PC-Cluster Simulator for Joint Infrastructure Interdependencies Studies

by

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Abstract

Rapid advances in network interface cards have facilitated the interconnection of inexpensive desktop computers to form powerful computational clusters, wherein independent simulations can be run in parallel. In this thesis, a hardware and software infrastructure is developed for the simulation of a complex system of interdependent infrastructures. A PC-Cluster is constructed by interconnecting 16 off-the-shelf computers via high speed Scalable Coherent Interface (SCI) network adapter cards. To enable synchronized data transfer between the cluster nodes with very low latencies, a special library comprised of communication routines was developed based on low-level functions of the SCI protocols. Different interrupt mechanisms for synchronous data transfer between the cluster computers (nodes) were investigated. A new method of implementing the interrupts is developed to achieve a 3.6μs latency for one directional data transfer, which is shown to be an improvement over the standard interrupt mechanisms. To facilitate distributed and concurrent simulation of Simulink models on different computers of the PC-Cluster, a special communication block with the appropriate GUI interface has been developed based on the Simulink S-Function and interfaced with the developed SCI library. A reduced-scale benchmark system comprised of some of the University of British Columbia's infrastructures including the Hospital, Substation, Power House, Water Station, and Steam Station has been chosen to investigate the operation of PC-Cluster and potential improvement in simulation speed compared to a single-computer simulation. The models of considered infrastructures are distributed to different computer nodes of the PC-Cluster and simulated in parallel to study their interdependencies in a case of emergency situations caused by an earthquake or a similar disturbing event. It is shown that an improvement of computational speed over a single computer can be achieved through parallelism using the SCI-based PC-Cluster. For the considered benchmark system, an increase in simulation speed of up to 5 times was achieved.
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UBC, August 2007

Siva Singupuram
To Madhavi, Aravind and Akhil
Chapter 1

Introduction

1.1 Research Motivation

The Joint Infrastructure Interdependencies Research Program (JIIRP) is part of ongoing national efforts to secure and protect Canadians from natural disasters and terrorist attacks [1]. Water utilities, communications, banking, transportation networks and hospitals have many complex interaction points and depend critically on each other to function properly. The interdependencies issue has only recently been recognized as a key factor for optimal decision making where the ultimate outcome of the JIIRP is to produce new knowledge-based practices to better assess, manage, and mitigate risks to the lives of Canadians [2]. The system interactions are complex and dynamic. Simulation of complex heterogeneous systems for coordinated decision making in the case of emergency situations such as an earthquake or a terrorist attack requires an intensive computational tool. Advance performance forecasting of the constituent elements and the system as a whole, is highly essential. This requires that many simulations are run very quickly to find the optimal solution – a sequence of mitigating actions. The development of a monitor to intuitively visualize and comprehend the information on the system dynamics for anticipation of incipient emergencies by the human operators, leads to many research challenges including powerful computing. If the simulation of complex systems is done on a single computer, the physical computing time (the CPU time) can be prohibitively long. In spite of continuous increase in the computational capability of computers, the need to model complex infrastructure systems at increasingly higher levels of detail to study their interrelationships makes the computing time a critical issue. This kind of computationally intensive simulation can be successfully done using a PC-Cluster simulator. The development and implementation of PC-Cluster based simulator is the subject of this thesis.
1.2 Related Work

In the domain of infrastructure modeling, numerous works and studies have focused on the modeling, simulation and analysis of single entity elements. Few projects, however, have attempted to combine multiple-infrastructure networks into one model with the specific intent of analyzing the interactions and interdependencies between the systems. The efforts are being made by Idaho National Engineering and Environment Laboratory (INEEL) to implement parallel simulation for infrastructure modeling at the Idaho Nuclear Technology and Engineering Center (INTEC) to study potential vulnerabilities, emergent behaviours from infrastructure outages, and to assist in the development of protective strategies [3]. The Message Passing Interface (MPI) was used to parallelize the code and facilitate the information exchange between processors [4]. However, the paradigm of computer networking used therein introduces a significant communication overhead leading to slower simulation speed. This limits the study of interdependencies of complex networks where large amount of data exchange between the processors would take place. To achieve low communication latency of the order of micro-seconds, one of the few available interconnect standards such as the Scalable Coherent Interface (SCI) [ANSI/IEEE Std 1596-1992] can be adopted [5].

1.3 System Interconnection Network

System interconnection networks have become a critical component of the computing technology of the late 1990s, and they are likely to have a great impact on the design, architecture, and the use of future high-performance computers. Not only the sheer computational speed but also the efficient integration of the computing nodes into tightly coupled multiprocessor system that distinguishes high performance clusters from desktop systems. Network adapters, switches, and device driver software are increasingly becoming performance critical components in modern supercomputers.

Due to the recent availability of fast commodity network adapter cards and switches, tightly integrated clusters of PCs or workstations can be built, to fill the gap between desktop
systems and supercomputers. These PC-Clusters can be used to perform powerful computation wherein independent tasks can be executed in parallel. The use of commercial off-the-shelf (COTS) technology for both computing and networking enables scalable computing at relatively low costs. Some may disagree, but even the world champion in high-performance computing, Sandia Lab’s Accelerated Strategic Computing Initiative (ASCI) Red Machine [6], may be seen as COTS system. Clearly the system area network plays a decisive role in overall performance.

1.4 Contributions

An approach of achieving a high simulation speed for complex infrastructures is to distribute the whole system over different computers in a system area network using fast network adapters connection that perform the simulation concurrently. Then, these computers can communicate with each other back and forth implementing the interdependencies among the relevant subsystems. As the CPU resource of any one computer is limited, implementing simulations on multi-computer networks offers considerable potential for reducing the overall computing time. However, in a PC-Cluster based simulation, additional time will be required to communicate the data between the computers.

This project represents an important milestone in the quest of a fast computing tool for complex simulations. A novel approach of building a PC-Cluster based simulator via high-speed Scalable Coherent Interface (SCI) network adaptor cards has been proposed, implemented and discussed in this thesis. In this research a 16 nodes PC-Cluster is developed using Dolphin SCI Adapter Cards [7]. The most suitable topology, including other topologies for networking is discussed and implemented. Next a communication library of user callable functions has been developed, in order to use the Dolphin SCI Adapter Card and the associated low-level functions [8]. Different interrupt mechanisms are investigated for back and forth synchronized data transfer. To reduce the communication overhead between the cluster nodes, a new interrupt mechanism has been developed and implemented. A user defined block, based on the communication libraries is developed in MATLAB/SIMULINK.
[9], to communicate between different processors. It is shown that an improvement of computational speed over a single computer can be achieved through parallelism using the SCI-based PC-Cluster. For the considered benchmark system, an increase in simulation speed of up to 5 times was achieved.

1.5 Composition of the Thesis

This thesis is outlined as follows. In Chapter 2, the concepts of system area network and point-to-point links of SCI adapter cards are presented. Several network topologies are examined. Based on preliminary review of SCI-based networks, a specific architecture for building the PC-Cluster at the UBC Power Lab is described that is best suited to accomplish the research objectives. Next, in Chapter 3, possible software protocols are examined and suitable low level functions are chosen to build the communication library of user callable functions. The communication latencies between two nodes of the SCI-based PC-Cluster are measured for different interrupt mechanisms of synchronized data transfer. A MATLAB/SIMULINK S-function block is developed for communicating between SIMULINK models in different nodes. The SIMULINK model of an interdependent reduced-scale UBC benchmark system of infrastructures is implemented and described in Chapter 4. In Chapter 5, the simulation studies and results are compared with those obtained from a conventional single processor implementation. The conclusions are summarized in Chapter 6, which also includes suggestions for the future research.
Chapter 2

SCI-Based PC-Cluster

Scalable Coherent Interface (SCI) is an innovative interconnect standard (ANSI/IEEE Std 1596-1992 [1]) that addresses the high performance computing and networking domain. SCI is used as a high-speed interconnection network (often called system area network) for compute clusters built from commodity workstation nodes. In this chapter, several communication networks for distributed simulations are described and compared qualitatively and quantitatively. SCI Interconnect, System Area Network for Clusters, Point-to-Point Link, and various network topologies evaluated and discussed. Finally the hardware infrastructure of 16 nodes UBC Power lab PC-Cluster is presented.

2.1 Communication Networks

Over the years, numerous communication networks and concepts have been proposed for building clusters or tightly-coupled multiprocessors like NUMA (non-uniform memory access) or CC-NUMA (cache coherent non-uniform memory access) systems. The Myricom’s Myrinet [2] is a high-speed system area and local area network that has its origins in the interconnect technology of a massively parallel machine. Network interface cards attaching to workstations’ I/O buses, high-speed links, switches, and a wealth of software, predominantly optimized message-passing libraries, are available to facilitate the construction of high performance compute clusters. In contrast to SCI, a shared address space across the nodes in a cluster is not provided by the technology. However, the adapter card hosts a programmable processor, which allows specific communication mechanisms to be implemented, among them abstractions emulating the distributed shared memory (DSM [3]). Also the network cost is relatively high because Myrinet requires crossbar switches for network connection [4].
SCRAMNet+ (Shared Common Random Access Memory Network) from Curtis-Wright Controls Embedded Computing (CWCEC) [5] is another cluster interconnects that supports high-bandwidth and low data latency. The SCRAMNet+ approach is a shared-memory network that has a common pool of memory accessible from multiple nodes through PCI cards. The software for Windows NT™ or Linux is commercially available; however, the software must be purchased for each node.

The advertised performance characteristics of all the above PCI-based networks including SCI (Scalable Coherent Interface) from Dolphin Interconnect Solutions, Inc. [6], [7], [8] are summarized in Table 2.1. The latency represents the time needed to transfer the data from one node to another in the associated with each network. In general, this latency depends on the amount of data, i.e. length of message to be transmitted, and is nonzero even for zero length messages.

<table>
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<tr>
<th>Scheme</th>
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<th>Performance</th>
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<td></td>
<td>Zero-Length</td>
<td>1-kB Length</td>
</tr>
<tr>
<td></td>
<td>Message (μs)</td>
<td>Message (μs)</td>
</tr>
<tr>
<td>Myrinet</td>
<td>*2.6 - 3.2</td>
<td>-</td>
</tr>
<tr>
<td>SCRAMNet+</td>
<td>*0.8</td>
<td>*62.0</td>
</tr>
<tr>
<td>SCI D350 Series—PCI Express™ Adapter Cards</td>
<td>*1.4</td>
<td>*4.0</td>
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* Advertised values by the corresponding companies

Table 2.1: Comparison of Myrinet, SCRAMNet+, and SCI PCI cards

As shown in Table 2.1, the Myrinet latency for zero-length message is longer than that of SCRMNet+ network. The reason for the increased latency is due to the number of layers in Myrinet software protocols. On the other hand, compared to SCRAMNet+ and Myrinet, the SCI networks allow sharing of local computer memory between two or more nodes with
relatively low latencies. Moreover, the SCI software for NT and Linux is freely distributed. In this project, D350 Series – SCI-PCI Express™ adapter card [9] was selected because of its advertised low communication latency, low price per card, and the fact that it can be used to implement either shared-memory or message-passing programming paradigms, thus offering more options and future flexibility for the UBC Cluster than other network cards. Features and benefits of the Dolphin PCI-SCI networks [10] include:

- Dolphin PCI-SCI is ANSI/IEEE 1596-1992 scalable and coherent interface compliant.
- Currently, the SCI software supports Linux, Windows NT™, Lynx and Solaris operating systems.
- It is possible to communicate on an SCI cluster using different operating systems on different nodes, for example between Windows NT™ and Linux.
- The SCI supports multiprocessing with very low latency and high data throughput.
- The SCI reduces the delay of inter-processor communication by an enormous factor compared to the newest and best interconnect technologies that are based on generation of networking and I/O protocols (Fiber Channel and ATM), because SCI eliminates the run-time layers of software protocol-paradigm translation [11].

### 2.2 SCI Interconnect

The use of SCI as a cache-coherent memory interconnects allows nodes to be tightly coupled. This application requires SCI to be attached to the memory bus of a node, as shown in Figure 2.1 [12]. At this attachment point, SCI can participate in and “export”, if necessary, the memory and cache coherence traffic on the bus and make the node’s memory visible and accessible to other nodes. The nodes’ memory address ranges (and the address mappings of processes) can be laid out to span a global (virtual) address space, giving processes transparent and coherent access to memory anywhere in the system. Typically this approach is adopted to connect multiple-bus-based commodity SMPs (symmetric multiprocessors) to form a large-scale, cache-coherent (CC) shared-memory system, often termed a CC-NUMA (cache-coherent non-uniform memory access) machine.
The inter-node memory interconnects are proprietary implementations of the SCI standard, with specific adaptations and optimizations incorporated to ease the implementation and integration with the node architecture and to foster overall performance.

![Figure 2.1: SCI-based CC-NUMA multiprocessor model [12]](image)

### 2.3 System Area Network for Cluster

Computer clusters, i.e., networks of commodity workstations or PCs, are becoming ever more important as cost-effective parallel and distributed computing facilities. An SCI system area network can provide high-performance communication capabilities for such a cluster. In this application, the SCI interconnect is attached to the I/O bus of the nodes (e.g., PCI) by a peripheral adapter card, very similar to a LAN; as shown in Figure 2.2[12]. In contrast to a LAN though (and most other system area networks as well), the SCI cluster network, by virtue of the common SCI address space and associated transactions, provides hardware-based physical distributed shared memory. Figure 2.2 shows a high-level view of the DSM (Distributed Shared Memory). An SCI cluster is thus more tightly coupled than a LAN-based cluster, exhibiting the characteristics of a NUMA (non-uniform memory access) parallel machine.
The SCI adapter cards, together with the SCI driver software, establish the DSM as depicted in Figure 2.3[12]. A node, willing to share memory with other nodes (e.g., A), creates shared memory segments in its physical memory and exports them to the SCI network (i.e., SCI address space). Other nodes (e.g., B) import these DSM segments into their I/O address space. Using on-board address translation tables (ATTs), the SCI adapters maintain the mappings between their local I/O addresses and the global SCI addresses. Processes on the nodes (e.g., i and j) may further map DSM segments into their virtual address spaces. The latter mappings are conventionally being maintained by the MMUs (Memory Management Units).
Once the mappings have been set up, internodes communication may be performed by the participating processes at user level, by simple load and store operations into DSM segments mapped from remote memories. The SCI adapters translate I/O bus transactions that result from such memory accesses into SCI transactions, and vice versa, and perform them on behalf of the requesting processor. Thus remote memory accesses are both transparent to the requesting processes and do not need intervention by the operating system. In other words, no protocol stack is involved in remote memory accesses, resulting in low communication latencies even for user-level software.

Currently there is one commercial implementation of such an SCI cluster networks, PCI-SCI adapter cards offered by Dolphin Interconnect Solutions. These adapter cards are used to build a 16 node cluster in this project for parallel computing.

There are two ways to transfer data to and from the SCI network. The first method works as described above: a node’s CPU actively reads data from (writes data to) a remote memory using load (store) operations into the DSM address window mapped from the remote node. This can be fully done on user level, resulting in very low round-trip latencies but occupying CPU for moving the data. The second method involves a direct memory access (DMA) engine in the SCI adapter that copies the data in and out of the node’s memory. While this method relieves the CPU, it has higher start-up costs since the SCI driver software has to be
involved to set up the DMA transfer. As we will be using low level data transfer in this project, the previous method is preferred.

An important property of such SCI cluster interconnect adapters is worth pointing out here. Since an SCI cluster adapter attaches to the I/O bus of a node, it cannot directly observe, and participate in, the traffic on the memory bus of the node. This therefore precludes caching and coherence maintenance of memory regions mapped to the SCI address space. In other words, remote memory contents are basically treated as non-cacheable and are always accessed remotely. Therefore, the SCI cluster interconnect hardware doesn’t implement cache coherence capabilities. Note that this property raises a performance concern: remote accesses (round-trip operations such as reads) must be used judiciously since they are still an order of magnitude more expensive than local memory accesses.

The basic approach to deal with the latter problem is to avoid remote operations that are inherently round-trip, i.e., read, as rare as possible. Rather remote writes are used which are typically buffered by the SCI adapter and therefore, from the point of view of processor issuing the write, experience latencies in the range of local accesses, which are several times faster than remote read operations.

### 2.4 Point-to-Point Links

An SCI interconnect is defined to be built only from unidirectional, point-to-point links between participating nodes. These links can be used for concurrent data transfers, in contrast to the one-at-a-time communications characteristics of buses. The number of links grows as the nodes are added to the system, increasing the aggregate bandwidth of the network. The links can be made fast and their performance can scale with the improvements in the underlying technology.

Most implementations today use parallel links over distances of up to few meters. The data transfer rates and lengths of shared buses are inherently limited due to signal propagation delays and signaling problems on the transmission lines, such as capacitive loads that have to
be driven by the sender, impedance mismatches, and noise and signal reflections on the lines. The unidirectional Point-to-Point SCI links avoid these signalling problems. High speeds are also fostered by low-voltage differential signals.

Furthermore, SCI strictly avoids back-propagating signals; even reverse flow control on the links, in favor of high signalling speeds and scalability. A reverse flow control signal would make timing of, and buffer space required for, a link dependent on the link’s distance [13]. Thus flow control information becomes part of the normal data stream in the reverse direction, leading to the requirement that an SCI node must at least have one outgoing link and one incoming link. The SCI cards include two unidirectional links where each speeds 500 Mbytes/s in system area networks (distances of a few meters) [14].

2.5 Network Topologies

In principle, SCI networks with complex topologies could be built. However, the standard anticipates simple topologies. For small systems, for instance, the preferred topology may be a small ring (a so-called ringlet) as shown in Figure 2.4 [15]. For larger systems, interconnecting the rings using a switch topology as shown in Figure 2.6 or multidimensional tori are feasible. In ring topology, we have the option called 1-D, 1-D with Interleaved connection (shown in Figure 2.5) and 2-D connection.

![Figure 2.4: 4-Node SCI Ring Topology [15]](image-url)
When interconnecting 1-D Adapter Cards with a switch as shown in Figure 2.6, each port of the switch has an input and an output connector, just like a regular node. One or more nodes can be connected to a switch port, following the same scheme as shown above in Figure 2.4 for regular ring topology.
As explained above, the SCI offers considerable flexibility in topology choices all based on the fundamental structure of a ring. However, since a message from one node in a ring must traverse every other node in that ring, this topology becomes inefficient as the number of nodes increases. Multi-dimensional topologies and/or switches are used to minimize the traffic paths and congestion in larger systems [15]. The multi-dimensional topologies assume an equal number of nodes in each dimension. Therefore, for a system with \( D \)-dimensions and \( n \) nodes in each dimension, the total number of nodes (i.e. system size) is equal to \( n^D \). With simple ring, all nodes must share a common communication path, thus limiting the scalability. Hence in this project the two dimensional torus topology (4x4) is considered to achieve optimum performance from the latency perspective.

### 2.5 PC-Cluster at UBC Power Lab

The PC-Cluster built at UBC Power Lab is shown in Figure 2.7 and Figure 2.8. The SCI-D352 PCI-Express™ type 2-D adapter cards are used in this cluster. These adapter cards have two additional SCI links and can be connected in a 2D-torus topology as shown in Figure 2.9 [16]. The nodes in the torus topology are organized in an X and Y direction as shown in Figure 2.10. This means that each node-processor has two SCI interfaces; one interface attaches to the horizontal ringlet (X-direction) and the other attaches to the vertical ringlet (Y-direction). The minimum number of nodes in a 2-D torus cluster is 4 (e.g., 2x2), and maximum number of nodes is 256 (e.g., 16x16). The Node Id is assigned to each node as specified in Figure 2.10 [16].

The cables are connected according to Figure 2.10 [16]. The SCI cables in the X direction should be connected to the main card connectors (SCI-Link 0) and the SCI cables in the Y direction should be connected to the daughter card connectors (SCI-Link 1) as shown in Figure 2.11. The detailed connection with all the node numbers are shown in Figure 2.12.
Figure 2.7: Front view of the 16-Node PC-Cluster at UBC Power Lab
Figure 2.8: Back view of the PC-Cluster showing the wiring interconnecting the 16-Nodes
Figure 2.9: 16 Node SCI Torus Topology (4x4)
Figure 2.10: 4x4-Torus Topology showing the Node Id and X-Y axes

PlugUp Board Connectors
SCI Link 1
SCI Link 0
Main Board Connectors

Figure 2.11: 2D PCI-to-SCI Adapter
Figure 2.12: Wiring of the SCI Network of 16 computers with Node numbers
Chapter 3

SCI Communication Software

Regardless of the selected network topology, the SCI standard is designed to support two main communication software protocols for programming of parallel tasks: Message-Passing and Shared Address Space. The SCI promises efficient implementation of both paradigms as it allows transport of messages with extremely low latency and high bandwidth, thus providing an ideal platform for the message-passing applications. The SCI also provides a direct access to remote memory using plain load-store transactions, thus providing an equally good base for the shared memory applications. In this chapter, the trade-offs between both programming protocols are briefly discussed. The fastest low-level functions [1] are chosen to develop a libraries composed of set of user-callable communication routines in order to implement the distributed simulation using PC-Cluster. The communication latencies for different interrupt mechanisms of synchronized data transfer are measured and compared. Additionally, the MATLAB/SIMULINK [2] S-function block has been developed to facilitate communication between SIMULINK models residing on different cluster nodes.

3.1 SCI Programming

The IEEE SCI standard [3] defines a shared memory interconnect from the physical layer to the transport layer. However, no standard user-callable software layer is defined. In a distributed shared memory environment, little software is required because once a distributed shared memory (DSM) system is set up all accesses can be performed by directly reading/writing from/to the appropriate shared memory spaces. It results in the lowest possible message passing latency and transaction overhead. No procedures or system services need to be called in order to exchange data between system nodes. Although there is no software required to perform DSM data exchange, there is a fair amount of software infrastructure necessary to create an appropriate shared memory segment and to export it into
the global shared address space or to import that global shared memory segment into the local address space of another process.

Transaction overhead and latency are very important features in complex distributed simulations, where distributed shared memory systems are widely used. Therefore the software interface standard for DSM (distributed shared memory) applications was designed in such a way that it does implement all necessary hardware abstraction functions but avoids any additional functionality that would increase the overhead.

In the shared-memory protocol, all the processors in the network share a global memory as shown in Figure 3.1[4]. The SCI DSM constitutes a shared physical address space only, disallowing caching of remote memory contents. It is much more challenging to devise and realize shared-memory or shared-objects abstractions in this environment than it is to implement message passing-programming models. The major challenge involved is to provide a global virtual address space that can be conveniently and efficiently accessed by processes or threads distributed in the cluster. Solutions will also have to address caching and consistency aspects of shared memory models.

Figure 3.1: Shared-memory communication model [4]
For simplicity reason, message passing communication model as shown in Figure 3.2 [4] is often used. Message-passing programming requires to explicitly coding of communication operations. In doing so, it is often simpler to take care of the performance aspects [5]. In this approach, each processor has its own address space and can only access a location in its own memory. The interconnection network enables the processors to send messages to other processors. Also special mechanisms are not necessary in message-passing protocol for controlling simultaneous access to data, thus reducing the computational overhead of a parallel program.

The Dolphin adapter cards allow direct mapping of memory accesses from the I/O bus of a machine to the I/O bus and into the memory of a target machine. This means that the memory in a remote node can be directly accessed by the CPU using store/load operations giving the possibility to bypass the time consuming driver calls in the applications. The high latency of accessing the remote memory as compared to local memory does not make it very attractive to share the program variable over the SCI, since maintaining cache coherence is not possible on the I/O bus. A remote CPU read will stall the CPU, but writes are posted such that the latency is minimized. Message passing model using write-only model fits very well into this scheme, offering a low-latency, high-bandwidth and reliable channel that makes it possible to implement an efficient message passing software interface. Thus message-passing method is adopted in this research because of its straightforward program structure.

Figure 3.2: Message-Passing communication model [4]
3.2 SCI Communication Library

To allow different hardware and software implementations, the IEEE Std P1596.9 “Physical layer Application Programming Interface (API) for the Scalable Coherent Interface (SCI PHY-API)” [6] defines an API to abstract the underlying SCI physical layer. An EU-funded project, named “Standard Software Infrastructures for the SCI-based Parallel Systems” (SISCI) [7] developed highly advanced state-of-the-art software environment and tools to exploit the unique hardware capabilities of the SCI communications. The SISCI API [8] supports:

- **Distributed shared memory (DSM)** whereby memory segment can be allocated on one node and mapped in the virtual address space of a process running on another node. Data is then moved using programmed I/O.
- **DMA (Direct Memory Access) transfers** to move data from one node to another without CPU intervention.
- **Remote interrupt** whereby a process can trigger interrupts on a remote node.
- **Fault tolerance** allowing a process to check if a transfer was successful and to catch asynchronous events (such as link failures) and take appropriate actions.

The low-level SCI software functional specifications were developed by Espirit Project 23174 [1]. This project (Software Infrastructure for SCI, “SISCI”) has defined a common Application Programming Interface to serve as a basis for porting major applications to heterogeneous multi vendor SCI platforms. However, from the requirement analysis [9] it has appeared clear that a unique API for all applications is not realistic.

The functional specification of the API is defined in ANSI C [10]. The following functionality of this SCI API are used here [9].

- Mapping a memory segment residing on a remote node in read/write mode
- Transferring data from remote node via DMA (Direct Memory Access)
- Transferring data to a remote node via DMA
- Getting some information about the underlying SCI system
Sending an interrupt to an application running on a remote node
> Checking if a function execution failed and why
> Checking data transmission errors
> Transferring data to/from a remote node in a blocking way
> Transferring data to/from a remote node in a non-blocking way
> Sending and receiving raw SCI packets
> All functions are thread-safe
> The API implementation doesn’t deteriorate the characteristic performance of the SCI technology (high bandwidth and low-latency)
> Implementing API on more than one platform and making SCI network to become a heterogeneous if the connected nodes are different in terms of platform

The mapping operation is in general very complex and needs to be split in several steps. Basically, when Node A wants to map a memory segment physically residing on Node B, the following sequence has to take place:

• Node B allocates the segment and, via one of its SCI interface, makes it available to Node A
• Node A, via one of its SCI interfaces, connects to the memory segment on Node B and maps it into its own memory

Making a segment available to every other node means essentially mapping this segment to the SCI address space, in such a way that it is visible to other nodes. Connecting to a segment is the complementary action and consists mainly in determining the address range of the segment inside the SCI address space. But it also means maintaining a relationship between the physical segment and its mapped companion. The connection then appears to be a very delicate aspect of the above procedure and it could have strong implications on the lower-level SCI software, i.e. the driver.

Several models can be adopted to abstract the connection management. For the design of SCI API (SCI Application Programming Interface) two of them have been considered [9]. They
differ mainly in how a connection is identified. In what is called the segment-oriented model, a connection is identified by three elements: the physical segment, the SCI adapter used by the creating node, the SCI adapter used by the connecting node. However this doesn't allow distinguishing between all the possible connections, for example in the case when two connections to the same physical memory segments have been performed via the same SCI adapters. The alternative, on which the other model, called connection-oriented, is based, is to assign a unique identifier (handle) to a connection, through which the connection can be referenced.

The connection-oriented model is evidently more flexible, but it is also more complex to manage. For this reason, the design of the API is based at the beginning on the segment-oriented model. There is however the intention to extend the design and then the specification towards the connection-oriented model. In particular the memory management and the connection management are well separated. The procedure described above is then extended as follows:

- On the creating node, an application first allocates a memory segment (SCICreateSegment), assigning to it a host-unique identifier, and prepares it to be accessible by the SCI interface (SCIPrepareSegment). What the preparation means depends on the platform where the software runs. For example, it could consist of making the segment contiguous and locked in physical memory. These two operations concern the memory management aspect of the job.
- Finally, the segment is made available to the external network via one or more SCI interfaces (SCISetSegmentAvailable). This operation represents the connection part of the job.

A segment can be made unavailable to new connections (SCISetSegmentUnavailable) without affecting the already established ones. A segment can be removed only when it is not used any more (SCIRemoveSegment). If the segment is still in use, locally or remotely, the remove function fails.
Figure 3.3 shows a diagram with the possible states that a local segment can occupy for each adapter card and the legal transitions between those states. The state of a local segment can be thought-of as the set of all possible combinations of the states of all the adapters. The function SCIRemoveSegment is a legal operation from all the states, but it fails if the segment is still in use.

![State diagram for a local segment](image)

**Figure 3.3: State diagram for a local segment**

- On the **connecting node**, an application first needs to connect to the remote segment via an SCI interface (SCIConnectSegment) and then it can map it into its own address space (SCIMapRemoteSegment). The two calls create and initialize respectively the descriptors sci_remote_segment and sci_map and for each return a reference handle.

When the segment is not any more in use, it has to be unmapped (SCIUnmapSegment) and disconnected (SCIDisconnectSegment). On the local node, the creation of a memory segment does not imply that it is mapped in the address space of the creating process. To do this, a specific action has to be performed (SCIMapLocalSegment), which also produces a descriptor of type sci_map. The same type descriptor is used for remote mapped segments.

Once the mapping is established, movement of data to or from the mapped segment is performed by a normal assignment to or from a memory location falling in the range where the segment has been mapped in an absolutely transparent way. This also means that there is no software intervention and the typical SCI performance is untouched [9]. Moreover, in
order to profit from hardware-dependent features, a specific function is provided to copy a piece of memory to a remote map (SCIMemCopy). This optimized SCI interface is used in the project to transfer data back and forth between two nodes and achieve low communication latency.

In order to use the SCI adapter cards and the associated low-level functions [1], a library of SCI user-callable functions has been developed. These communication functions are compiled into one static library that can be linked to applications such as C-based MATLAB/Simulink [2] or the Fortran-based languages such as ACSL (Advanced Continuous Simulation Language) [11] and possibly MicroTran [12]. The developed routines implement message passing transactions and are summarized in Table 3.1 and Table 3.2.

The first function that must be called in a user program is scistart (). This function initializes the SCI-library, opens the virtual device and verifies the SCI cards installed on a node. A unique positive integer handle h is returned by this function represents the first connection. This handle is the identifier of the node-to-node connection. In case of failure of virtual device or SCI-library, the handle returned by this function is zero. Next, the function sciinit () is called with respect to the handle h, local Node ID, remote Node ID, and the vector size of exchange variables. This function initializes the network connection corresponding to the handle h, by creating and mapping memory segments and then connecting to remote segments. After initialization of the network corresponding to handle h, data can be transferred between the nodes back and forth using two other functions called scisend () and scireceive (). Once the data transfer is done, the particular network connection link with handle h can be closed using the function sciend (). This unmaps and removes the memory segments corresponding to handle h. Finally, the sciclose () function is called to close the virtual device on the given Node and free the allocated resources and thus terminate from the SCI environment.
### Function Description

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>scistart (int *h)</code></td>
<td>Initializes the SCI-library, opens virtual device and returns handle h = zero, in case of failure.</td>
</tr>
</tbody>
</table>
| `sciinit (int *h, int lnId, int rnId, int vecsize)` | Initializes the network connection corresponding to the handle h.  

- `lnId` - local node ID  
- `rnId` - remote node ID  
- `vecsize` - size of the vectors (number of exchange variables in doubles). Last memory segment is reserved for the message(1) to tell the remote node, that the data transfer is done  
- Creates local segment  
- Prepares the local segment  
- Maps the local segment to user space  
- Sets the local segment available to other nodes  
- Connects to remote segment  
- Maps remote segment to user space |
| `scisend (int *h, double *invec, int vecsize)` | Transfers data from invec (localBuffer) to remote Map using SCIMemCpy. Also sends interrupt message telling remote node that data transfer is done |
| `scireceive (int *h, double *outvec, int vecsize)` | Checks for the interrupt message received from the sending node that transferred the data. Once this message is received, it puts data in outvec, from the localMapAddr. |
| `sciend (int *h)`                 | Closes the network connection corresponding to the handle h.  

- Unmaps the remote segment.  
- Disconnects the remote segment.  
- Unmaps the local segment.  
- Removes the local segment. |
| `sciclose ()`                     | Closes the virtual device and frees the allocated resources. |

Table 3.1: SCI communication library
### Sending Node

**NODE 8**

- **Initialize SCI Library**
- Creates Local Segment+
  Maps Local Segment to User Space(LocalMapAddr)
- Connect to Remote Segment
- Maps Remote Segment to User Space(RemoteMapAddr)
- Transfers Data from LocalBuffer(Invec) to RemoteMap using SCIMemCpy + Sends Interrupt Message to indicate Node4, that Data Transfer is Done
- Unmap Remote Segment
- Disconnect Remote Segment
- Unmap Local Segment
- Remove Local Segment
- Close the Virtual Device
- Free the Allocated resources using SCI Terminate

### Receiving Node

**NODE 4**

- **Initialize SCI Library**
- Creates Local Segment+
  Maps Local Segment to User Space(LocalMapAddr)
- Connect to Remote Segment
- Maps Remote Segment to User Space(RemoteMapAddr)
- Waits Until Receives Interrupt from Node 8 to Know that Data Transfer is done to the LocalMapAddr
- If Receives interruptMessage then the correct Data is transferred to LocalMapAddr + Reads Data into Outvec(LocalBuffer) from LocalMapAddr.
- Unmap Remote Segment
- Disconnect Remote Segment
- Unmap Local Segment
- Remove Local Segment
- Close the Virtual Device
- Free the Allocated resources using SCI Terminate

---

Table 3.2: Sequence of actions between node 8 and node 4
3.3 Measured Communication Latencies

The communication library shown in Table 3.1 can be used in all nodes participating in SCI-Network. Different interrupt mechanisms can be used to obtain a synchronized data transfer. This is investigated in this section and the best method for synchronized back and forth data transfer is used. The communication time between two nodes is measured using the C programs as described in Figure 3.4.

One possible way is to use interrupt mechanism available in SCI-API (SCI application programming interface) [1]. It is possible to send and interrupt to an application running on a remote Node. If an application running on Node A wants to trigger an interrupt for an application running on Node B, the procedure is the following:

- Application B creates an interrupt resource on its node, assigning to it a host-unique identifier (SCICreateInterrupt); a descriptor of type sci_local_interrupt_t is initialized.
- Application A connects to the remote interrupt (SCIConnectInterrupt) that initializes a descriptor of type (sci_remote_interrupt) and after that, can trigger it (SCITriggerInterrupt).

When application A does not need any more to trigger, it disconnects from the remote interrupt (SCIDisconnectInterrupt). Application B frees the interrupt resource with SCIRemoveInterrupt. Application B can catch the interrupt either specifying a call-back function at creation time or using a blocking wait function (SCIWaitForInterrupt).

An alternate method of synchronized data transfer is proposed in this thesis. In the proposed approach, a special bit message flag in the memory segment of the sending node is used. The receiving node checks for that message flag bit to determine whether or not a new message was send. Once the message is received, the data can be accepted into local map address. This kind of polling method does not involve any low-level SCI-API interrupt functions and is very fast.
### Node 4 (computer 1)

```c
#include <stdio.h>
#include <string.h>
#include <stdlib.h>
#include <sci_comm_lib.h>

void main ()
{
    int localNodeId=4; /*computer 1 */
    int remoteNodeId=8; /*computer 2*/
    int vecsize = 100;
    int h = 1;

    double y [100], x [100];
    int i;

    scistart (&h);
    sciinit(&h,localNodeId,
            remoteNodeId, vecsize);
    for (i=0;i<vecsize-1;i++) {
        x[i] = (double)i;
    }
    scireceive (&h, y, vecsize);
    scisend (&h, x, vecsize);

    sciend (&h);
    sciclose (&h);
}
```

### Node 8 (computer 2)

```c
#include <stdio.h>
#include <string.h>
#include <stdlib.h>
#include <sci_comm_lib.h>

void main ()
{
    int localNodeId=8; /*computer 1 */
    int remoteNodeId=4; /*computer 2*/
    int vecsize = 100;
    int h = 1;

    double y [100], x [100];
    int i;

    scistart (&h);
    sciinit(&h,localNodeId,
            remoteNodeId, vecsize);
    for (i=0;i<vecsize-1;i++) {
        x[i] = (double)i;
    }
    scireceive (&h, y, vecsize);
    scisend (&h, x, vecsize);

    sciend (&h);
    sciclose (&h);
}
```

Figure 3.4: C programs in Computer 1 and Computer 2 using communication library

The measured two-way communication latencies are shown in Figure 3.5 for both the standard SCI interrupt handling and the new message polling method of synchronized data transfer. The horizontal axis in Figure 3.5 corresponds to the number of double precision variables exchanged back and forth between the nodes. The vertical axis corresponds to the measured CPU time.
The slope of each line represents the rate of change of the CPU time with respect to the number of double-precision variables exchanged. The vertical-intercept represents the communication overhead associated with the interrupts and other SCI-API functions.

![CPU Time versus Double Precision Variables](image)

**Figure 3.5:** CPU time versus number of doubles using two different methods of synchronized data transfer

As can be seen here, using the API interrupt function for two-way synchronized data transfer, the communication overhead is on the order of 40 µs for the two-way zero-length message. This relatively long time may be due to internal Windows operating system handling of the interrupt mechanism. In the case of message segment method of synchronization, the communication overhead is 7.2 µs for the two-way zero-length data transfer between the two computers. Hence, in this project, the proposed method of synchronization for transferring the data is chosen to build the communication library functions, `scisend()` and `scireceive()`. 
From Figure 3.5, the communication time between two computers for the SCI network for both types of synchronized data transfer, can be approximated as follows:

\[
T_{\text{comp1}} = (0.105 * d + 7.2) \mu s \\
T_{\text{comp2}} = (0.120 * d + 40) \mu s
\]

Where \(d\) is the number of doubles transferred back and forth between two nodes. \(T_{\text{comp1}}\) and \(T_{\text{comp2}}\) are the CPU times for the proposed method of synchronized data transfer and low-level SCI interrupt function method of data transfer.

### 3.4 Simulink S-function Communication Block

Various methods are available for distributed and parallel simulation within Simulink. One of them is the DS Toolbox (Distributed Simulation Toolbox) for Simulink and Stateflow that enables the realization and simulation of distributed Simulink models [13]. In this case, the communication is based on the CORBA/TCP/IP [14]. In analogy to the decomposition concept of Simulink using subsystem blocks with port sets, it provides blocks (called EXITE blocks) with the same structuring functionality but with additional features for parallel and distributed simulation. The user can create their models in the common way and distribute these models on several computers which are interconnected via a standard TCP-IP network. During the simulation, all connected models on all computers run truly in parallel (co-simulation). However this kind of distributed simulation introduces a significant communication overhead, which will slow down the simulation speed of large complex systems. Hence, the SCI communication library described in Section 3, with optimum low-level API functions for achieving low latency is used to develop Simulink S-function user interface block.

The functional diagram of the developed Simulink block is shown in Figure 3.6. This block is then used for distributing a complex Simulink model and simulating it concurrently on the SCI PC cluster. At the beginning of the simulation, the initialization of SCI-library has to
take place. This includes opening of the virtual device and initialization of network connection corresponding to handle $h$ is done using $\text{scistart}()$ and $\text{sciinit}(\&h)$. After that, the data can be transferred to other Simulink models residing on different computers using $\text{scisend}()$ and $\text{scireceive}()$ during every time-step. Finally, at the end of simulation, and after completion of data transfer, the closing of SCI network, virtual device and releasing of the allocated resources is executed using $\text{sciend}()$ and $\text{sciclose}()$. Thus, $\text{scistart}()$, $\text{sciinit}()$, $\text{sciend}()$ and $\text{sciclose}()$ are used only once while $\text{scisend}()$ and $\text{scireceive}()$ are used in every time-step during the whole process of simulation.

Figure 3.6: Distributed Simulink model using SCI-communication S-function Block
Chapter 4

System Description

It should be acknowledged that development of the infrastructure cell models is an on-going research mostly carried out by other graduate students working on the JIIRP project in the UBC Power Lab; whereas the focus of this thesis remains to be software and hardware of the SCI-based PC-Cluster. In this chapter, a reduced-scale UBC test case for joint infrastructure interdependencies is briefly described for completeness. The Simulink models of these infrastructures [1] including the S-function communication block introduced in Chapter 3 are also presented here. The overall reduced-scale benchmark system will be used in this thesis for demonstration and evaluation of distributed simulation using the SCI-based PC-Cluster.

4.1 Reduced-Scale UBC Benchmark System Description

The reduced-scale version of the UBC infrastructures connected with each other form a complex system shown in Figure 4.1. The five infrastructures include Sub-Station, Hospital, Water Station, Power House, and Steam Station. This system is selected because it represents many attributes of a small city [2], which is sufficiently complex to investigate the improvement in the simulation speed that can be obtained if instead of using a single computer this system is distributed over 5 computers of the PC-Cluster. For this comparison, the Simulink models of the infrastructure cells are taken from [1]. The description of a system of infrastructures requires at least two layers: the physical layer and the human layer [3]. In this thesis, only the physical layer model has been considered for distributed simulation.
4.1.1. System components

In describing the model of infrastructures, we use a particular ontology developed by the UBC JIIRP researchers. The components of the physical layer exchange quantities such as power, water, steam, gas and oil, which are defined as tokens [4], [5], and [6]. Generally speaking, the tokens are goods or services that are provided by some entity to another entity that uses them. The infrastructures: Sub-Station, Hospital, Water Station, Power Station, and Steam Station are therefore represented as cells that exchange appropriate tokens. A cell is an entity that performs a function [3]. For example, a Hospital is a cell that uses input tokens:
electricity, water, medicines, etc. and produces output tokens: e.g., beds served. A group of one or multiple cells can form a Node. A Node is a Generator Node if the tokens are produced by its cells or taken out from its storing facilities or exported to other nodes. Similarly, a Node can be a Load Node if it receives tokens from other nodes and then delivers to its internal cells for immediate use or storage. In the reduced-scale UBC benchmark system considered in this thesis, a Node consists of one cell only and it is a Generator Node with respect to some tokens and at the same time a Load Node with respect to other tokens. The nodes are connected by the Transportation Channels. The Transportation Channels are the means by which tokens flow from Generator Node to Load Node. If the channel is broken down, no tokens can be transmitted through that channel.

4.1.2. System equations

Sometimes in modeling a system where goods are produced and consumed, mathematical economics concepts are used [7]. In mathematical economics, similar to systems engineering, the logical relationships between entities and quantities are expressed in terms of mathematical symbols. Due to the complex problem of representing the dynamics of infrastructures during the disasters and due to the highly non-linear nature of the underlying relationships, the mathematical approach is considered to model the system of infrastructures [3], on which the models used in this thesis are based.

4.1.3. Cell modeling

A cell's function can be characterized using an input-output model [3], [8]. For example, a Water Station cell shown in Figure 4.2 uses a number of input tokens: electricity, external-water, and oil to produce an output token: water. A given token in the cell will then be denoted as $x_{kj}$. The first subscript indicates the cell number and the second subscript indicates the type number of token. Assuming Water Station is cell $k = 3$, the tokens in the Water Station cell as shown in Figure 4.2 can be expressed as follows:

$x_{31} = \text{electricity}, x_{32} = \text{external-water}, x_{33} = \text{oil}, x_{34} = \text{water}$
Figure 4.2: Water Station Cell with input and output tokens

Here it is assumed that the relationship among the tokens can be described by some non-linear function as:

\[ x_{34}(t) = f(x_{31}(t), x_{32}(t), x_{33}(t), x_{34}(t)) \]  \hspace{1cm} (4.1)

\( x_{kj} \) = token \textit{j used or produced in k}

\( x_{31} = electricity \textit{used} \)

\( x_{32} = \textit{external - water used} \)

\( x_{33} = oil \textit{used} \)

\( x_{34} = \textit{water produced} \)

The Water Station as a cell is connected to all the networks that provide the tokens it needs to output the product (water). The diagram in Figure 4.3 illustrates these relationships for the UBC Water Station. The Hospital cell connection can also be represented in a similar fashion.
4.1.4. Channel modeling

In general, the tokens needed at the nodes for the cells to do their jobs come from other nodes (apart from local storage) [6]. The tokens travel through the transportation network from Generator Nodes to Load Nodes. A given token may come out of multiple Generator Nodes, and likewise travel to multiple Load Nodes. Thus, there may be multiple channels linking Generator Nodes to Load Nodes. For a given Generator Node, dispatching decisions will determine how the node’s token production will be distributed among the channels coming out of the node. Once the tokens are in the channels, they may be affected by the channel’s capacity and transportation delays. How many token units received by a Load Node at a
given instant of time depends on many factors including the amount of token generated in the system, the dispatching decisions, and the channels capacity and possible delays.

For studying the interdependencies in the UBC benchmark system, specific scenarios can be implemented using the cell and channels features. For example, the water pipe channel may be broken and then repaired after sometime during the study [1]. Also, the channels' gain may be changed to simulate different capacity of the channels. For example, the channel 32 illustrated in Figure 4.4 [3] is the means for transportation of water from Node 3 to Node 2. Here, \( m_{32} \) denotes the magnitude and \( D_{32}(t-k) \) the corresponding time delay. If no water is lost during the transportation, then \( m_{32} \) becomes 1. Also, if the trip takes 20 minutes (assuming one time delay is one hour) then

\[
x_{332}(t) = D_{32}(t - 0.33)
\]

Here subscript \( \lambda \) stands for "link". This equations states that the quantity of water arriving at Node 2 at a given time \( t \) is same as the amount of water sent from Node 3 twenty minutes earlier.

![Figure 4.4: Transportation Channel model](image)

The channel model concept is derived from wave propagation in electrical transmission lines [6]. According to this approach, a wave injected at the sending end of transmission line will arrive at the receiving end after the elapsed traveling time. The maximum channel capacity should be considered as a limit in the dispatch block \( D_{32} \). Broken channels or reduced capacity links during disaster situations can be modeled easily with the model depicted in
Figure 4.4. For example, if an electric power line has to be disconnected for four hours due to a fault, the link model for the line would include the transmission line maximum power capacity plus a delay time of four hours.

4.2 Substation

The Substation is one of the key infrastructure components of the UBC campus. To have a better appreciation for the real physical infrastructure, a picture of the Substation is shown in Figure 4.5. This Substation is the link between BC Hydro transmission network and the UBC campus power network [9]. The electricity is transmitted from BC Hydro via two high voltage 64KV overhead lines. The key components of the Substation are transformers, switch gear equipments, circuit breakers, feeders etc.

Figure 4.5: UBC Substation feeding power to whole campus

The Substation model is built to assess its performance in normal state and in disaster situations that may arise due to an earthquake or a terrorist attack. After an earthquake, the
Substation service may be deteriorated and require repairs to the damaged components in order to restore its normal operating condition. The model takes into consideration the restoration process after the disaster, based on the repairing process as per predefined response plan and appropriate schedules. The UBC Substation Simulink model developed by Liu, L. [1] includes distributor, channels, and tokens. For the distributed simulation using the PC-Cluster, the Substation model is implemented on one computer and interfaced with other components using the S-function communication block as shown in Figure 4.6.

![Figure 4.6: Simulink model of the UBC Substation Cell interfacing the PC Cluster](image)

**4.3 Hospital**

Hospital is considered to be a critical infrastructure. On the one hand, in the event of a disaster like an earthquake, many people would be injured leading to a dramatic increase in the number of patients requiring immediate medical assistance/service. On the other hand, the functionality of the Hospital may be impaired due to the direct and indirect impact of this kind of unforeseen circumstances. For optimum health care services during this situation, proper decision making based on the awareness of the real capability of the infrastructure during emergency situations is essential. A Simulink Hospital model has been developed in [1] to evaluate its performance during normal and abnormal situations where different hypothetical scenarios may be considered. The output of this model is the service it can provide to the patients on urgent, short-term and long-term basis. The Simulink model of the
UBC Hospital Cell includes input tokens, output tokens (number of beds), and the channels. For the distributed simulation using the PC-Cluster, the Hospital Cell has been interfaced with the cluster nodes using the S-function communication block as shown in Figure 4.7.

Figure 4.7: Simulink model of the UBC Hospital interfaced with the PC-Cluster

### 4.4 Water Station

The water system is also a critical infrastructure as its failure may jeopardize the human life. In this project, the Water Station is one of the components of the reduced-scale UBC campus benchmark system, which was originally developed in Simulink [1]. The UBC water supply system gets the water from the reservoir located outside the campus – external source. The Water Station pumps the water to every building through appropriate system of pipes. In the model considered here, the water is a token and pipes are the channels as described earlier in Section 4.1.1. The Simulink implementation of the Water Station cell includes the channels and appropriate tokens. The Water Station model has been interfaced with the PC Cluster using the S-function communication block as shown in Figure 4.8. To enable parallel simulation, this model will be connected with other cell models that reside on different computers.
4.5 Power House

In the UBC campus, the Power House plays a vital role in the case of emergency as it supplies the power and oil to the Water Station and Steam Station. The Power House receives power from the Substation, and it has the oil storage facility which can be used for back up generator in case of power failure. Thus, the Power House cell has one input token and two output tokens. The output tokens are delivered to the other cells including the Water Station and Steam Station through different channels. The Simulink model of the Power House has been interfaced with the PC Cluster using two S-function communication blocks as shown in Figure 4.9. This model is then used for distributed simulation of the UBC benchmark system.
Similar to the other cells of the UBC benchmark system, the Steam Station cell also plays a major role in daily activities on campus. The Steam Station supplies steam to the whole campus including some critical facilities such as Hospital. The steam generation process requires several pieces of equipments including steam pump, boiler, hot well, air fan, and deaerator. This cell receives input tokens including the electricity and water, and then delivers output tokens which are the steam and gas. A Simulink model of the Steam Station has been developed using input-output logical tables [1]. In this thesis, the Steam Station has been interfaced with the PC Cluster using the S-function communication block as shown in Figure 4.10. This final model is then used for distributed simulation of the UBC benchmark system.
4.7 Combined Model of the UBC Benchmark System

The interconnection of the overall subsystems of the reduced-scale UBC campus benchmark is shown in Figure 4.11. As can be seen here, the Substation sends electricity token to Hospital and Power House. Similarly, Hospital receives tokens: electricity, steam, gas, and water from different cells. The Substation Cell, Hospital Cell, Water Station Cell, Power House Cell and Steam Station Cell models have been first simulated on a single computer as shown in the Figure 4.12.
Figure 4.11: Overall model of the reduced-scale UBC benchmark system
4.8 Distribution of Models to PC Cluster Nodes

The overall model of the reduced-scale UBC benchmark system has been also implemented using 5 computers in the PC-Cluster. The corresponding computer screens showing the Simulink model interfaces are depicted in Figure 4.13. The individual computers simulating the cell models in parallel are shown in Figures 4.14, 4.15, 4.16, 4.17, and 4.18, respectively. The assignment of the cell models to the cluster nodes is as follows:

Computer 1: simulates Substation as shown in Figure 4.14
Computer 2: simulates Hospital as shown in Figure 4.15
Computer 3: simulates Water Station as shown in Figure 4.16
Computer 4: simulates Power House as shown in Figure 4.17
Computer 5: simulates Steam Station as shown in Figure 4.18
The interdependencies among the cells are realized through the exchange of tokens between the subsystems, which are implemented using the communication interfaces between the corresponding cluster nodes. Each cell shown in Figures 4.14 – 4.18 has sophisticated graphical user interface (GUI) that enables simultaneous monitoring of several variables and/or tokens during the simulation as well as the user control of the dispatch. This feature is particularly useful as it allows several users to dynamically interact with the model as the simulation proceeds and displays the results and flow of the tokens.

The Substation cell in computer 1 receives power token through the transmission line channels from BC Hydro. Then, it dispatches the electricity token to the Hospital cell in computer 2 and to the Power House cell in computer 4. The Hospital cell in computer 2 receives 4 tokens: electricity, water, steam and gas from Substation in computer 1, Water Station in computer 3, and Steam Station in computer 4. Then, the Hospital cell in turn outputs the services in the form of number of beds serviced for urgent patients, short-term patients, and long-term patients. The Water Station cell in computer 3 receives water token from an external source, electricity token from the Power House cell in computer 4, and outputs water token through water pipe channel to the Hospital in computer 2 and the Steam Station in computer 5, respectively. The Power House cell in computer 4 gets input electricity token from Substation in computer 1 and dispatches electricity and oil tokens to Water Station and Steam Station in computer 3 and computer 5, respectively. Finally, the Steam Station cell in computer 5 receives electricity token from the Power House and sends output tokens gas and steam to the Hospital cell in computer 2.
Figure 4.13: Parallel simulation of 5 cells of reduced UBC test case on PC-Cluster

Figure 4.14: User interface on Computer 1 simulating the Substation Cell
Figure 4.15: User interface on Computer 2 simulating the Hospital Cell

Figure 4.16: User interface on Computer 3 simulating Water Station Cell
Figure 4.17: User interface on Computer 4 simulating Power House Cell

Figure 4.18: User interface on Computer 5 simulating Steam Station Cell
Chapter 5

PC-Cluster Simulator Studies

In this chapter, the PC-Cluster simulator is used to simulate the reduced-scale UBC benchmark system described in Chapter 4[1]. The interdependency aspect is explained in detail showing that some of the parameters can be observed on-line while the simulation is running.

For the studies presented here, the simulation time step $\Delta t$ of 1 minute was used. The results of the distributed simulation are superimposed in Figures 5.1 through 5.6 against those obtained from a single computer simulation, which validates the proposed solution methodology. The CPU times for different methods of simulation are investigated and presented. Finally, it is observed that using the proposed distributed simulation approach as implemented on the UBC PC Cluster, a remarkable increase in the overall simulation speed and reduction of the CPU time can be achieved.

5.1 Case Scenario

In order to establish the simulation speed for both the single computer implementation and the PC-Cluster implementation, the following computer study was conducted. In this study, each cell model was initialized corresponding to its normal conditions. The model was run for 1500 minutes (25 hours) of the total time. The following disturbances were simulated.

- Initially, up to 460 minutes, there was no disturbance in the system and all five cells were running in normal mode.
- Then, at $t = 461$ minutes, there was a total failure of the power supply from BC Hydro through both feeders 1 and 2. This disturbance was assumed due to some emulated disaster event.
• At \( t = 700 \) minutes, a disturbance in water supply system was simulated where the water pipe to the Hospital was broken.
• At \( t = 900 \) minutes, the power lines were fixed and the normal power was restored. However, at the same time, irregularities in steam supply to the Hospital were also created resulting in reduction of the steam production by 50%.
• At \( t = 1200 \) minutes, the water pipe to the Hospital was repaired and the water supply was restored to normal. At the same time, the steam supply to the Hospital was completely cut off to zero.
• At \( t = 1400 \) minutes, the steam production was finally restored to its nominal production capacity.

Due to the stated above sequence of disturbances, the situation at the UBC campus unfolded as follows: From \( t = 461 \) minutes to \( t = 900 \) minutes, there was no electricity supply from Substation to Hospital and Power House as shown in Figure 5.1 and Figure 5.2. Immediately following this disturbance the backup generators automatically came into service in the Hospital, Power House and Water Station. As a result of backup generators, there were no interruptions in the Hospital service for urgent, short-term and long-term patients due to the failure of power supply from substation as observed in Figure 5.5. However, the disruption in water supply from Water Station as shown in Figure 5.4, has affected the Hospital service for 30 minutes starting from the time \( t = 700 \) minutes when the water pipe was broken. After 30 minutes the backup water system came into service and normal operation of the Hospital service was restored.

However, the disturbance in the Steam Station shown in Figure 5.6 had a greater implication on the Hospital service. The long-term service was particularly badly affected due to the interruptions in steam supply. As we can observe when the steam was reduced to 50%, the long-term beds service was also reduced to half of normal capacity, while urgent- and short-term patient services were less influenced. In case of complete cut-off of the steam token, the long-term service was completely unavailable, while sufficient portion of other two types of beds were still served.
SubStation Cell Sending Power Token to Hospital Cell

Figure 5.1: Electricity token dispatched from Substation to the Hospital

SubStation Cell Sending Power Token to Power House Cell

Figure 5.2: Electricity token dispatched from Substation to the Power House
Figure 5.3: Simulation results for power token dispatch from Power House to the Water Station

Figure 5.4: Water token dispatch from Water Station to the Hospital
Figure 5.5: Simulation results for Hospital output tokens in terms of number of beds in service
5.2 Analysis of PC-Cluster Simulation

All the above scenario conditions were simulated using the PC-Cluster and the results were compared with those obtained from a single computer. The simulation results were shown in Figures 5.1 to 5.6. As can be seen in these figures, the distributed simulation results coincide exactly with those obtained from the single computer simulation. This shows the accuracy of the distributed simulation on the PC-Cluster. Thus, the PC-Cluster simulation results were validated against the benchmark system of the Simulink model simulation in a single computer.

The CPU-time for the simulation of five cells model is summarized in Table 5.1. As can be seen, a single-computer simulation took about 4.4 seconds. The distributed simulation of the five cells UBC test case with five computers of PC-Cluster was five times faster than the
original model in a single computer where message flag is used as a method of synchronized data transfer. But for the same distributed simulation test using the standard low-level SCI interrupt functions for synchronized data transfer the simulation speed was only 3.86 times faster. Thus, the PC Cluster simulation with the developed synchronization method is more efficient. This is an expected result based on the previously investigated interrupt mechanisms as explained in Chapter 3.

<table>
<thead>
<tr>
<th>Simulation Method</th>
<th>Processor Type</th>
<th>CPU-time in seconds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single Computer</td>
<td>AMD Athlon 64 Processor 4000+ 2.41 GHz</td>
<td>4.25</td>
</tr>
<tr>
<td>Distributed Simulation using 5 Computers in PC-Cluster</td>
<td>AMD Athlon 64 Processor 4000+ 2.41 GHz</td>
<td>1.1</td>
</tr>
<tr>
<td>where sci low level interrupt message is used for</td>
<td></td>
<td></td>
</tr>
<tr>
<td>synchronized data transfer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distributed Simulation using 5 Computers in PC-Cluster</td>
<td>AMD Athlon 64 Processor 4000+ 2.41 GHz</td>
<td>0.85</td>
</tr>
<tr>
<td>where message segment flag is used for synchronized</td>
<td></td>
<td></td>
</tr>
<tr>
<td>data transfer</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 5.1: CPU-time for different methods of simulation

To further investigate the benefits of distributed simulation, we have considered different number of cells. As shown in Table 5.2 and Figure 5.7, there was a little improvement in computational speed with distributed simulation in case of only two cells (Substation and Power House), compared to a single processor simulation. In case of three cells (Substation, Power House and Water Station) there was a more pronounced rise in simulation speed (2.37 times) with PC-Cluster as compared to the single computer method. Computational
efficiency was further increased for the four cells (Substation, Power House, Water Station and Steam Station) distributed simulation with four computers. Ultimately, this observation showed that for the UBC test case with five cells (Substation, Power House, Water Station, Steam Station and Hospital), the PC-Cluster simulation is about five times faster than its single-computer version. This concludes that PC-Cluster Simulator can be used as a powerful tool in investigating complex infrastructure interdependencies such as complete UBC case where more than 50 cells will be used for representing the scenarios of disaster conditions such as earth quake, terrorist attack, etc.

<table>
<thead>
<tr>
<th>Combination of different Cells</th>
<th>CPU-time for Single Computer Simulation</th>
<th>CPU-time for Distributed Simulation with the same number of computers as that of Cells</th>
<th>Improvement in simulation speed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substation + Power House</td>
<td>0.3280 seconds</td>
<td>0.3200 seconds (2-computers)</td>
<td>2.5%</td>
</tr>
<tr>
<td>Substation + Power House + Water Station</td>
<td>0.5690 seconds</td>
<td>0.2400 seconds (3-computers)</td>
<td>137%</td>
</tr>
<tr>
<td>Substation + Power House + Water Station + Steam Station</td>
<td>0.9100 seconds</td>
<td>0.2500 seconds (4-computers)</td>
<td>264%</td>
</tr>
<tr>
<td>Substation + Power House + Water Station + Steam Station + Hospital</td>
<td>4.25 seconds</td>
<td>0.85 seconds (5-computers)</td>
<td>400%</td>
</tr>
</tbody>
</table>

Table 5.2: Simulation speed for different number of cells
Figure 5.7: CPU-time for different number of cells with single computer and distributed simulation.
Chapter 6

Conclusions and future work

In this thesis, a SCI-based PC-Cluster simulator for studying the interdependent nature of critical infrastructures was developed where the computational superiority of a clustered multi processor PC-Cluster over a sequential single processor system was demonstrated. A practical scaled test case of the University of British Columbia (UBC) campus was used. The UBC model included a substation, hospital, water station, power house and steam station.

The test model was distributed among five computers of the PC-Cluster. The PC-Cluster used in this research consists of 16 computers networked via high speed SCI interconnect adapter cards. A set of C-language based communication libraries based on optimum SCI low-level functions was developed, in order to use the hardware interconnect efficiently. Furthermore, a Simulink S-function block using these communication functions was built for parallel simulation of Simulink models distributed in different computers of the PC-Cluster. With the implementation of SCI low-level interrupt function for synchronized back and forth zero-length data transfer between nodes, the measured communication overhead was 40μseconds. In case of message polling method of bi-directional synchronized zero-length data transfer the communication latency was 7.2μseconds.

The PC-Cluster based distributed simulation plot results for the UBC test case were identical as those obtained from sequential simulation on a single computer. The message polling synchronized data transfer based distributed simulation on PC-Cluster was nearly five times faster than the simulation of the original model on a single computer. On the other hand, SCI low-level interrupt function synchronized data transfer method of distributed simulation is 3.86 times faster than that obtained on a single processor. There is a negligible improvement in PC-Cluster simulation speed compared to a single node simulation in case of models with less complexity such as two cells: substation and power house. When the number of cells
increases to five, the distributed parallel computing showed a remarkable improvement in computation speed compared to that of one processor simulation. Hence PC-Cluster simulator is proposed for better computational efficiency and feasibility of investigating the interdependencies of complex and critical infrastructures.

Future research would include the implementation of the OVNI (Object Virtual Network Integrator) method and I2SIM (Infrastructure Interdependencies Simulator) on SCI-based PC-Cluster using the MATE ("Multi-Area Thévenin Equivalent") network partitioning technique [1], [2], [3]. Here one master node contains a memory segment to which all other sub-systems residing on other nodes will send Thévenin Equivalent data. Then the master node solves the reduced system represented by the links subsystem. The updated link currents are then communicated back to the sub-systems and the whole process starts again. For this, a method of sending data from different nodes to a single node with a common memory segment should be developed [3] on SCI. Additionally, this distributed PC-Cluster simulation with different Δt's (simulation time step) should be investigated in order to achieve better computational efficiency due to less communication overhead.
References

Chapter 1


Chapter 2


Chapter 3


**Chapter 4**


Chapter 5


Chapter 6


Appendix

Computer Programs

/*---------------------------------------------*/
/*
/* FILE NAME : COMPUTER1.C
/* DESCRIPTION : Measurement of Communication Latency between two
* Computers using two different methods of synchronous
* back and forth data transfer
/*
/*
*/
/*---------------------------------------------*/

#include <stdio.h>
#include <string.h>
#include <stdlib.h>
#include <sci_comm_lib.h>

void main ()
{
    int localNodeId = 4;    /*computer 1 */
    int remoteNodeId = 8;   /*computer 2 */
    int vecsize = 100;
    int h = 1;
    
    double y [100], x [100];
    int i;
    
    scistart (&h);
    sciinit (&h, localNodeId, remoteNodeId, vecsize);
    for (i=0; i<vecsize-1; i++) {
        x [i] = (double) i;
    }
    scireceive (&h, y, vecsize);
    scisend (&h, x, vecsize);
    sciend (&h);
    sciclose (&h);
}
/* FILE NAME : COMPUTER2.C */
/* DESCRIPTION : Measurement of Communication Latency between two Computers using two different methods of synchronous back and forth data transfer */

#include <stdio.h>
#include <string.h>
#include <stdlib.h>
#include <sci_coram_lib.h>

void main () {
  int localNodeId = 8; /*computer 2 */
  int remoteNodeId = 4; /*computer 1 */
  int vecsize = 100;
  int h = 1;

  double y [100], x [100];
  int i;
  scistart (&h);
  sciinit (&h, localNodeId, remoteNodeId, vecsize);
  for (i=0; i<vecsize-1; i++) {
    x [i] = (double) i;
  }
  scireceive (&h, y, vecsize);
  scisend (&h, x, vecsize);
  sciend (&h);
  sciclose (&h);
}
#define S_FUNCTION_NAME SCI_COMM_LIB
#define S_FUNCTION_LEVEL 2

#define NUM_INPUTS 2

/* Input Port 0 */
#define IN_PORT_0_NAME u0
#define INPUT_0_WIDTH 3
#define INPUT_DIMS_0_COL 1
#define INPUT_0_DTYPE real_T
#define INPUT_0_COMPLEX COMPLEX_NO
#define IN_0_FRAME_BASED FRAME_NO
#define IN_0_DIMS 1-D
#define INPUT_0_FEEDTHROUGH 1
#define IN_0_ISSIGNED 0
#define IN_0_WORDLENGTH 8
#define IN_0_FIXPOINTSCALING 1
#define IN_0_FRACTIONLENGTH 9
#define IN_0_BIAS 0
#define IN_0_SLOPE 0.125

/* Input Port 1 */
#define IN_PORT_1_NAME u1
#define INPUT_1_WIDTH 1
#define INPUT_DIMS_1_COL 1
#define INPUT_1_DTYPE real_T
#define INPUT_1_COMPLEX COMPLEX_NO
#define IN_1_FRAME_BASED FRAME_NO
#define IN_1_DIMS 1-D
#define INPUT_1_FEEDTHROUGH 1
#define IN_1_ISSIGNED 0
#define IN_1_WORDLENGTH 8
#define IN_1_FIXPOINTSCALING 1
#define IN_1_FRACTIONLENGTH 9
#define IN_1_BIAS 0
```c
#define IN_1_SLOPE 0.125

#define NUM_OUTPUTS 0

/* Output Port 0 */
#define OUT_PORT_0_NAME y0
#define OUTPUT_0_WIDTH 3
#define OUTPUT_DIMS_0_COL 1
#define OUTPUT_0_DTYPE real_T
#define OUTPUT_0_COMPLEX COMPLEX_NO
#define OUT_0_FRAME_BASED FRAME_NO
#define OUT_0_DIMS 1-D
#define OUT_0_ISSIGNED 1
#define OUT_0_WORDLENGTH 8
#define OUT_0_FIXPOINTSCALING 1
#define OUT_0_FRACTIONLENGTH 3
#define OUT_0_BIAS 0
#define OUT_0_SLOPE 0.125

#define NPARAMS 0

#define SAMPLE_TIME_0 0
#define NUM_DISC_STATES 0
#define DISC_STATES_IC [0]
#define NUM_CONT_STATES 0
#define CONT_STATES_IC [0]

#define SFUNWIZ_GENERATE_TLC 1
#define SOURCEFILES "_SFB_scilib.lib _SFB_sisci_api.lib"
#define PANELINDEX 6
#define USE_SIMSTRUCT 0
#define SHOW_COMPILE_STEPS 0
#define CREATE_DEBUG_MEXFILE 0
#define SAVE_CODE_ONLY 1
#define SFUNWIZ_REVISION 3.0
#define u_width 1
#define y_width 1

#if defined (MATLAB_MEX_FILE)
#include "tmwtypes.h"
#include "simstruc_types.h"
#else
```
#include "rtwtypes.h"
#endif

#include <stdio.h>
#include <string.h>
#include <math.h>
#include "sci_comm_lib_version.h"
#include "simstruc.h"

int localNodeId = 4; /* NodeId of Computer 1 */
int remoteNodeId = 8; /* NodeId of Computer 2 */
int h = 1;
int vecsize = 4;

static void mdlInitializeSizes (SimStruct *S)
{
    DECL_AND_INIT_DIMSINFO (inputDimsInfo);
    DECL_AND_INIT_DIMSINFO (outputDimsInfo);
    ssSetNumSFcnParams (S, NPARAMS);
    if (ssGetNumSFcnParams(S) != ssGetSFcnParamsCount(S)) {
        return; /* Parameter mismatch will be reported by Simulink */
    }

    ssSetNumContStates (S, NUM_CONT_STATES);
    ssSetNumDiscStates (S, NUM_DISC_STATES);

    if (!ssSetNumInputPorts (S, NUM_INPUTS)) return;

    /*Input Port 0*/
    ssSetInputPortWidth (S, 0, INPUT_0_WIDTH);
    ssSetInputPortDataType (S, 0, SS_DOUBLE);
    ssSetInputPortComplexSignal (S, 0, INPUT_0_COMPLEX);
    ssSetInputPortDirectFeedThrough(S,0,
        INPUT_0_FEEDTHROUGH);
    ssSetInputPortRequiredContiguous (S, 0, 1); /*direct input signal access*/
/*Input Port 1*/

ssSetInputPortWidth (S, 1, INPUT_1_WIDTH);
ssSetInputPortDataType (S, 1, SS_DOUBLE);
ssSetInputPortComplexSignal (S, 1, INPUT_1_COMPLEX);
ssSetInputPortDirectFeedThrough (S, 1,
    INPUT_1_FEEDTHR0UGH);
ssSetInputPortRequiredContiguous (S, 1, 1); /*direct
    input
    signal
    access*/

if (!ssSetNumOutputPorts (S, NUM_OUTPUTS)) return;

/*Output Port 0*/

ssSetOutputPortWidth (S, 0, OUTPUT_0_WIDTH);
ssSetOutputPortDataType (S, 0, SS_DOUBLE);
ssSetOutputPortComplexSignal (S, 0, OUTPUT_0_COMPLEX);

ssSetNumSampleTimes (S, 1);
ssSetNumRWork (S, 0);
ssSetNumIWork (S, 0);
ssSetNumPWork (S, 0);
ssSetNumModes (S, 0);
ssSetNumNonsampledZCs (S, 0);

ssSetOptions (S, (SS_OPTION_EXCEPTION_FREE_CODE |
    SS_OPTION_USE_TLC_WITH_ACCELERATOR |
    SS_OPTION_WORKS_WITH_CODE_REUSE));

}  

#define MDL_SET_INPUT_PORT_FRAME_DATA

static void mdlSetInputPortFrameData (SimStruct *S,
    int_T port,
    Frame_T frameData)
{
    ssSetInputPortFrameData (S, port, frameData);
}

static void mdlInitializeSampleTimes (SimStruct *S)
{
    ssSetSampleTime (S, 0, SAMPLE_TIME_0);
ssSetOffsetTime (S, 0, 0.0);
}

#define MDL_SET_INPUT_PORT_DATA_TYPE
static void mdlSetInputPortDataType (SimStruct *S, int port, DTypeId dType)
{
    ssSetInputPortDataType (S, 0, dType);
    ssSetInputPortDataType (S, 1, dType);
}

#define MDL_SET_OUTPUT_PORT_DATA_TYPE
static void mdlSetOutputPortDataType (SimStruct *S, int port, DTypeId dType)
{
    ssSetOutputPortDataType (S, 0, dType);
}

#define MDL_SET_DEFAULT_PORT_DATA_TYPES
static void mdlSetDefaultPortDataTypes (SimStruct *S)
{
    ssSetInputPortDataType (S, 0, SS_DOUBLE);
    ssSetInputPortDataType (S, 1, SS_DOUBLE);
    ssSetOutputPortDataType (S, 0, SS_DOUBLE);
}

#define MDL_INITIALIZE_CONDITIONS
#if defined (MDL_INITIALIZE_CONDITIONS)

static void mdlInitializeConditions (SimStruct *S)
{
}
#endif

#define MDL_START
#if defined (MDL_START)

static void mdlStart (SimStruct *S)
{
    /*SCI functions to start and initialize the SCI environment */
    scistart (&h);
    sciinit (&h, localNodeId, remoteNodeId, vecSize);
}
static void mdlOutputs (SimStruct *S, int_T tid)
{
    const real_T *u0 = (const real_T*) ssGetInputPortSignal (S,0);
    real_T     *y0  = (real_T *) ssGetOutputPortRealSignal (S,0);

double x [4], y [4];

    u0 [0] = x [0];
    u0 [1] = x [1];
    u0 [2] = x [3];

    scisend (&h, x, vecsize);
    scireceive (&h, y, vecsize);

    y0 [0] = y [0];
    y0 [1] = y [1];
    y0 [2] = y [2];
}

/* Function: mdlTerminate * Abstract:

* In this function, you should perform any actions that
* are necessary at the termination of a simulation.
* For example, if memory was allocated in mdlStart,
* this is the place to free it.
*/

static void mdlTerminate (SimStruct *S)
{
    /* SCI functions to close the network and free the
    * allocated resource and terminate from SCI environment
    */
    sciend (&h);
    sciclose ();
} 

#ifdef MATLAB_MEX_FILE /* Is this file being compiled as a MEX-file? */
#include "simulink.c" /* MEX-file interface mechanism */
#endif