.20070920bsc
   - For RHEL 5.1, create the zImage file as explained in the following steps.
     You must perform these steps on the POWER technology-based system that contains the file system and kernel that you want to mount onto the BladeCenter QS21.
     i. Make sure the correct boot configuration is stored in the zImage kernel. That is, BOOTPROTO=dhcp must be in /etc/sysconfig/network-scripts/ifcfg-eth0.
     ii. Create the initrd image by using the following command:
         # mkinitrd --with=tg3 --rootfs=nfs --net-dev=eth0
         --rootdev=<nfs server>:/<path to nfsroot> \
         ~/initrd-<kernel-version>.img <kernel-version>
     iii. Create the zImage file by using the following command:
         # mkzimage /boot/vmlinuz-<kernel-version>
         /boot/config-<kernel-version> \
         /boot/System.map-<kernel-version> <initrd> \
         /usr/share/ppc64-utils/zImage.stub <zImage>

2. Apply your created or downloaded zImage file to the exported directory of your TFTP server.

Additional files to change: In addition to changing the basic network configuration files on the root file system, you must change a few files to enable NFS root as explained in the examples in 10.2.4, “Example of installation from network storage” on page 550.
Applying the zImage file and root file system

You have now obtained the most important components that are needed to boot up a BladeCenter QS21: the root file system to be mounted and the zImage file. You must establish how to pass these components onto the BladeCenter QS21.

When the BladeCenter QS21 system boots up, it first must access the kernel. Therefore, you must load the zImage file. After the kernel is successfully loaded and booted up, the root file system is mounted via NFS.

You can provide the zImage file to a DHCP server that sends the file to the BladeCenter QS21 by using TFTP as explained in the following steps:

1. Ensure that the DHCP and TFTP packages are installed and the corresponding services are enabled on the server.
2. Place the zImage file in a directory that will be exported. Ensure that this directory is exported and the TFTP server is enabled by editing the /etc/xinet.d/tftp file:

   ```
   disable = no
   server_args             = -s <directory to export> -vvvvv
   ```

3. Edit the /etc/dhcpd.conf file to reflect your settings for the zImage by editing the filename argument.

   ```
   Fedora 7: If you are booting up a Fedora 7 file system, you must add the following option entry to the /etc/dhcpd.conf file:
   
   option root-path "<NFS server>:<path to nfsroot>";
   ```

Starting the BladeCenter QS21

After you have created, modified, and configured the root file system for being exported, and placed the zImage file on a TFTP server, you can start the QS21 blade server.

You can start the BladeCenter QS21 blade server from the following locations:

- The optical drive of the BladeCenter unit media tray
- A SAS storage device, typically one or more hardisks attached to the BladeCenter unit
- A storage device attached to the network

To start the blade server through a device attached to the network, ensure that the boot sequence for the BladeCenter QS21 is set to Network. This configuration can be established through the AMM Web browser by selecting **Blade Tasks → Configuration → Boot Sequence**. You can now start your BladeCenter QS21 system.
10.2.4  Example of installation from network storage

In this section, we take you through an example based on the steps in 10.2.3, “Installation from network storage” on page 547. In this example, we cover specific modifications to the root file system in order for it to successfully start on the BladeCenter QS21 blade server. We implement these steps, mainly by using bash scripts, to show how these steps can be applied in an automated fashion.

In 10.5, “Example of a DIM implementation on a BladeCenter QS21 cluster” on page 577, we show an example of implementing a cluster of QS21 blade servers through the use of the DIM tool.

In this example, we have already installed the Linux distribution of choice on a POWER technology-based system. First, we establish the parameters that will be used for this example as shown in Example 10-2.

Example 10-2  Network settings specific to this example

<table>
<thead>
<tr>
<th>Linux Distribution: RHEL5.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kernel Version: 2.6.18-53.el5</td>
</tr>
<tr>
<td>POWER based system with initial install IP &amp; hostname:</td>
</tr>
<tr>
<td>192.168.170.30/POWERbox</td>
</tr>
<tr>
<td>NFS, PTFTP &amp; DHCP Server: 192.168.170.50</td>
</tr>
<tr>
<td>QS21 Hostnames: qs21cell51-62</td>
</tr>
<tr>
<td>QS21 Private Network IP Address (eth0) : 192.168.170.51-62</td>
</tr>
<tr>
<td>NFS root path: /srv/netboot/QS21/RHEL5.1/boot/192.168.170.51</td>
</tr>
</tbody>
</table>

Note: We use the NFS, TFTP, and DHCP server as the same system. Ideally, you want them to be the same, but the NFS server can be a different machine if preferred for storage purposes.

We use the cell_build_master.sh script, as shown in the following example, to copy the root tree directory from the RHEL 5.1 installation on a POWER technology-based system to the NFS server. This same script modifies some files in this master root file system to prepare it for nfsroot.

root@dhcp-server# ./cell_build_master.sh 192.168.170.50 \ /srv/netboot/QS21/RHEL5.1/master
Example 10-3 shows the complete cell_build_master.sh script.

Example 10-3  cell_build_master.sh script

#!/bin/bash

# QS21 Cell Build Master Root Tree
#
#
#

### Show help information

usage()
{
    cat << EOF
${0##*/} - Creates master root tree for Cell QS21.
usage: $0 [POWER_MACHINE] [DESTINATION_PATH]

Arguments:
    POWER_MACHINE       Server where '/' will be copied from
    DESTINATION_PATH    Path where the master directory will be stored

Example:
    ./cell_build_master 10.10.10.1 /srv/netboot/qs21/RHEL5.1/master

This will copy the root directory from machine '10.10.10.1' and store it in /srv/netboot/qs21/RHEL5.1/master

EOF
    exit 1
}

if [ $# != 2 ]; then
    usage
fi

POWER_MACHINE=$1
DESTINATION_DIR=$2
RSYNC_OPTS="-avp -e ssh -x"

set -u
set -e
### Check if master tree already exists ###########################
test -d $DESTINATION_DIR || mkdir -p $DESTINATION_DIR
### Copy root filesystem from POWER-based machine to NFS server ###
rsync $RSYNC_OPTS $POWER_MACHINE:/ $DESTINATION_DIR
### Remove 'swap', '/' and '/boot' entries from /etc/fstab ########
grep -v "swap" $DESTINATION_DIR/etc/fstab | grep -v " / " \
| grep -v " /boot" > $DESTINATION_DIR/etc/fstab.bak
### Ensure SELinux is disabled ####################################
sed -i "s%^\(SELINUX\=\).*%\\1disabled%" \
$DESTINATION_DIR/etc/selinux/config
The /etc/fstab file has changes that are placed in the /etc/fstab.bak file. This
backup file will eventually overwrite the /etc/fstab file. For now, we have a master
copy for this distribution, so that we can apply it to multiple BladeCenter QS21
blade servers.
Next, we use a copy of this master root file system to edit some files and make it
more specific to the individual BladeCenter QS21s. We use the
cell_copy_rootfs.sh script:
root@dhcp-server# ./cell_copy_rootfs.sh \
/srv/netboot/qs21/RHEL5.1/master /srv/netboot/qs21/RHEL5.1/boot/ \
192.168.70.30 192.168.70.51-62 -i qs21cell 51 - 62
In this case, we copy the master root file system to 12 individual BladeCenter
QS21 blade servers. In addition, we configure some files in each of the copied
root file systems to accurately reflect the network identity of each corresponding
BladeCenter QS21. Example 10-4 shows the complete cell_copy_rootfs.sh
script.
Example 10-4 cell_copy_rootfs.sh script

#!/bin/bash
##################################################################
# QS21 Cell Copy Master Root Filesystem
#
#
#
#
#
##################################################################
### Show help information ########################################
usage()
{
cat << EOF
${0##*/} - Copy Master Root Tree for individual BladeCenter QS21.

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usage: $0 [MASTER] [TARGET] [NFS_IP] [QS21_IP] -i [QS21_HOSTNAME]

Arguments:
MASTER              Full path of master root filesystem
TARGET              Path where the root filesystems of the blades will be stored.
NFS_IP              IP Address of NFS Server
QS21_IP             IP Address of QS21 Blade(s). If creating for multiple blades, put in range form: 10.10.2-12.10
-i [QS21_HOSTNAME]     Hostname of BladeCenter QS21. If creating root filesystems for multiple blades, use:
                        
                        -i <hostname> <first> - <last>
<h                     Show this message

Example:
                        
                        ./cell_copy_master.sh /srv/netboot/qs21/RHEL5.1/master \
/srv/netboot/qs21/RHEL5.1 10.10.10.50 10.10.10.25-30 \
-i cell 25 - 30
                        
                        This will create root paths for QS21 blades cell25 to cell30, with IP addresses ranging from 10.10.10.25 to 10.10.10.25. These paths will be copied from /srv/netboot/qs21/RHEL5.1/master into /srv/netboot/qs21/RHEL5.1/cell<25-30> on NFS server 10.10.10.50

EOF
                        
                        exit 1

}

### Process QS21 IP Address passed ###############################################

proc_ip (){}

PSD_QS21_IP=( `echo "$1"` )
QS21_IP_ARY=( `echo $PSD_QS21_IP | sed "s\#.\# g\"` )
QS21_IP_LENGTH=${#QS21_IP_ARY[*]}
for i in $(seq 0 $((${#QS21_IP_ARY[*]} - 1)))
do
    PSD_RANGE=`echo ${QS21_IP_ARY[i]} | grep ":="`
    if [ "$PSD_RANGE" != "" ]; then
        RANGE=`echo ${QS21_IP_ARY[i]} | \ 
    sed "s#-# #"
    QS21_IP_ARY[i]=new

EOF
                        
                        exit 1

}

### Process QS21 IP Address passed ###############################################

proc_ip (){}

PSD_QS21_IP=( `echo "$1"` )
QS21_IP_ARY=( `echo $PSD_QS21_IP | sed "s\#.\# g\"` )
QS21_IP_LENGTH=${#QS21_IP_ARY[*]}
for i in $(seq 0 $((${#QS21_IP_ARY[*]} - 1)))
do
    PSD_RANGE=`echo ${QS21_IP_ARY[i]} | grep ":="`
    if [ "$PSD_RANGE" != "" ]; then
        RANGE=`echo ${QS21_IP_ARY[i]} | \ 
    sed "s#-# #"
    QS21_IP_ARY[i]=new

EOF
                        
                        exit 1

}

### Process QS21 IP Address passed ###############################################

proc_ip (){}

PSD_QS21_IP=( `echo "$1"` )
QS21_IP_ARY=( `echo $PSD_QS21_IP | sed "s\#.\# g\"` )
QS21_IP_LENGTH=${#QS21_IP_ARY[*]}
for i in $(seq 0 $((${#QS21_IP_ARY[*]} - 1)))
do
    PSD_RANGE=`echo ${QS21_IP_ARY[i]} | grep ":="`
    if [ "$PSD_RANGE" != "" ]; then
        RANGE=`echo ${QS21_IP_ARY[i]} | \ 
    sed "s#-# #"
    QS21_IP_ARY[i]=new

EOF
                        
                        exit 1

}
QS21_TEMP=`echo ${QS21_IP_ARAY[*]}`
for a in `seq $RANGE`; do
  NEW_IP=`echo ${QS21_TEMP[*]} | \sed "s#new#$a#" | sed "s# #.#g"`
  QS21_IP[b]=$NEW_IP
  ((b++))
done
echo ${QS21_IP[*]}
break
fi

if [ -z "$QS21_TEMP" ]; then
echo $PSD_QS21_IP
fi

### Show usage if no arguments passed ###################################
if [ $# = 0 ]; then
  usage
fi

MASTER=$1
TARGET=$2
NFS_IP=$3
QS21_IP=$4
shift 4

QS21_IP=( `proc_ip "$QS21_IP"` )

### Capture QS21 Hostname(s) ############################################
while getopts hi: OPTION; do
  case $OPTION in
    i)
      shift
      BLADES=( $* )
      ;;
    h|?)
      usage
      ;;
esac
done

### If a range of blades is provided, process them here ###################
if [ "${BLADES[2]}" = "-" ]; then
BASE=${BLADES[0]}
FIRST=${BLADES[1]}
LAST=${BLADES[3]}
BLADES=()
for i in `seq $FIRST $LAST`;
do
  BLADES[a]=${BASE}${i}
  ((a++))
done

### Ensure same number of IP and Hostnames have been provided #####
if [ "${#BLADES[*]}" != "${#QS21_IP[*]}" ]; then
  echo "Error: Mismatch in number of IP Addresses & Hostnames"
  exit 1
fi

### Creation & configuration of individual Blade paths ###############
for i in $(seq 0 $((${#BLADES[*]} - 1)))
do
  ### Check if master root filesystem already exists ###########
test -d "${TARGET}${BLADES[i]}" && 
  { echo "target \"${TARGET}/${BLADES[i]}\" exists"; exit 1; }
  ### Copy master root filesystem for a specific blade ####
/usr/bin/rsync -aP --delete ${MASTER}/* 
${TARGET}${BLADES[i]}
  #### Edit /etc/fstab for specific machine ################
echo "$NFS_IP:${TARGET}${BLADES[i]}
  nfs tcp,nolock 1 1" >>
$TARGET/${BLADES[i]}/etc/fstab.bak
echo "spufs /spu
  spufs defaults 0 0" >>
$TARGET/${BLADES[i]}/etc/fstab.bak
cp -f $TARGET/${BLADES[i]}/etc/fstab.bak
$TARGET/${BLADES[i]}/etc/fstab
mkdir $TARGET/${BLADES[i]}/spu
  
  
  

### Setup Network ####################################
echo "Now configuring network for target machine...."
SUBNET=`echo $NFS_IP | cut -f1-2 -d.`
sed -i "s%^${SUBNET}.*%${QS21_IP[i]} ${BLADES[i]}
${BLADES[i]}%" $TARGET/${BLADES[i]}/etc/hosts
sed -i "s%^\(HOSTNAME\=\).*%\\1${BLADES[i]}%" \
$TARGET/${BLADES[i]}/etc/sysconfig/network

\

done
echo "Tree build and configuration completed."
Two Ethernet connections: There might be situations where you want two
Ethernet connections for the BladeCenter QS21 blade server. For example, on
eth0, you have a private network and you want to establish public access to
the BladeCenter QS21 via eth1. In this case, make sure that you edit the
ifcnfg-eth1 file in /etc/sysconfig/network-scripts/. In our example, we edit
/srv/netboot/QS21boot/192.168.170.51/etc/sysconfig/
network-scripts/ifcfg-eth1, so that it reflects the network settings for the
BladeCenter QS21 blade server.
Our root file system is modified and ready to be NFS mounted to the
BladeCenter QS21 blade server. However, first, we must create a corresponding
zImage. For this purpose, we use the BLUEcell_zImage.sh script, which will run
on the POWER technology-based system where the RHEL 5.1 installation was
done:
root@POWERbox# ./cell_zImage 192.168.70.50 2.6.18-53.el5
/srv/netboot/qs21/RHEL5.1/boot/ -i cell 51 - 62
In Example 10-5, we create 12 zImage files, one for each BladeCenter QS21
blade server.
Example 10-5 cell_zImage.sh script

#/bin/bash
##################################################################
# QS21 zImage Creation Script
#
#
#
#
#
#
#
##################################################################
set -e
### Show help information ########################################

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usage()
{
  cat << EOF
${0##*/} - creates zImage for Cell QS21.

usage: $0 [NFS_SERVER] [KERN_VERSION] [NFS_PATH] -i [IDENTIFIER]

Arguments:
  NFS_SERVER          Server that will mount the root filesystem
to a BladeCenter QS21.
  KERN_VERSION        Kernel version zImage will be based on
  NFS_PATH            Path on NFS_SERVER where the root
filesystems for blades will be stored

-i [identifier]     Directory name that will identify specific
blade on NFS_SERVER. If creating zImage
for multiple blades, use:

                   -i <hostname> <first> - <last>

-h                   Show this message

Example:

./cell_zImage.sh 10.10.10.1 2.6.18-53.el5 \
/srv/netboot/qs21/RHEL5.1 -i cell 25 - 30

This will create zImages for QS21 blades cell21 to cell34, based
on kernel 2.5.18-53.el5, and whose NFS_PATH will be
10.10.10.1:/srv/netboot/qs21/RHEL5.1/cell<25-30>

EOF
  exit 1
}

if [ $# = 0 ]; then
  usage
fi

NFS_SERVER=$1
KERN_VERSION=$2
NFS_PATH=$3

shift 3
while getopts hi: OPTION; do
  case $OPTION in

### Ensure to set BOOTPROTO=dhcp on eth0 config files ###############

```bash
sed -i "s%^\(BOOTPROTO\=\).*%\1dhcp%" \
/etc/sysconfig/network-scripts/ifcfg-eth0
```

### If a range of blades is provided, process them here ###############

```bash
if [ "$\{BLADES[2]\}" = "-" ]; then
    BASE=\$\{BLADES[0]\}
    FIRST=\$\{BLADES[1]\}
    LAST=\$\{BLADES[3]\}
    BLADES=()
    for i in `seq $FIRST $LAST`;
    do
        BLADES[a]=\$\{BASE\}$\{i\}
        ((a++))
    done
fi
```

### Create the initrd and zImages ################################

```bash
for QS21 in \$\{BLADES\[\*\]\}; do
    ### Create initrd image ##################################
    mkinitrd --with "tg3" --net-dev "eth0" \
    --rootdev=\$\{NFS_SERVER\}:\$\{NFS_PATH\}/\$\{QS21\} --rootfs=nfs \n    initrd-nfs-\$\{QS21\}-\$\{KERN_VERSION\}.img \$\{KERN_VERSION\}

    ### Create kernel zImage #################################
    mkzimage vmlinux-\$\{KERN_VERSION\} config-\$\{KERN_VERSION\} \n    System.map-\$\{KERN_VERSION\} \n    initrd-nfs-\$\{QS21\}-\$\{KERN_VERSION\}.img \n    /usr/share/ppc64-utils/zImage.stub \n    zImage-nfs-\$\{QS21\}-\$\{KERN_VERSION\}.img
```
We copy the zImage file to the DHCP server:

root@POWERbox# scp /boot/zImage.POWERbox-2.6.18-53.el5 root@192.168.170.50:/srv/netboot/QS21/RHEL5.1/images/

Now that we have placed the root file system in our NFS server and the zImage file in our DHCP server, we must ensure that they are accessible by the BladeCenter QS21 blade server. We edit the /etc/exports file in the NFS server to give the proper access permissions as follows:

```
/srv/netboot/QS21/RHEL5.1/boot/192.168.70.51 192.168.170.0(rw,sync,no_root_squash)
```

We then edit the /etc/xinet.d/tftp file. We are mostly interested in the server_args and disable parameters. Example 10-6 shows how our file looks now.

```
Example 10-6   Sample /etc/xinet.d/tftp configuration file

#default: off
# description: The tftp server serves files using the trivial file
# transfer protocol. The tftp protocol is often used to boot diskless
# workstations, download configuration files to network-aware printers,
# and to start the installation process for some operating systems.
service tftp
{
    socket_type = dgram
    protocol = udp
    wait = yes
    user = root
    server = /usr/sbin/in.tftpd
    server_args = -vvv -s /srv/netboot/QS21/images
    disable = no
    per_source = 11
    cps = 100 2
    flags = IPv4
}
```

We pay attention to our /etc/dhcpd.conf file in our DHCP server. In order for changes to the dhcpd.conf file to take effect, we must restart the dhcpd service.
each time we change the dhcp.conf file. To minimize this need, we create a soft
link that points to the zImage file as shown in the following example:

root@192.168.170.50# ln -snf \\
/srv/netboot/QS21/RHEL5.1/images/POWERbox/zImage-POWERbox-2.6.18-53\._el5 \\
/srv/netboot/QS21/images/QS21BLADE

In the future, if we want to change the zImage for the BladeCenter QS21 blade
server, we change where the soft link points instead of editing the dhcpd.conf file
and restarting the dhcpd service.

Next, we ensure that the /etc/dhcpd.conf file is properly set. Example 10-7 shows
how the file looks now.

Example 10-7   Sample entry on /etc/dhcpd.conf file

subnet 192.168.170.0 netmask 255.255.255.0 {
  next-server 192.168.170.50;
}
host QS21Blade{  
  filename "QS21Blade";
  fixed-address 192.168.170.51;
  option host-name "POWERbox";
  hardware ethernet 00:a1:b2:c3:d4:e5;
}

Notice for the filename variable, we call it “QS21Blade”. This file is named in
reference to the directory that is specified on the variable server_args and is
defined in the /etc/xinet.d/tftp file. “POWERbox” is a soft link that we assigned
earlier that points to the appropriate zImage file of interest.

Let us assume that we edited our /etc/dhcpd.conf file. In this case, we must
restart the dhcpd service as shown in the following example:

root@192.168.170.50# service dhcpd restart

We have now completed this setup. Assuming that our BladeCenter QS21 blade
server is properly configured for starting via a network, we can restart it.

10.3 Installing SDK 3.0 on BladeCenter QS21

The BladeCenter QS21 and SDK 3.0 supports the following operating systems:

- RHEL 5.1
- Fedora 7
Furthermore, while SDK 3.0 is available for both of these Linux distributions, SDK 3.0 support is available only for RHEL 5.1.

Additionally, the following SDK 3.0 components are not available for RHEL 5.1:

- Crash SPU commands
- Cell performance counter
- OProfile
- SPU-Isolation
- Full-System Simulator and Sysroot Image

Table 10-1 and Table 10-2 on page 562 provide the SDK 3.0 ISO images that are provided for each distribution, their corresponding contents, and how to obtain them.

<table>
<thead>
<tr>
<th>Product set</th>
<th>ISO name</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Product package is intended for production purposes. This package contains access to IBM support and all of the mature technologies in SDK 3.0.</td>
<td>CellSDK-Product-RHEL 3.0.0.1.0.iso</td>
<td><a href="http://www.ibm.com/software/hotbuy/passportadvantage">http://www.ibm.com/software/hotbuy/passportadvantage</a></td>
</tr>
<tr>
<td>The Extras package contains the latest technologies in the SDK. These packages are usually less mature or contain technology preview code that may or may not become part of the generally available product in the future.</td>
<td>CellSDK-Extra-RHEL 3.0.0.1.0.iso</td>
<td><a href="http://www.ibm.com/developerworks/power/cell/downloads.html">http://www.ibm.com/developerworks/power/cell/downloads.html</a></td>
</tr>
</tbody>
</table>
Table 10-2  Fedora 7 ISO images

<table>
<thead>
<tr>
<th>Product set</th>
<th>ISO name</th>
<th>Locations</th>
</tr>
</thead>
<tbody>
<tr>
<td>The Developer package is intended for evaluation of the SDK in a development environment and contains all of the mature technologies in SDK 3.0.</td>
<td>CellSDK-Devel-Fedora_3.0.0.1.0.iso</td>
<td><a href="http://www.ibm.com/developerworks/power/cell/downloads.html">http://www.ibm.com/developerworks/power/cell/downloads.html</a></td>
</tr>
<tr>
<td>The Extras package contains the latest technologies in the SDK. These packages are usually less mature or contain technology preview code that may or may not become part of the generally available product in the future.</td>
<td>CellSDK-Extra-Fedora_3.0.0.0.1.0.iso</td>
<td><a href="http://www.ibm.com/developerworks/power/cell/downloads.html">http://www.ibm.com/developerworks/power/cell/downloads.html</a></td>
</tr>
</tbody>
</table>

In addition to these packages, there is a set of open source SDK components that are usually accessed directly through YUM from a directory on the Barcelona Supercomputing Center at the following address:

http://www.bsc.es/

If your BladeCenter QS21 blade server does not have outside Internet access, you can download these components into one of your local servers that has external access and install the RPMs manually on the BladeCenter QS21 blade server.

The SDK open source components for RHEL 5.1 of the GCC toolchain and numactl are available for download from the following address:


The following SDK open source components for RHEL 5.1 are included with distribution:

- LIPSPE/LIBSPE2
- netpbm

The following SDK open source components for RHEL 5.1 are not available:

- Crash CPU commands
- OProfile
- Sysroot image
All SDK 3.0 open source components are available for Fedora 7 from the following Web address:


### 10.3.1 Pre-installation steps

To install the SDK 3.0 on a BladeCenter QS21 blade server, we must first complete the following pre-installation steps:

1. Ensure that the BladeCenter QS21 blade server has the appropriate firmware level, which is QB-01.08.0-00 or higher. For more information about firmware, refer to 10.4, “Firmware considerations” on page 565.

2. Ensure that the YUM updater daemon is disabled because it cannot be running when installing SDK. To turn off the YUM updater daemon, type the following command:

   # /etc/init.d/yum-updatesd stop

3. (For RHEL 5.1 only) Complete the following steps:

   a. If you plan on installing FDPR-Pro, ensure that compat-libstdc++ RPM is installed first. Otherwise, the FDPR-Pro installation will fail.

   b. Ensure that the following LIBSPE2 libraries provided in the RHEL 5.1 Supplementary CD are installed:
      
      - libspe2-2.2.0.85-1.el5.ppc.rpm
      - libspe2-2.2.0.85-1.el5.ppc64.rpm

   c. Ensure that the elfspe utility is provided by installing the following elfspe2 rpms as provided in the RHEL 5.1 Supplementary CD:
      
      - elfspe2-2.2.0.85-1.el5.rpm
      - libspe2-devel-2.2.0.85-1.el5.ppc.rpm (if application development applies)
      - libspe2-devel-2.2.0.85-1.el5.ppc64.rpm (if application development applies)

      After installing this rpm, ensure that spufs is loaded correctly:

      # mkdir -p /spu
      # mount /spu

      If you want to mount spufs immediately, enter the following command:

      # mount /spu

      To ensure it automatically mounts on boot, place the following line under /etc/fstab:

      spufs /spu spufs defaults 0 0
4. Ensure that the rsync, sed, tcl, and wget packages are installed in your BladeCenter QS21 blade server. The SDK Installer requires these packages.

5. If you plan to install SDK 3.0 through the ISO images, create the /tmp/sdkiso directory and place the SDK 3.0 ISO images in this directory.

You are now ready to proceed with the installation.

10.3.2 Installation steps

The installation for SDK 3.0 is mostly covered by the SDK Installer, which you obtain after installing the cell-install-<rel>-<ver>.noarch rpm. After you install this rpm, you can install SDK 3.0 by using the cellsdk script with the iso option:

```
# cd /opt/cell
# ./cellsdk --iso /tmp/cellsdkiso install
```

Tip: If you prefer to install SDK 3.0 with a graphical user interface (GUI), you can add the --gui flag when executing the cellsdk script.

Make sure to read the corresponding license agreements for GPL and LGPL and additionally, either the International Programming License Agreement (IPLA) or the International License Agreement for Non-Warranted Programs (ILAN). If you install the “Extras” ISO, you are be presented with the International License Agreement for Early Release of Programs (ILAER).

After you read and accept the license agreements, confirm to begin the YUM-based installation of the specified SDK.

10.3.3 Post-installation steps

You have now installed the default components for SDK 3.0. The only necessary steps at this point are related to configuration updates for YUM.

First, if you do not intend on installing additional packages, you must unmount the ISOs that were automatically mounted when you ran the installation, by entering the following command:

```
# ./cellsdk --iso /tmp/cellsdkiso unmount
```
Note: If you must install additional packages that are included in the SDK 3.0 but are not part of the default installation, you must mount these ISOs again by using the following command:

```
# ./cellsdk --iso /tmp/cellsdkiso mount
```

Then run the following command:

```
'yum install <package_name>'
```

Next, edit the `/etc/yum.conf` file to prevent automatic updates from overwriting certain SDK components. Add the following clause to the `[Main]` section of this file to prevent a YUM update from overwriting SDK versions of the following runtime RPMs:

```
exclude=blas kernel numactl oprofile
```

Re-enable the YUM updater daemon that was initially disabled before the SDK 3.0 installation:

```
# /etc/init.d/yumupdater start
```

You have now completed all of the necessary post-installation steps. If you are interested in installing additional SDK 3.0 components for development purposes, refer to the `Cell Broadband Engine Software Development Kit 2.1 Installation Guide Version 2.1.2` Refer to this same guide for additional details about SDK 3.0 installation and installing a supported distribution on a BladeCenter QS21 system.

### 10.4 Firmware considerations

Firmware for the BladeCenter QS21 comes primarily through two packages, one is through the BMC firmware, and the other through the basic input/output system (BIOS) firmware. In a more detailed manner, each firmware package supports the following functions:

- **BMC firmware**
  - Communicates with advanced management module
  - Controls power on
  - Initializes board, including the Cell/B.E. processors and clock chips
  - Monitors the physical board environment

---

2 See note 1 on page 541.
System firmware
   - Takes control after BMC has successfully initialized the board
   - Acts as BIOS
   - Includes boot-time diagnostics and power-on self test
   - Prepares the system for operating system boot

It is important that both of these packages match at any given time in order to avoid problems and system performance issues.

10.4.1 Updating firmware for the BladeCenter QS21

When updating the firmware for the BladeCenter QS21, we highly recommend that both the BMC and system firmware are updated, with the system firmware being updated first. You can download both of these packages from the following Web address:


Checking current firmware version

There are two ways to check the firmware level on a BladeCenter QS21 blade server. The first way is through the AMM, while the other is through the command line.

The AMM gives information about the system firmware and the BMC firmware. Through this interface, under Monitors → Firmware VPD, you can view the build identifier, release, and revision level of both firmware types.

You can also view the system firmware level through the command line by typing the following command:

# xxd /proc/device-tree/openprom/ibm,fw-vernum_encoded

In the output, the value of interest is the last field, which starts with “QB.” This is your system firmware level.

Updating system firmware

To update the system firmware, you can download the firmware update script from the online IBM support site. After you download the script to your BladeCenter QS21 server, enter the following command on your running BladeCenter QS21:

# ./<firmware_script> -s
This command automatically updates the firmware silently and reboots the machine. You can also extract the .bin system firmware file into a directory by using the following command:

```
# ./<firmware_script> -x <directory to extract to>
```

After you extract the .bin file into a chosen directory, you can update the firmware by using the following command on Linux:

```
# update_flash -f <rom.bin file obtained from IBM support site>
```

**Machine reboot:** Both of the previous two commands cause the BladeCenter QS21 machine to reboot. Make sure that you run these commands when you have access to the machine's console, either via a serial connection or SOL connection.

Now that you have updated and successfully started the new system firmware, ensure that you have a backup copy of this firmware image on your server. There are always two copies of the system firmware image on the blade server:

- **TEMP**
  This is the firmware image that is normally used in the boot process. When you update the firmware, the TEMP image is updated.

- **PERM**
  This is a backup copy of the system firmware boot image. Should your TEMP image be corrupt, you can recover to a working firmware from this copy. We provide more information about this process later in this section.

You can check from which image the blade server has booted by running the following command:

```
# cat /proc/device-tree/openprom/ibm,fw-bank
```

If the output returns “P”, this means that you have booted on the PERM side. Usually, you usually boot on the TEMP side unless that particular image is corrupt.

After you have successfully booted up your blade server from the TEMP image with the new firmware, you can copy this image to the backup PERM side by typing the following command:

```
# update_flash -c
```

Additionally, you can accomplish this task of copying from TEMP to PERM by typing the following command:

```
# echo 0 > /proc/rtas/manage_flash
```
Refer to “Recovering to a working system firmware” on page 568 if you encounter problems with your TEMP image file.

Updating the BMC firmware
After you obtain and uncompress the .tgz file, you have the BMC firmware image whose file name is BNBT<version number>.pkt. To update this image:

1. Log into the corresponding BC-H Advanced Management Module through a browser.
2. Turn off the BladeCenter QS21 whose firmware you’re going to update.
3. Select **Blade Tasks → Firmware Update**.
4. Select the blade server of interest and then click **Update** and **Continue** on the following window.

You should now be set with your updated BMC firmware. You can start your BladeCenter QS21 blade server.

Updating firmware for the InfiniBand daughter card
If you have the optional InfiniBand daughter card on your BladeCenter QS21, you might have to occasionally update the firmware. To complete this task, the openib-tvflash package must be installed.

To update the firmware for this card:

1. Obtain the .tgz packaged file from the IBM support site.
2. Extract it on your BladeCenter QS21 blade server:
   
   ```bash
   # tar xzvf cisco-qs21-tvflash.tgz
   ```
3. Enter the following command:
   
   ```bash
   # ./tvflash -p <firmware .bin file>
   ```

Recovering to a working system firmware
The system firmware is contained in two separate images of the flash memory, which are the temporary (TEMP) image and the permanent (PERM) image. Usually, when a BladeCenter QS21 blade server starts, it starts from the TEMP image. However, in instances where the TEMP image is corrupt or damaged, the system starts from the PERM image.

To choose which image to start from, access the SMS utility. On the main menu, select **Firmware Boot Side Options**.
To check which image your machine is currently booted on, type the following command:

```bash
# cat /proc/device-tree/openprom/ibm,fw-bank
```

A returned value of “P” means that you have booted from the PERM side. If you booted from the PERM side and want to boot from the TEMP side instead:

1. Copy the PERM image to the TEMP image by running the following command:

   ```bash
   # update_flash -r
   ```

2. Shut down the blade server.
3. Restart the blade system management processor through the AMM.
4. Turn on the BladeCenter QS21 blade server again.

## 10.5 Options for managing multiple blades

As previously mentioned, the characteristic of the BladeCenter QS21 blade server of being a diskless system implies the need for additional configurations to ensure initial functionality. Some of these steps, more specifically for RHEL 5.1, require individual BladeCenter QS21 configurations, which can become mundane when amplified to a cluster network of BladeCenter QS21 blade servers.

In this section, we introduce two tools that can be implemented to simplify the steps for scalable purposes and to establish cluster a environment in an organized fashion. The two tools are Extreme Cluster Administration Toolkit (xCAT) and Distributed Image Management (DIM) for Linux Clusters.

### 10.5.1 Distributed Image Management

DIM for Linux Clusters is a tool that was developed for scalable image management purposes. This tool allows diskless blades, in particular, to run a Linux distribution over the network. Additionally, traditional maintenance tasks can be easily and quickly applied to a multitude of blades at the same time. DIM for Linux Clusters was first implemented for use in the IBM MareNostrum supercomputer, which consists of over 2,500 IBM JS21 blades.

DIM for Linux Clusters is a cluster image management utility. It does not contain tools for cluster monitoring, event management, or remote console management. The primary focus of DIM is to address the difficult task of managing Linux distribution images for all nodes of a fairly sized cluster.
DIM includes the following additional characteristics:

- Offers an automated IP and DHCP configuration through an XML file that describes the cluster network and naming taxonomy
- Allows setup of multiple images (that is, it can have Fedora and RHEL 5.1 images setup for one blade) for every node in parallel
- Allows fast incremental maintenance of file system images, changes such as user IDs, passwords, and RPM installations
- Manages multiple configurations to be implemented across a spectrum of individual blades:
  - IP addresses
  - DHCP
  - NFS
  - File system images
  - Network boot images (BOOTP/PXE)
  - Node remote control

While DIM is not a comprehensive cluster management tool, if needed, it can be complemented by other cluster management tools such as xCAT.

DIM has been tested on BladeCenter QS21 blade servers, and the latest release supports DIM implementation on Cell blades.

**DIM implementation on BladeCenter QS21 blade servers**

The most recent release of DIM supports and documents how to implement it for BladeCenter QS21 node clusters. DIM has the ability to provide configuration, setup and provisioning for all these nodes through one or more image servers. We show generalized steps for implementing DIM on a cluster of QS21 blade servers and follow it with an example implementation.

Before installing and setting up DIM, the following prerequisites are required:

- A single, regular disk-based installation on a POWER technology-based machine
- A POWER technology-based or x86-64 machine to be the DIM server
- An image server to store the master and node trees (can be the same as the DIM server)
  
  We recommend that you have at least 20 GB of storage space plus .3 GB per node.
- The following software:
  - BusyBox
  - Perl Net-SNMP
– PERL XML-Simple
– PERL XML-Parser

We assume that the POWER technology-based installation has been already established. Ensure that when you do the initial installation, the Development Tools package is included. We also assume that the DIM server in this case contains the distribution that will be deployed to the BladeCenter QS21 blade servers.

We recommend that you establish a local YUM repository to ease the process of package installation and for instances where your BladeCenter QS21 does not have external Internet access. Assuming that you will apply the rpms from a distribution installation DVD, use the following commands:

```
# mount /dev/cdrom /mnt
rsync -a /mnt /repository
umount /dev/cdrom
rpm -i /repository/Server/createrepo-*.noarch.rpm
createrepos /repository
```

Next, create the /etc/yum.repos.d/local.repo configuration file to reflect this newly created repository:

```
[local]
name=Local repository
baseurl=file:///repository
enabled=1
```

### Setting up the DIM server

On your DIM server, ensure the following packages are installed and enabled:

- dhcp
- xinetd
- tftp-server

Because the DIM loop mounts the BladeCenter QS21 images, you must increase the number of allowed loop devices. Additionally, you want to automatically start DIM_NFS upon startup of the DIM server. You can accomplish both of these tasks by editing the /etc/rc.d/rc.local file so that it is configured as shown in Example 10-8.

**Example 10-8  The /etc/rc.d/rc.local file for DIM server**

```bash
#!/bin/sh
#
# This script will be executed *after* all the other init scripts.
# You can put your own initialization stuff in here if you don't
# want to do the full Sys V style init stuff.
```
DIM_SERVER=<DIM_IP_ADDRESS>
touch /var/lock/subsys/local
if grep -q "^/dev/root / ext" /proc/mounts; then
  # commands for the DIM Server
  modprobe -r loop
  modprobe loop max_loop=64
  if [ -x /opt/dim/bin/dim_nfs ]; then
    /opt/dim/bin/dim_nfs fsck -f
    /opt/dim/bin/dim_nfs start
  fi
else
  # commands for the DIM Nodes
  ln -sf /proc/mounts /etc/mtab
test -d /home || mkdir /home
  mount -t nfs $DIM_SERVER:/home /home
  mount -t spufs spufs /spu
fi
if [ -x /etc/rc.d/rc.local.real ]; then
  . /etc/rc.d/rc.local.real
fi

You must replace the variable DIM_SERVER with your particular DIM server IP address. Next, run the following commands to edit /etc/init.d/halt:

root@dim-server# perl -pi -e \n's#loopfs|autofs#/\readonly\|loopfs|autofs#' /etc/init.d/halt

root@dim-server# perl -pi -e \n's#/\dev/\ram#/(^\dev/\ram|\readonly)/#' /etc/init.d/halt

Editing this file prevent the reboot command from failing on BladeCenter QS21 nodes. Without this fix, the QS21 blade server tries to unmount its own root file system when switching into run level 6.

Now you can reboot your DIM server and proceed to install the DIM.

**Installing the DIM software**

To install DIM on your designated DIM server:

1. Obtain the latest DIM rpm from the IBM alphaWorks Web site at the following address:
   
   http://www.alphaworks.ibm.com/tech/dim
2. Download the additional required software:
   - For Busybox:
     http://www.busybox.net/download.html
   - For Perl Net-SNMP:
     http://www.cpan.org/modules/by-module/Net/Net-SNMP-5.1.0.tar.gz
   - For Perl XML-Parser:
     http://www.cpan.org/modules/by-module/XML/XML-Parser-2.34.tar.gz
   - For Perl XML-Simple

3. Place all of the downloaded packages and the DIM rpm into a created /tmp/dim_install directory. Install the DIM rpm from this directory.

4. Add /opt/dim/bin to the PATH environment variable:
   root@dim-server# echo 'export PATH=$PATH:/opt/dim/bin' >> $HOME/.bashrc
   root@dim-server# . ~/.bashrc

5. Install the additional PERL modules that are required for DIM (Net-SNMP, XML-Parser, XML-Simple). Ignore the warnings that might be displayed.
   root@dim-server# cd /tmp/dim_install
   make -f /opt/dim/doc/Makefile.perl
   make -f /opt/dim/doc/Makefile.perl install

6. Build the Busybox binary and copy it to the DIM directory:
   root@dim-server# cd /tmp/dim_install
   tar xzf busybox-1.1.3.tar.gz
   cp /opt/dim/doc/busybox.config \ 
   busybox-1.1.3/.config
   patch -p0 < /opt/dim/doc/busybox.patch
   cd busybox-1.1.3
   make
   cp busybox /opt/dim/dist/busybox.ppc
   cd /

**Busybox binary:** The Busybox binary must be built on a POWER technology-based system. Otherwise the kernel zimage will not start on a BladeCenter QS21 blade server.

At this point DIM and its dependent packages are installed on your DIM server. Continue with the steps to set up the software.
Setting up the DIM software

In this section, we begin by identifying specific configuration files where modifications can be made to meet your specific network needs. Then we explain the basic setup steps and the options that are available for them.

You must modify the following DIM configuration files to meet your specific network setup:

- `/opt/dim/config/dim_ip.xml`
  
  You must create this file by copying one of the template examples named “dim_ip.xml.<example>” and modifying it as you see fit. The purpose of this file is to set a DIM IP configuration for your Cell clusters. It defines your Cell DIM nodes, their corresponding host names, and IP addresses. It also defines the DIM server.

- `/opt/dim/config/dim.cfg`
  
  This file defines the larger scale locations of the NFS, DHCP, and DIM configuration files. The default values apply in most cases. However, the values for the following configuration parameters frequently change:

  - **DIM_DATA**
    
    In this directory, you store all of the data that is relevant to the distribution that you are deploying across your network. This data includes the zImages, the master root file system, and the blade filesystem images.

  - **NFS_NETWORK**
    
    Here, you define your specific IP address deployment network.

  - **PLUGIN_MAC_ADDR**
    
    Here, you define the IP address of your BladeCenter H. Through this variable, the DIM accesses the BladeCenter H chassis to obtain MAC address information about the QS21 nodes along with executing basic, operational, blade-specific commands.

- `/opt/dim/config/<DISTRO>/dist.cfg`
  
  This file defines variables that are specific to your distribution deployment. It also defines the name of the DIM_MASTER machine, should it be different. You might need to change the following variables:

  - **KERNEL_VERSION**
    
    This is the kernel version that you want to create bootable zImages on your particular distribution.
- **KERNEL_MODULES**
  These are modules that you want and are not enabled by default on your kernel. The only module that you need for BladeCenter QS21 to function is “tg3.”

- **DIM_MASTER**
  This variable defines the machine that contains the master root file system. In cases where the distribution that you want to deploy in your cluster is in a different machine, you specify from which machine to obtain the root file system. Otherwise, if it is in the same box as your DIM server, you can leave the default value.

- **/opt/dim/<DIST>/master.exclude**
  This file contains a list of directories that are excluded from being copied to the master root file system.

- **/opt/dim/<DIST>/image.exclude**
  This file contains a list of directories that are excluded from being copied to the image root file system of each BladeCenter QS21 blade server.

Before you proceed with using the DIM commands to create your images, ensure that you adapt the files and variables defined previously to meet your particular network needs and preferences.

Now you can execute the DIM commands to create your distribution and node specific images. You can find the extent of the DIM commands offered under the /opt/dim/lib directory. Each of these commands should have an accessible manual page on your system:

```
man <dim_command>
```

To create your distribution and node-specific images:

1. Create the DIM master directory for your distribution of interest:
   ```
   root@dim-server# dim_sync_master -d <distribution>
   ```

2. Build the DIM network boot image:
   ```
   root@dim-server# dim_zImage -d <distribution>
   ```

3. Create the read-only and “x” read-write DIM images that represent the number of nodes in your network:
   ```
   root@dim-server# dim_image -d <distribution> readonly
dim-server[y]-blade[[1..x]]
   ```

   Here “y” is the corresponding DIM server number (if there is more than one), and “x” represents the number of BladeCenter QS21 nodes.
4. Add all of these DIM images to /etc/exports:

   root@dim-server# dim_nfs add -d <distribution> all

5. Mount and confirm all the DIM images for NFS exporting:

   root@dim-server# dim_nfs start
   dim_nfs status

**Important:** Ensure that you have enabled the maximum number of loop devices on your dim-server. Otherwise, you will see an error message related to this when running the previous command. To increase the number of loop devices, enter the following command:

   root@dim-server# modprobe -r loop
   modprobe loop max_loop=64

Set the base configuration of DHCPD with your own IP subnet address:

   root@dim-server# dim_dhcp add option -O
   UseHostDeclNames=on:DdnsUpdateStyle=none
   dim_dhcp add subnet -O Routers=<IP_subnet> \ dim-server[1]-bootnet1
   dim_dhcp add option -O DomainName=dim

The following steps require the BladeCenter QS21 blade server to be connected in the BladeCenter H chassis. The previous steps did not require this.

6. Add each QS21 blade to the dhcp.conf file by using the following command:

   root@dim-server# dim_dhcp add node -d <distribution> \ dim-server[y]-blade[x]

   Again “y” represents the dim-server number (if there is more than one dim server) and “x” represents the QS21 blade.

7. After you add all of the nodes of interest, restart the DHCP service:

   root@dim-server# dim_dhcp restart dhcpd

You should now be ready to boot up your QS21 to your distribution as configured under DIM.
After your setup is complete, you can eventually add additionally software to your nodes. For these purposes, apply software maintenance on the original machine where the root file system was copied from. Then you can sync the master root file system and DIM images:

```
root@powerbox# dim_sync_master -d <distribution> <directory updated>..<directory updated>..
```

```
root@powerbox# dim_sync_image -d <distribution> <directory updated>..<directory updated>..
```

Notice that the directories are listed to sync on the master and DIM images, depending on which directories are affected by your software installation.

**Example of a DIM implementation on a BladeCenter QS21 cluster**

In this section, we show an example of applying DIM on a small cluster of BladeCenter QS21 blade servers. This example addresses the issue where POWER technology-based servers are in limited availability. We have two initial installations on a POWER technology-based machine, while the dim-server is an IBM System x machine.

Our example uses the following variables:

- **Distributions:** Fedora 7 and RHEL 5.1
- **Fedora 7 Server:** f7-powerbox
- **RHEL 5.1 Server:** rhel51-powerbox
- **DIM Boot Image Creation Server:** POWER-based
  *qs21-imagecreate(192.168.20.70)*
- **Dim Server:** System x qs21-dim-server(192.168.20.30)
- **Kernel Version:** 2.6.22-5.20070920bsc and 2.6.18-53.el5
- **Dim Nodes:** qs21cell{1..12}(192.168.20.71..82)
- **BladeCenter H IP address:** 192.168.20.50

We chose to use both Fedora 7 and RHEL 5.1 distributions to split our cluster in half between RHEL 5.1 and Fedora. Also, note that we have our DIM server and our image server be two different machines.

Our dim-server is a System x server, while our DIM Boot Image server is a POWER technology-based system. We chose a a POWER technology-based system for our boot image server because such as system is required to create the kernel zImage file. Because we are working with two different distributions, we must copy the root file system from two different a POWER technology-based system installations to our System x server.
We install the DIM only on the dim server. When we create a zImage, we mount the DIM directory to our boot image server. We specify the machine if a procedure is to be applied to only one of these two servers.

**Alternative:** If you do not have a POWER technology-based system, you can install your distribution on a BladeCenter QS21 blade server by using USB storage. Copy this installed file system to your non-a POWER technology-based dim-server, and use the USB storage strictly to create the required boot zImage files.

We do not show the steps for the initial hard-disk installation on a POWER technology-based system. However we indicate that we included the Development Tools package in our installation.

1. We want to give our dim-server the ability to partially manage the BladeCenter QS21 blade server from the dim-server's command prompt. To achieve this, we access the AMM through a Web browser. Then we select **MM Control → Network Protocols → Simple Network Management Protocol (SNMP)** and set the following values:
   - SNMPv1 Agent: enable
   - Community Name: public
   - Access Type: set
   - Hostname or IP: 192.168.20.50

2. Since one of our distributions is Fedora7, we install the kernel rpm that is provided from the Barcelona Supercomputing Center on our Fedora 7 machine:

   ```bash
   root@f7-powerbox# wget \ http://www.bsc.es/projects/deepcomputing/linuxoncell/cellsimulator/sdk3.0/kernel-2.6.22-5.20070920bsc.ppc64.rpm
   rpm -ivh --force --noscripts kernel-2.6.22-5.20070920bsc.ppc64.rpm
   depmod -a 2.6.22-5.20070920bsc
   ```

3. We increase the number of maximum loop devices and start the NFS service by default by editing the file as shown in Example 10-8 on page 571. The file looks the same, except that we assign the variable DIM-Server to the IP address in our example, 192.168.20.70.
4. We make the two changes that are required on the /etc/init.d/halt file on both the f7-powerbox and rhel51-powerbox:

```
root@f7-powerbox# perl -pi -e \\
's#loopfs\|autofs#\/readonly\|loopfs\|autofs#'
/etc/init.d/halt
perl -pi -e \\
's#/^\/dev\/ram/#/(^\/dev\/ram|\/readonly)/#' \
/etc/init.d/halt
```

5. We add all of the QS21 blade host names under /etc/hosts. This applies only to the dim-server:

```
root@dim-server# echo "192.168.20.50 mm mm1" >> /etc/hosts
echo "192.168.20.70 dim-server" >> /etc/hosts
for i in `seq 71 82`; do echo "192.168.20.$i cell$i b$i"; done >> /etc/hosts
```

6. Now we have established all the steps before proceeding to the DIM software installation. We now install the DIM software:

a. For the DIM software installation, first we download the DIM RPM from the IBM alphaWorks Web site at the following address and the required dependent software for DIM:

   http://www.alphaworks.ibm.com/tech/dim

b. We create a /tmp/dim_install directory and place our downloaded DIM rpm in that directory. This applies to both the dim-server and the dim-storage machine:

   ```
   root@dim-server # mkdir /tmp/dim_install
   ```

   **Prerequisite:** Ensure that the following packages are installed before running the following dim_install.sh script:

   - gcc
   - expat-devel

   ```
   root@dim-server # /tmp/dim_install
   ```

   c. We run the following dim_install.sh script on both servers to set up the procedures for installing our DIM software:

   ```
   root@dim-server # ./dim_install.sh
   ```

   Example 10-9 on page 580 shows the dim_install.sh script.
Example 10-9  dim_install.sh script

```
#/bin/bash

set -e

### Download the additional sofware needed by DIM ################
cd /tmp/dim_install
wget http://www.busybox.net/downloads/busybox-1.1.3.tar.gz
wget http://www.cpan.org/modules/by-module/Net/Net-SNMP-5.1.0.tar.gz
wget http://www.cpan.org/modules/by-module/XML/XML-Parser-2.34.tar.gz

### Add /opt/dim/bin to PATH environment variable ################
echo "export PATH=$PATH:/opt/dim/bin" >> $HOME/.bashrc
chmod u+x ~/.bashrc
~/.bashrc

### Install perl modules required for DIM ################
cd /tmp/dim_install
make -f /opt/dim/doc/Makefile.perl
make -f /opt/dim/doc/Makefile.perl install

### Add /opt/dim/bin to PATH environment variable ################
echo "export PATH=$PATH:/opt/dim/bin" >> $HOME/.bashrc
~/.bashrc
echo "Completed"
```
d. The required busybox binary must be built on a POWER technology-based system. Because of this, we build this binary on our boot image creation POWER machine:

```
root@qs21-imagecreate# cd /tmp/dim_install
    tar xzf busybox-1.1.3.tar.gz
    cp /opt/dim/doc/busybox.config \
    busybox-1.1.3/.config
    patch -p0 < /opt/dim/doc/busybox.patch
    cd busybox-1.1.3
    make
    scp busybox \
    root@qs21-dim-server:/opt/dim/dist/busybox.ppc
```

7. With the DIM installed on our server, we must make some modifications to set up DIM:

a. We use one of the example templates (dim_ip.xml.example4) and modify the /opt/dim/config/dim_ip.xml file as needed on both servers.

```
root@dim-server# cp /opt/dim/config/dim_ip.xml.example4 \
    /opt/dim/config/dim_ip.xml
```

Example 10-10 shows the xml file.

```
Example 10-10 The /opt/dim/config/dim_ip.xml file

<?xml version="1.0" ?>
<!-- $Id: dim_ip.xml.example4 1654 2007-09-02 09:03:44Z morjan $ -->
<!--
Simple DIM IP configuration file for CELL
 - boot network 192.168.70.0/24 (eth0)
 - user network 10.0.0.0/8    (eth1, optional)
 - 13 DIM Nodes   (blade 1-13)
 - 1  DIM Server  (blade 14)

Examples:
DIM Component IP-Address Hostname Comment
===============================================================================================
dim-server[1]-bootnet1 192.168.70.14 s1 DIM Server JS21 Slot 14 eth0
dim-server[1]-blade[1]-bootnet1 192.168.70.1 cell1 Cell Blade 1 Slot 1 eth0
dim-server[1]-blade[13]-bootnet1 192.168.70.13 cell13 Cell Blade 13 Slot 13 eth0
dim-server[1]-blade[1]-usernet1 10.0.0.1 cell1u Cell Blade 1 Slot 1 eth1
-->
<dim_ip xmlns:xsi="http://www.w3.org/2001/XMLSchema-instance"
xsi:noNamespaceSchemaLocation="dim_ip.xsd">
    <configuration name="dim">
        <component name="server" min="1" max="1">
```

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The main changes made in the example template revolve around the IP addresses and host names for both the DIM server and the individual BladeCenter QS21 blade servers. This file is open to be modified to meet your specific hardware considerations.

b. We modify /opt/dim/config/dim.cfg to fit our needs as shown in Example 10-11.

Example 10-11 The /opt/dim/config/dim.cfg file

#----------------------------------------------------------------------
# (C) Copyright IBM Corporation 2004
# All rights reserved.
#
#
#----------------------------------------------------------------------

# DIM data directory (no symbolic links !)
DIM_DATA    /var/dim

# DHCPD config file
DHCPD_CONF  /etc/dhcpd.conf
In the file in Example 10-11, we changed the NFS_NETWORK, TFTP_ROOT_DIR, and PLUGIN_MAC_ADDR variables to reflect our settings. The file is shown so that you can see the additional changes that can be configured.
8. Because we are working with two distributions, we must create entries for both distributions:

```
root@qs21-storage# mkdir /opt/dim/config/Fedora7
mkdir /opt/dim/dist/Fedora7
mkdir /opt/dim/config/RHEL51
mkdir /opt/dim/dist/RHEL51

cp /opt/dim/CELL/* /opt/dim/config/Fedora7
cp /opt/dim/dist/CELL/* /opt/dim/dist/Fedora7/
cp /opt/dim/CELL/* /opt/dim/config/RHEL51
cp /opt/dim/dist/CELL/* /opt/dim/dist/RHEL51
```

As you can see, we copied all the files under /opt/dim/CELL and /opt/dim/dist/CELL into our distribution config directories. The files copied from /opt/dim/CELL were dist.cfg, image.exclude, and master.exclude. We configure these files to meet our needs. For the files under /opt/dim/dist/CELL, we do not need to change them.

Example 10-12 and Example 10-13 on page 586 show the dist.cfg file for Fedora7 and RHEL 5.1 respectively.

```
Example 10-12   /opt/dim/config/Fedora7/dist.cfg file

#----------------------------------------------------------------------
# (C) Copyright IBM Corporation 2006
# All rights reserved.
#
#----------------------------------------------------------------------
# $Id: dist.cfg 1760 2007-10-22 16:30:11Z morjan $
#----------------------------------------------------------------------

# Image file size for readwrite part in Megabytes
IMAGE_SIZE_RW     200

# Image file size for readonly part in Megabytes
IMAGE_SIZE_RO     5000

# Read only directories
DIR_RO             bin boot lib opt sbin usr lib64

# Read write directories
DIR_RW             root dev etc var srv selinux

# Create additional directories
DIR_ADD            media mnt proc readonly sys spu huge tmp
home
```
# Ethernet DHCP trials
ETHERNET_DHCP_TRIALS 3

# Reboot delay in seconds on failure
REBOOT_DELAY 60

# Zimage command line options
ZIMAGE_CMDLINE not used

# Mount options readwrite tree
MOUNT_OPTIONS_RW        rw,tcp,nolock,async,rsize=4096,wsize=4096

# Mount options readonly tree
MOUNT_OPTIONS_RO        ro,tcp,nolock,async,rsize=4096,wsize=4096

# Boot method
BOOT_METHOD bootp

# Name of zimage linuxrc file
LINUX_RC linuxrc

# Kernel release (uname -r)
KERNEL_VERSION 2.6.22-5.20070920bsc

# Kernel modules
KERNEL_MODULES tg3.ko sunrpc.ko nfs_acl.ko lockd.ko nfs.ko

# DIM master node       (hostname | [user@]hostname[:port]/module)
DIM_MASTER f7-powerbox

# DIM server            (hostname | [user@]hostname[:port]/module
[,...])
DIM_SERVER localhost

The only variables that we have changed are the KERNEL_VERSION, KERNEL_MODULES, and DIM_MASTER. We additionally modified the MOUNT_OPTIONS_RW and MOUNT_OPTIONS_RO variables to include "tcp".

The rest of the variables have their default values. In this example, the only difference between the Fedora7 and RHEL 5.1 configuration file are the KERNEL_VERSION, KERNEL_MODULES, and DIM_MASTER variables.
Example 10-13  /opt/dim/config/RHEL51/dist.cfg file

#----------------------------------------------------------------------
# (C) Copyright IBM Corporation 2006
# All rights reserved.
#
#----------------------------------------------------------------------
# $Id: dist.cfg 1760 2007-10-22 16:30:11Z morjan $
#----------------------------------------------------------------------

# Image file size for readwrite part in Megabytes
IMAGE_SIZE_RW           200

# Image file size for readonly part in Megabytes
IMAGE_SIZE_RO           5000

# Read only directories
DIR_RO                      bin boot lib opt sbin usr lib64

# Read write directories
DIR_RW                      root dev etc var srv selinux

# Create additional directories
DIR_ADD                      media mnt proc readonly sys spu huge tmp
                home

# Ethernet DHCP trials
ETHERNET_DHCP_TRIALS    3

# Reboot delay in seconds on failure
REBOOT_DELAY            60

# Zimage command line options
ZIMAGE_CMDLINE         not used

# Mount options readwrite tree
MOUNT_OPTIONS_RW        rw,nolock,async,rsize=4096,wsize=4096

# Mount options readonly tree
MOUNT_OPTIONS_RO        ro,nolock,async,rsize=4096,wsize=4096

# Boot method
BOOT_METHOD             bootp

# Name of zimage linuxrc file
LINUX_RC                linuxrc

# Kernel release (uname -r)
KERNEL_VERSION          2.6.18-53.el5

# Kernel modules
KERNEL_MODULES          tg3.ko fscache.ko sunrpc.ko nfs_acl.ko lockd.ko
nfs.ko

# DIM master node       (hostname | [user@]hostname[:port]/module)
DIM_MASTER              rhel51-powerbox

# DIM server            (hostname | [user@]hostname[:port]/module [,...])
DIM_SERVER              localhost

9. The image.exclude and master.exclude text files contain a list of directories to be excluded from image root file systems and the master root file system. We leave the default directories on these files.

10. We create the master image:
   
a. We copy the root file systems from the initial Fedora 7 and RHEL 5.1 machines into our qs21-storage server. We already specified these initial machines in the /opt/dim/config/<DIST>/dist.cfg file.

b. We create public ssh public key authentication for each machine:

   root@dim-server# ssh-keygen -t dsa
   root@dim-server# cat ~/.ssh/id_dsa.pub | ssh root@f7-powerbox
   "cat \ - >> ~/.ssh/authorized_keys"
   root@dim-server# cat ~/.ssh/id_dsa.pub | ssh root@rhel51-powerbox
   "cat - >> ~/.ssh/authorized_keys"

c. We can create the master images:

   root@dim-server# dim_sync_master -d Fedora7
dim_sync_master -d RHEL5.1

11. We create the zImage file. In this case, because our dim-server is a System x machine, we cannot run the command on the dim-server machine. Instead, we mount the /opt/dim, /var/dim, and /srv/tftpboot directories from the System x dim server onto a POWER technology-based machine, which in this case is our boot image server:

   root@dim-server# echo "/opt/dim" >> /etc/exports
   root@dim-server# echo "/var/dim" >> /etc/exports
   root@dim-server# echo "/srv/tftpboot" >> /etc/exports
12. Now that these directories are exportable from the dim-server, we mount them on the POWER technology-based boot image server:

```
root@qs21-imagecreate# mkdir /opt/dim
mkdir /var/dim
mkdir /srv/tftpboot
mount dim-server:/opt/dim /opt/dim
mount dim-server:/var/dim /var/dim
mount dim-server:/srv/tftpboot /srv/tftpboot

echo "export PATH=$PATH:/opt/dim/bin" >> $HOME/.bashrc
```

13. We create the zImage files on the dim boot image server and afterward umount the directories:

```
root@qs21-imagecreate# dim_zimage -d Fedora7
dim_zimage -d RHEL5.1
umount /opt/dim
umount /var/dim
umount /srv/tftpboot
```

14. We return to the dim-server and create the images:

```
root@dim-server# dim_image -d Fedora7 readonly \
dim-server[1]-blade[{1..5}]
    dim_image -d RHEL5.1 readonly \
dim-server[1]-blade[{6..12}]
dim_nfs add -d Fedora7 all
dim_nfs add -d RHEL5.1 all
dim_nfs start
```

15. We add the base configuration to our dhcpd.conf file, this will apply to our dim-server only:

```
root@dim-server# dim_dhcp add option -O \nUseHostDeclNames=on:DdnsUpdateStyle=none
    dim_dhcp add subnet -O Routers=192.168.20.100 \ndim-server[1]-bootnet1
    dim_dhcp add option -O DomainName=dim
```

16. We add the entries into our /etc/dhcpd.conf file:

```
root@dim-server# for i in ‘seq 1 5’; do dim_dhcp add node -d \Fedora7\ dim-server[1]-blade[i]; done
    for i in ‘seq 6 12’; do dim_dhcp add node -d \RHEL5.1 dim-server[1]-blade[i]; done
dim_dhcp restart
```
17. We can use DIM to ensure each blade is configured to boot up via network and also boot up each QS21 blade node:

```
root@dim-server# for i in 'seq 1 12'; do dim_bbs -H mm1 i network;\
        done
        for i in 'seq 1 12'; do dim_bctool -H mm1 i on;\n        done
```

We have now completed implementing DIM on 12 of our QS21 nodes, using both Fedora7 and RHEL 5.1 as deployment distributions.

### 10.5.2 Extreme Cluster Administration Toolkit

The Extreme Cluster Administration Toolkit (xCAT) is a toolkit used for deployment and administration of Linux clusters, with many of its features taking advantage of the System x hardware.

xCAT is written entirely by using scripting languages such as korn, shell, perl, and Expect. Many of these scripts can be altered to reflect the needs of your particular network. xCAT provides cluster management through four main branches, which are automated installation, hardware management and monitoring, software administration, and remote console support for text and graphics.

xCAT supports the following offerings:

- Automated installation
  - Network booting with PXE or Etherboot/GRUB
  - Red Hat installation with Kickstart
  - Automated parallel installation for Red Hat and SUSE®
  - Automated parallel installation via imaging with Windows or other operating systems
  - Other operating system installation using imaging or cloning
  - Automatic node configuration
  - Automatic errata installation
  - Hardware management and monitoring
Hardware management and monitoring
- Supports the Advanced Systems Management features on the System x server
  - Remote power control (on/off/state) via IBM Management Processor Network or APC Master Switch
  - Remote Network BIOS/firmware update and configuration on extensive IBM hardware
  - Remote vital statistics (fan speed, temperatures, voltages)
  - Remote inventory (serial numbers, BIOS levels)
  - Hardware event logs
  - Hardware alerts via SNMP
  - Creation and management of diskless clusters
- Supports remote power control switches for control of other devices
  - APC MasterSwitch
  - BayTech Switches
  - Intel EMP
- Traps SNMP alerts and notify administrators via e-mail

Software administration
- Parallel shell to run commands simultaneously on all nodes or any subset of nodes
- Parallel copy and file synchronization
- Provides installation and configuration assistance of the high-performance computing (HPC) software stack
  - Message passing interface: Build scripts, documentation, automated setup for MPICH, MPICH-GM, and LAM
  - Maui and PBS for scheduling and queuing of jobs
  - GM for fast and low latency interprocess communication using Myrinet

Remote console support for text and graphics
- Terminal servers for remote console
  - Equinox ELS and ESP™
  - iTouch In-Reach and LX series
  - Remote Console Redirect feature in System x BIOS
- VNC for remote graphics
As can be seen, the offerings from xCAT are mostly geared toward automating many of the basic setup and management steps for small and larger, more complex clusters.

As previously mentioned, xCAT’s offerings are broad to the extent that they can complement the cluster management offerings that DIM does not provide. Additionally, xCAT provides its own method of handling diskless clusters. While we do not go into further detail about xCAT’s full offerings and implementation, we briefly discuss xCAT’s solution to diskless clusters.

**Diskless systems on xCAT**

Similar to DIM, a stateless cluster solution can be implemented with xCAT that provides an alternative method for installing and configuring diskless systems.

Warewulf is a tool that is used in conjunction with xCAT to provide a stateless solution for HPC clusters. It was originally designed and implemented by the Lawrence Berkley National Laboratory as part of the Scientific Cluster Support (SCS) program to meet the need of a tool that would allow deployment and management of a large number of clusters. Warewulf provides ease of maintaining and monitoring stateless images and nodes as well as allowing such tasks to be applied in a scalable fashion.

Warewulf provides the following offerings:

- Master/slave relationship
  - Master nodes
    - Supports interactive logins and job dispatching to slaves
    - Gateway between outside world and cluster network
    - Central management for all nodes
  - Slave nodes
    - Slave nodes optimized primarily for computation
    - Only available on private cluster network or networks

- Multiple physical cluster network support
  - Fast Ethernet administration
  - High-speed data networks (that is Myricom, InfiniBand, and GigE)

- Modular design that facilitates customization
  - Can change between kernel, linux distribution, and cluster applications
Network booting

- **Boot image built from Virtual Node File System (VNFS)**
  - A small chroot Linux distribution that resides on the master node
  - Network boot image created using VNFS as a template
  - Destined to live in the RAM on the nodes
- **Nodes boot using Etherboot**
  - Open source project that facilitates network booting
  - Uses DHCP and TFTP to obtain boot image
- **Implements RAM-disk file systems**
  - All nodes capable of running diskless
- **Account user management**
  - Password file built for all nodes
  - Standard authentication schemes (files, NIS, LDAP)
  - Rsync used to push files to nodes

The extent of Warewulf’s offerings rest on providing customization and scalability on a small or large cluster. To learn more about Warewulf, see the following Web site:

http://www.perceus.org/portal/project/warewulf

There are many similarities between Warewulf and DIM, among them, mostly revolving around easing the process of installation and management in an efficient, scalable manner. Both of these tools automate many of the initial installation steps that are required for a BladeCenter QS21 blade server to start.

The primary difference between these two node installation and management tools rests on the machine state. Warewulf provides solutions for stateless machines through the use of RAM-disk file systems that are shared and read only. DIM preserves the state of individual nodes by using NFS root methods to provide read/write images on each node along with making certain components of the file system read only.

Warewulf version 2.6.x in conjunction with xCAT version 1.3 has been tested on BladeCenter QS21s. DIM version 9.14 has been tested on BladeCenter QS21 blade servers.

Both of these offerings are good solutions whose benefits depend on the needs of the particular private cluster setup. In instances where storage is a limited resource and maintaining node state is not important, Warewulf might be better suited for your needs. If storage is not a limited resource and maintaining the state of individual nodes is important, then DIM might be the preferred option.
10.6 Method for installing a minimized distribution

The diskless characteristic of the BladeCenter QS21 blade server requires a common need to have the operating system use a minimal amount of resources. Achieving this process saves on storage space and minimizes memory usage.

This solution of minimal usage of resources by the operating system can be further extended by decreasing the size of the zImage kernel that is loaded on the BladeCenter QS21 blade server. Additionally, the storage footprint of the root file system that is used by each BladeCenter QS21 can be minimized by decreasing the directories to be mounted or making the some directories read only.

We do not cover these two topics in this section because they are beyond the scope of this documentation. Additionally, DIM addresses both of these issues and provides a resource efficient root file system and zImage kernel.

We cover additional steps that you can take during and briefly after installation to remove packages that are not necessary in most cases for a BladeCenter QS21. This example is shown for RHEL 5.1 only, but can closely be applied to Fedora 7 installations as well.

**Note:** These steps are all to be implemented on a POWER technology-based system that contains actual disk storage. The product of this process is then mounted on to a BladeCenter QS21 blade server.
10.6.1 During installation

In this section, we discuss the packages that are installed. While this example is for RHEL 5.1, we do not explain the detailed steps of the installation on RHEL 5.1.

During the package installation step, you must select **Customize Software Selection** as shown in Figure 10-1.

![Figure 10-1 Package Selection step of RHEL 5.1 Installation Process](image)

On the Package Group Selection display (Figure 10-2 on page 595), you are prompted to specify which group of software packages you want to install. Some are already selected by default. Be sure to deselect all of the packages.
Welcome to Red Hat Enterprise Linux Server

+----------¦ Package Group Selection +----------+
¦           |
¦ Please select the package groups you would like to have installed. |
¦           |
¦ [ ] Administration Tools ? |
¦ [ ] Authoring and Publishing ? |
¦ [ ] DNS Name Server |
¦ [ ] Development Libraries |
¦ [ ] Development Tools |
¦ [ ] Editors ? |
¦ [ ] Engineering and Scientific ? |
¦ [ ] FTP Server ? |
¦ [ ] GNOME Desktop Environment |
¦ [ ] GNOME Software Development |
¦ [ ] Games and Entertainment |
¦ [ ] Graphical Internet ? |
¦ [ ] Graphics |
¦ [ ] Java Development |
¦ [ ] KDE (K Desktop Environment) ? |
¦ [ ] KDE Software Development |
¦ [ ] Legacy Network Server |
¦ [ ] Legacy Software Development ? |
¦ [ ] Mail Server ? |
¦ [ ] MySQL Database |
¦ [ ] Network Servers |
¦ [ ] News Server ? |
¦ [ ] Office/Productivity |
¦ [ ] PostgreSQL Database ? |
¦ [ ] Printing Support ? |
¦ [ ] Server Configuration Tools |
¦ [ ] Sound and Video |
¦ [ ] System Tools ? |
¦ [ ] Text-based Internet |
¦ [ ] Web Server ? |
¦ [ ] Windows File Server ? |
¦ [ ] X Software Development |
¦ [ ] X Window System ? |
+----+                +----+
¦ OK ¦                ¦ Back ¦
+----+                +----+

+---------------------------------------------------------------------+
| <Space>,<+>,<-> selection       | <F2> Group Details       | <F12> next screen |
+---------------------------------------------------------------------+

*Figure 10-2  Package Group Selection display*
After you ensure that none of the package groups are selected, you can proceed with your installation.

10.6.2 Post-installation package removal

Now we focus on removing packages that were installed by default. To ease the process of package removal and dependency check, we recommend that you establish a local yum repository in your installed system.

In determining which packages to remove from our newly installed system, our approach is to keep in mind the common purpose of a BladeCenter QS21 server. We remove packages that are related to graphics, video, sound, documentation or word processing, network or e-mail, security, and other categories that contain packages for the BladeCenter QS21 blade server that might be unnecessary.

Graphics and audio
A majority of the packages that are installed by default are related to Xorg and GNOME. Despite the fact we selected to not install these particular package groups during our installation process, nevertheless packages are installed that are related to these tools.

1. Remove the Advanced Linux Sound Architecture (ALSA) library by using the following command statement:

   # yum remove alsa-lib

   The removal of this particular package through YUM also removes other dependent packages, which are listed in Table 10-3.

   \textit{Table 10-3 Packages removed with alsa-lib}

   \begin{tabular}{|l|c|c|c|c|}
   \hline
   Package & GNOME & Sound & Other graphics & Doc & Other \\
   \hline
   alsa-utils & & X & & & \\
   antlr & & & X & & \\
   esound & & X & & & \\
   firstboot & & & X & & \\
   gjdoc & & & X & & \\
   gnome-mount & X & & & & \\
   gnome-python2 & X & & & & \\
   gnome-python2-bonobo & X & & & & \\
   gnome-python2-canvas & X & & & & \\
   \hline
   \end{tabular}
2. Remove the ATK library by using the following command, which adds accessibility support to applications and GUI toolkits:

```
# yum remove atk
```

Removing this package also removes the packages listed in Table 10-4 through YUM.

**Table 10-4 Packages removed with atk**

<table>
<thead>
<tr>
<th>Package</th>
<th>GNOME</th>
<th>Other graphics</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>gnome-python2-extras</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gnome-python2-gconf</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gnome-python2-gnomevfs</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gnome-python2-gtkhtml2</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gnome-vfs2</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>gtkhtml2</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>gtkhtml2-ppc64</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>java-1.4.2-gcj-compat</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>libbonoboui</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>libgcj</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>libgcj-ppc64</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>libgnome</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>libgnomeui</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>rhn-setup-gnome</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>sox</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Package</td>
<td>GNOME</td>
<td>Other graphics</td>
<td>Other</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-------</td>
<td>----------------</td>
<td>-------</td>
</tr>
<tr>
<td>bluez-utils</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>gail</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>gail-ppc64</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>gnome-keyring</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>gtk2</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>gtk2-ppc64</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>gtk2-engines</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libglade2</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libglade2-ppc64</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libgnomecanvas</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libgnomecanvas-ppc64</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libnotify</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>libwnck</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>metacity</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>notification-daemon</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>notify-python</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>pygtk2</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>pygtk2-libglade</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>redhat-artwork</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>usermode-gtk</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>xsri</td>
<td></td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>
3. Remove the X.org X11 runtime library by entering the following statement:
   
   ```bash
   # yum remove libX11
   ```

   Removing this package also removes the packages listed in Table 10-5.

   **Table 10-5 Packages removed with libX11**

<table>
<thead>
<tr>
<th>Package</th>
<th>X.org</th>
<th>Other graphics</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>libXcursor</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXcursor-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXext</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXext-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXfixes</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXfixes-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXfontcache</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXft</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXft-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXi</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXi-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXinerama</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXinerama-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXpm</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXrandr</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXrandr-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXrender</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXrender-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXres</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXtst</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXtst-ppc64</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXv</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXxf86dga</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXxf86misc</td>
<td>X</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
4. Remove the X.org X11 libICE runtime library:

   # yum remove libICE

Removing this package also removes the packages listed in Table 10-6.

*Table 10-6 Packages removed with libICE*

<table>
<thead>
<tr>
<th>Package</th>
<th>X.org</th>
<th>Other graphics</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>libXxf86vm</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>libXxf86vm-ppc64</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>libxkbfile</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>mesa-libGL</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>mesa-libGL-ppc64</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>tclx</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>tk</td>
<td></td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Package</th>
<th>X.org</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>libSM</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libSM-ppc64</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libXTrap</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libXaw</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libXmu</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libXt</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>libXt-ppc64</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>startup-notification</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>startup-notification-ppc64</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>xorg-x11-server-utils</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>xorg-x11-xkb-utils</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>
5. Remove the vector graphics library cairo:

```
# yum remove cairo
```

Removing this package also removes the packages listed in Table 10-7.

*Table 10-7 Packages removed with cairo*

<table>
<thead>
<tr>
<th>Package</th>
<th>Doc</th>
<th>Printing</th>
<th>Other graphics</th>
</tr>
</thead>
<tbody>
<tr>
<td>cups</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>pango</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pango-ppc64</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>paps</td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>pycairo</td>
<td></td>
<td></td>
<td>X</td>
</tr>
</tbody>
</table>

6. Remove the remaining X.org packages:

```
# yum remove libfontenc
```

The following command also removes the libXfont and xorg-x11-font-utils package:

```
# yum remove x11-utils
```

The following command also removes the xorg-x11-utils package:

```
# yum remove libFS libXau libXdmcp xorg-x11-filesystem
```

7. Remove the remaining graphics packages:

```
# yum remove libjpeg
```

The following command also removes the libtiff package:

```
# yum remove libart_lgpl fbset libpng gnome-mime
```

8. Wrap up the audio packages removal:

```
# yum remove audiofile libvorbis talk
```

There are other graphics related packages that are removed by default due to dependencies on other non-graphics packages.
Documentation, word processing, and file manipulation packages
Remove packages that are related with documentation, word processing, and file manipulation:

1. Remove the diffutils file comparison tool:

   # yum remove diffutils

   Removing this package also removes the packages listed in Table 10-8.

   **Table 10-8 Packages removed with diffutils**

<table>
<thead>
<tr>
<th>Packages</th>
<th>X.org</th>
<th>SELinux</th>
<th>GNOME</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>chkfontpath</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>policycoreutils</td>
<td></td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>rhpxl</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>sabayon-apply</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>selinux-policy</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>selinux-policy-targeted</td>
<td></td>
<td></td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>urw-fonts</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>xorg-x11-drv-evdev</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-drv-keyboard</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-drv-mouse</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-drv-vesa</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-drv-void</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-fonts-base</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-server-Xnest</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-server-Xorg</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>xorg-x11-xfs</td>
<td></td>
<td>X</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

2. Remove the relevant packages that are left in this category:

   # yum remove aspell aspell-en ed words man man-pages groff bzip2
   zip\ unzip
Network, e-mail, and printing packages
Now you can move forward and remove any network, printing and e-mail related packages from your default distribution installation:

1. Remove the network packages:
   # yum remove ppp
   The following command removes the rp-pppoe and wvdial packages:
   # yum remove avahi avahi-glib yp-tools ypbind mtr lftp
   NetworkManager
2. Remove the printing related packages:
   # yum remove cups-libs mgetty
3. Remove the e-mail related packages:
   # yum remove mailx coolkey

Security, management, and DOS-related packages
Because we cannot use SELinux for the BladeCenter QS21, we can remove these packages from our default installation:

1. Enter the following command:
   # yum remove checkpolixy setools libsemanage
2. Remove the machine management-related packages:
   # yum remove conman anacron dump vixie-cron
3. Remove the DOS-related packages:
   # yum remove mtools unix2dos dos2unix dosfstools

USB and others
Now you can remove USB packages and other packages that are most likely not needed for the typical BladeCenter QS21 implementation:

# yum remove cccid usbutils

Remove the remaining packages from the RHEL 5.1 default installation:

# yum remove bluez-libs irda-utils eject gpm hdparm hicolor \ ifd-egate iprutils parted pcsc-lite pcsc-lite-libs smartmontools \ wpa_supplicant minicom

At this point, we have removed the packages that most relevant to graphics, audio, word-processing, networking and other tools. This process should reduce the number of installed packages by about 50%.
10.6.3 Shutting off services

The final step in making the system faster and more efficient is to provide a small list of remaining services that can be turned off. This list saves run-time memory and speeds up the BladeCenter QS21 boot process. Use the `chkconfig` command as shown in the following example to turn off the services that are specified:

```bash
# chkconfig --level 12345 atd off
chkconfig --level 12345 auditd off
chkconfig --level 12345 autofs off
chkconfig --level 12345 cpuspeed off
chkconfig --level 12345 iptables off
chkconfig --level 12345 ip6tables off
chkconfig --level 12345 irqbalance atd off
chkconfig --level 12345 isdn off
chkconfig --level 12345 mcstrans off
chkconfig --level 12345 rpcgssd off
chkconfig --level 12345 rhnsd off
```

With the distribution stripped down to a lower amount of installed packages and a minimum amount of services running, you can copy this root file system to a master directory and forward it to individual BladeCenter QS21 blade servers for deployment. As stated previously, DIM takes steps at implementing further resource efficiency. Such implementation can be complemented with the process shown in this section.
Appendixes

In this part, we provide additional information particularly in regard to the references made in this book and the code examples in Chapter 4, “Cell Broadband Engine programming” on page 75. This part contains the following appendixes:

- Appendix A, “Software Developer Kit 3.0 topic index” on page 607
- Appendix B, “Additional material” on page 615
This appendix contains a cross-reference of programming topics relating to Cell Broadband Engine (Cell/B.E.) application development to the IBM Software Developer Kit (SDK) 3.0 documentation that contains that topic. The information in Table A-1 is subject to change when a new SDK is released by IBM.

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The Cell/B.E. SDK 3.0 documentation is installed as part of the install package regardless of the product selected. The following list is the online documentation.

- Software Development Kit (SDK) 3.0 for Multicore Acceleration
  - *Cell Broadband Engine Software Development Kit 2.1 Installation Guide Version 2.1*
  - *Software Development Kit for Multicore Acceleration Version 3.0 Programming Tutorial*
  - *Cell Broadband Engine Programming Handbook*
  - *Security SDK Installation and User’s Guide*

- Programming tools and standards
  - *C/C++ Language Extensions for Cell Broadband Engine Architecture*
  - *IBM Full-System Simulator Users Guide and Performance Analysis*
  - *IBM XL C/C++ single-source compiler*
  - *SPU Application Binary Interface Specification*
  - *SIMD Math Library Specification*
  - *Cell BE Linux Reference Implementation Application Binary Interface Specification*
  - *SPU Assembly Language Specification*
► Programming library documentation
  – *ALF Programmer's Guide and API Reference For Cell For Hybrid-x86 (P)*
  – *BLAS Programmer's Guide and API Reference (P)*
  – *Data Communication and Synchronization Library for Cell Broadband Engine Programmer's Guide and API Reference*
  – *DACS Programmer's Guide and API Reference- For Hybrid-x86 (prototype)*
  – *Example Library API Reference*
  – *Monte Carlo Library API Reference Manual (prototype)*
  – *SPE Runtime Management Library*
  – *SPE Runtime Management Library Version 1.2 to 2.2 Migration Guide*
  – *SPU Timer Library (prototype)*

► Hardware documentation
  – *PowerPC User Instruction Set Architecture - Book I*
  – *PowerPC Virtual Environment Architecture - Book II*
  – *PowerPC Operating Environment Architecture - Book III*
  – *Vector/SIMD Multimedia Extension Technology Programming Environments Manual*
  – *Cell Broadband Engine Architecture*
  – *Cell Broadband Engine Registers*
  – *Synergistic Processor Unit (SPU) InSample caption*
Additional material

This book refers to additional material that can be downloaded from the Internet as described below.

Locating the Web material

The Web material associated with this book is available in softcopy on the Internet from the IBM Redbooks Web server. Point your Web browser to the following address:

ftp://www.redbooks.ibm.com/redbooks/SG247575

Alternatively, you can go to the IBM Redbooks Web site at:

ibm.com/redbooks

Select the Additional materials and open the directory that corresponds with the IBM Redbooks form number, SG247575.
Using the Web material

The additional Web material that accompanies this book includes the SG247575_addmat.zip file.

How to use the Web material

Create a subdirectory (folder) on your workstation, and extract the contents of the Web material compressed file into this folder.

Additional material content

The additional materials file for this book is structured as shown in Figure B-1.

![Folder Content]

Figure B-1  Contents of additional materials file SG247575_addmat.zip

In the rest of this appendix, we describe the contents of the additional material examples in more detail.
Data Communication and Synchronization programming example

In this section, we describe the Data Communication and Synchronization (DaCS) code example that is related to 4.7.1, “Data Communication and Synchronization” on page 291.

DaCS synthetic example

The *dacs_hello_ide.tar* file contains an IDE project in a format that is suitable for import by using the **File → Import** menu options in Eclipse. The code in this file calls almost all the DaCS functions and is both an introduction to the API and a good way to check that DaCS is functioning correctly. Although this is an IDE project, it can also be compiled outside Eclipse by using the **make** command.

Example B-1 shows the contents of the dacs_hello.tar file.

```plaintext
Example: B-1  dacs_hello.tar contents

dacs_hello/.cdtproject
dacs_hello/.project
dacs_hello/.settings/org.eclipse.cdt.core.prefs
dacs_hello/Makefile
dacs_hello/Makefile.example
dacs_hello/README.txt
dacs_hello/dacs_hello.c
dacs_hello/dacs_hello.h
dacs_hello/spu/Makefile
dacs_hello/spu/Makefile.example
dacs_hello/spu/dacs_hello_spu.c
```

Task parallelism and PPE programming examples

In this section, we describe a code example that is related to 4.1, “Task parallelism and PPE programming” on page 77.

Simple PPU vector/SIMD code

The *ppe_vectors* directory contains code that demonstrates simple vector/SIMD instructions on a PPU program. This code is related to Example 4-1 on page 81.
Running a single SPE

The *simple_single_spe directory* contains code that demonstrates how a PPU program can initiate a single SPE thread. This code is related to the following examples:

- Example 4-2 on page 86
- Example 4-3 on page 86
- Example 4-4 on page 89

Running multiple SPEs concurrently

The *simple_two_spes directory* contains code that demonstrate how a PPU program can initiate multiple SPE threads concurrently. The program executes two threads but can be easily extended to use more threads. This code is related to the following examples:

- Example 4-5 on page 90
- Example 4-6 on page 93

Data transfer examples

In this section, we describe code examples that are related to 4.3, “Data transfer” on page 110.

Direct SPE access ‘get’ example

The *ppe_direct_spe_access_get directory* contains a PPU program that uses ordinary load and store instructions to directly access the problem state of some SPEs and initiates the direct memory access (DMA) get command. This code is related to Example 4-14 on page 107.

SPU initiated basic DMA between LS and main storage

The *spe_simple_get_put directory* contains code that demonstrates how the SPU program initiated DMA transfers between the local storage (LS) and main storage. It also shows how to use the stall-and-notify mechanism of the DMA and implementing event handler on the SPU program to handle the stall-and-notify events. This code is related to the following examples:

- Example 4-16 on page 124
- Example 4-17 on page 125
SPU initiated DMA list transfers between LS and main storage

The `spe_dma_list directory` contains code that demonstrates how the SPU program initiates DMA list transfers between LS and main storage. This example also shows how to use the stall-and-notify mechanism of the DMA and implementing the event handler on the SPU program to handle the stall-and-notify events. This code is related to the following examples:

- Example 4-19 on page 129
- Example 4-20 on page 131

PPU initiated DMA transfers between LS and main storage

The `ppe_simple_get_put directory` contains a PPU program that initiates DMA transfers between LS and main storage. This program is related to the following examples:

- Example 4-22 on page 142
- Example 4-23 on page 144

Direct PPE access to LS of some SPEs

The `ppe_direct_ls_access directory` contains a PPU program that uses ordinary load and store instructions to directly access the local storage of some SPEs. This program is related to Example 4-24 on page 146.

Multistage pipeline using LS-to-LS DMA transfer

The `dma_ls_to_ls directory` contains code that uses DMA transfer between LS and LS to implement a multistage pipeline programming mode. This example does not provide the most optimized multistage pipeline model. Rather it demonstrates the potential usage of the LS-to-LS transfer and potentially a starting point for developing a highly optimized multistage pipeline model. This code is not related to any example.
SPU software managed cache

The *sw_cache directory* contains a program that demonstrates how the SPU program initiates the software managed cache and later uses it either to perform synchronous data access by using safe mode or to perform asynchronous data access by using unsafe mode.

This program is related to the following examples:

- Example 4-25 on page 153
- Example 4-26 on page 154
- Example 4-27 on page 155

Double buffering

The *spe_double_buffer directory* contains an SPU program that implements a double-buffering mechanism. This program is related to the following examples:

- Example 4-29 on page 161
- Example 4-30 on page 162
- Example 4-31 on page 164

Huge pages

The *simple_huge directory* contains a PPU program that uses buffers that are allocated on huge pages. This program is related to Example 4-33 on page 170.

Inter-processor communication examples

In this section, we describe the code examples that are related to 4.4, “Inter-processor communication” on page 178.

Simple mailbox

The *simple_mailbox directory* contains a simple PPU and SPU program that demonstrates how to use the SPE inbound and outbound mailboxes. This program is related to the following examples:

- Example 4-35 on page 186
- Example 4-36 on page 188
- Example 4-39 on page 199
**Simple signals**

The `simple_signals directory` contains a simple PPU and SPU program that demonstrates how to use the SPE signal notification mechanism. This program is related to the following examples:

- Example 4-37 on page 195
- Example 4-38 on page 197
- Example 4-39 on page 199
- Example 4-40 on page 202

**PPE event handler**

The `ppe_event_handler directory` contains a PPU program that implements an event handler that handles several SPE events. This program is related to Example 4-41 on page 209.

**SPU programming examples**

In this section, we describe the code examples that are related to 4.6, “SPU programming” on page 244.

**SPE loop unrolling**

The `spe_loop_unroll directory` contains an SPU program that demonstrates how to do the loop unrolling technique to achieve better performance. This program is related to Example 4-52 on page 266.

**SPE SOA loop unrolling**

The `spe_soa_unroll directory` contains an SPU program that demonstrates how to do the loop unrolling by using structure of arrays (SOA) data organization to achieve better performance. This program is related to Example 4-53 on page 269.
SPE scalar-to-vector conversion using insert and extract intrinsics

The `spe_vec_scalar` directory contains an SPU program that demonstrates how to cluster several scalars into a vector, by using `spu_insert` intrinsics, how to perform SIMD operations on them, and how to extract them back to their scalar shape by using the `spu_extract` intrinsic. This program is related to Example 4-59 on page 280.

SPE scalar-to-vector conversion using unions

The `spe_vec_scalar_union` directory contains an SPU program that demonstrates how to cluster several scalars into a vector by using unions. This program is related the following examples:

- Example 4-60 on page 281
- Example 4-61 on page 282
Related publications

The publications listed in this section are considered particularly suitable for a more detailed discussion of the topics covered in this book.

IBM Redbooks

This is the first IBM Redbooks publication regarding the Cell Broadband Engine (Cell/B.E.) Architecture (CBEA). There are no other related IBM Redbooks or Redpaper publications at this time.

Other publications

These publications are also relevant as further information sources:

   [http://www.eecs.berkeley.edu/Pubs/TechRpts/2006/EECS-2006-183.pdf](http://www.eecs.berkeley.edu/Pubs/TechRpts/2006/EECS-2006-183.pdf)
   [http://portal.acm.org/citation.cfm?id=1299042.1299047&coll=GUIDE&dl=](http://portal.acm.org/citation.cfm?id=1299042.1299047&coll=GUIDE&dl=)
http://www.math.sci.hiroshima-u.ac.jp/~m-mat/MT/DC/dgene.pdf
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http://charm.cs.uiuc.edu/papers/Ce11PMUP06.pdf
http://www.ibm.com/support/search.wss?rs=201&tc=SSVKBV&q=mass_cbepp u_docs&rankfile=08


http://sti.cc.gatech.edu/Slides/Acton-070619.pdf

http://cacs.usc.edu/education/cs596/Williams-OptMultiCore-SC07.pdf

http://www.insomniacgames.com/tech/articles/0807/files/dynamic_spu_code.txt


**Online resources**

These Web sites are also relevant as further information sources:

- Cell/B.E. resource center on IBM developerWorks with complete documentation
  http://www-128.ibm.com/developerworks/power/cell/

- Distributed Image Management for Linux Clusters on alphaWorks
  http://alphaworks.ibm.com/tech/dim

- Extreme Cluster Administration Toolkit (xCAT)
  http://www.alphaworks.ibm.com/tech/xCAT/

- OProfile on sourceforge.net
  http://oprofile.sourceforge.net

- IBM Dynamic Application Virtualization
  http://www.alphaworks.ibm.com/tech/dav

- MASS
  http://www.ibm.com/software/awdtools/mass
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Programming the Cell Broadband Engine™ Architecture
Examples and Best Practices

Understand and apply different programming models and strategies

Make the most of SDK 3.0 debug and performance tools

Use practical code development and porting examples

In this IBM Redbooks publication, we provide an introduction to the Cell Broadband Engine (Cell/B.E.) platform. We show detailed samples from real-world application development projects and provide tips and best practices for programming Cell/B.E. applications.

We also describe the content and packaging of the IBM Software Development Kit (SDK) version 3.0 for Multicore Acceleration. This SDK provides all the tools and resources that are necessary to build applications that run IBM BladeCenter QS21 and QS20 blade servers. We show in-depth and real-world usage of the tools and resources found in the SDK. We also provide installation, configuration, and administration tips and best practices for the IBM BladeCenter QS21. In addition, we discuss the supporting software that is provided by IBM alphaWorks.

This book was written for developers and programmers, IBM technical specialists, Business Partners, Clients, and the Cell/B.E. community to understand how to develop applications by using the Cell/B.E. SDK 3.0.

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