$\begin{array}{c} \textbf{Parallel Scientific Computing} \\ \textbf{in } C++ \ \textbf{and } MPI \end{array}$

 $A\ seamless\ approach\ to\ parallel\ algorithms\ and\ their\ implementation$

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Preface

Scientific computing is by its very nature a practical subject - it requires tools and a lot of practice. To solve realistic problems we need not only fast algorithms but also a combination of good tools and fast computers. This is the subject of the current book, which emphasizes equally all three: algorithms, tools, and computers. Often times such concepts and tools are taught *serially* across different courses and different textbooks, and hence the interconnection between them is not immediately apparent. We believe that such a close integration is important from the outset.

The book starts with a heavy dosage of C++ and basic mathematical and computational concepts, and it ends emphasizing advanced parallel algorithms that are used in modern simulations. We have tried to make this book fun to read, to somewhat demystify the subject, and thus the style is sometimes informal and personal. It may seem that this happens at the expense of rigor, and indeed we have tried to limit notation and theorem proofing. Instead, we emphasize concepts and useful tricks-of-the-trade with many code segments, remarks, reminders, and warnings throughout the book.

The material of this book has been taught at different times to students in engineering, physics, computer science, and applied mathematics at Princeton University, Brown University, and MIT over the last 15 years. Different segments have been taught to undergraduates and graduates, to novices as well as to experts. To this end, on all three subjects covered, we start with simple introductory concepts and proceed to more advanced topics - bandwidth, we believe, is one strength of this book.

We have been involved in large-scale parallel computing for many years from benchmarking new systems to solving complex engineering problems in computational mechanics. We represent two different generations of computational science and supercomputing, and our expertise are both overlapping and complementary. The material we selected to include in this book is based on our experiences and needs as computational scientists for high-order accuracy, modular code, and domain decomposition. These are necessary ingredients for pushing the envelope in simulation science and allow one to test new theories and concepts or solve very large specific engineering problems accurately.

In addition to integrating C++ and MPI concepts and programs into the text, we also provide with this book a *software suite* containing all the functions and programs discussed. It is our belief, as stated earlier, that mastery of this subject requires both a knowledge of the tools and substantial practice using the tools. Part of the integration that we are attempting to achieve is attained when the reader is able to go immediately from the textbook to the computer to experiment with the concepts which have been presented. We envision the software suite allowing the reader to do the following: to verify the concepts presented in the book by using the programs that are provided, to extend the programs in the book to implement concepts that may have been discussed but not programmed, and to tackle different problems than those presented using the software provided.

How to Use This Book

The current book is appropriate for use by students in engineering and physics, computer science, and applied mathematics. It is designed to be more like a textbook and less of a research monograph. The material can be used to fill two semesters with the following breakdown: The first semester will cover chapters 1 to 5 at the senior undergraduate or first year graduate level. The second semester will cover the remainder of the book in a first or second year graduate course. Chapters 1 to 5 cover all the basic concepts in algorithms, C++, and MPI. Chapters 6 to 10 cover discretization of differential equations and corresponding solvers, and present more advanced C++ and MPI tools. The material in chapter 3 on approximation of functions and discrete data is fundamental and precedes other topics. In the basic material on discretization, we separated *explicit* from *implicit* approaches because the parallel computational complexity of the two is fundamentally different.

A lighter course, e.g. a quarter course or a lower level undergraduate course, could be based on chapters 1 to 5 by leaving out the MPI material and possibly other advanced topics such as wavelets, advanced quadrature rules, and systems of nonlinear equations. There are other possibilities as well. A graduate level course on numerical linear algebra can be based on sections 4.1.6, 4.1.7 and chapters 7 to 10. Assuming that the student has a C++ background or even another high performance language then the addition of MPI material in sections 2.3, 3.4, 4.3 and 5.13 to the above will constitute one full semester course on parallel numerical linear algebra. Another possibility for a quarter course is to simply teach the algorithms in chapters 5 to 8 covering traditional numerical analysis. Supplementary notes from the instructor, e.g. theorem proofs and more case studies, can make this a full semester course.

The book is designed so that it can be used with or without the C++ and MPI tools and associated concepts but we strongly encourage the instructor to teach the course as a seamless integration of both algorithms and tools.

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Chapter 1

Scientific Computing and Simulation Science

1.1 What is Simulation?

Science and engineering have undergone a major transformation at the research as well as at the development and technology level. The modern scientist and engineer spend more and more time in front of a laptop, a workstation, or a parallel supercomputer and less and less time in the physical laboratory or in the workshop. The virtual wind tunnel and the virtual biology lab are not a thing of the future, they are here! The old approach of "cut-and-try" has been replaced by "simulate-and-analyze" in several key technological areas such as aerospace applications, synthesis of new materials, design of new drugs, chip processing and microfabrication, etc. The new discipline of nanotechnology will be based primarily on large-scale computations and numerical experiments. The methods of scientific analysis and engineering design are changing continuously, affecting both our approach to the phenomena that we study as well as the range of applications that we address. While there is a lot of software available to be used as almost a "black-box," working in new application areas requires good knowledge of fundamentals and mastering of effective new tools.

In the classical scientific approach, the physical system is first simplified and set in a form that suggests what type of phenomena and processes may be important, and correspondingly what experiments are to be conducted. In the absence of any known-type governing equations, dimensional inter-dependence between physical parameters can guide laboratory experiments in identifying key parametric studies. The database produced in the laboratory is then used to construct a simplified "engineering" model which after field-test validation will be used in other research, product development, design, and possibly lead to new technological applications. This approach has been used almost invariably in every scientific discipline, i.e., engineering, physics, chemistry, biology, etc.

The simulation approach follows a parallel path but with some significant differences. *First*, the phase of the physical model analysis is more elaborate: The physical system is cast in a form governed by a set of partial differential equations, which represent continuum approximations to microscopic models. Such approximations are not possible for all systems, and sometimes the microscopic model should be used directly. *Second*, the laboratory exper-

iment is replaced by simulation, i.e., a numerical experiment based on a discrete model. Such a model may represent a discrete approximation of the continuum partial differential equations, or it may simply represent a statistical representation of the microscopic model. Finite difference approximations on a grid are examples of the first case, and Monte Carlo methods are examples of the second case. In either case, these algorithms have to be converted to software using an appropriate computer language, debugged, and run on a workstation or a parallel supercomputer. The output is usually a large number of files of a few Megabytes to hundreds of Gigabytes, being especially large for simulations of time-dependent phenomena. To be useful, this numerical database needs to be put into graphical form using various visualization tools, which may not always be suited for the particular application considered. Visualization can be especially useful during simulations where interactivity is required as the grid may be changing or the number of molecules may be increasing.

The simulation approach has already been followed by the majority of researchers across disciplines in the last few decades. The question is if this is a new science, and how one could formally obtain such skills. Moreover, does this constitute fundamental new knowledge or is it a "mechanical procedure," an ordinary skill that a chemist, a biologist or an engineer will acquire easily as part of "training on the job" without specific formal education. It seems that the time has arrived where we need to reconsider boundaries between disciplines and reformulate the education of the future *simulation scientist*, an inter-disciplinary scientist.

Let us re-examine some of the requirements following the various steps in the simulation approach. The *first task* is to select the right representation of the physical system by making consistent assumptions in order to derive the governing equations and the associated boundary conditions. The conservation laws should be satisfied; the entropy condition should not be violated; the uncertainty principle should be honored. The second task is to develop the right algorithmic procedure to discretize the continuum model or represent the dynamics of the atomistic model. The choices are many, but which algorithm is the most accurate one, or the simplest one, or the most efficient one? These algorithms do not belong to a discipline! Finite elements, first developed by the famous mathematician Courant and rediscovered by civil engineers, have found their way into every engineering discipline, physics, geology, etc. Molecular dynamics simulations are practiced by chemists, biologists, material scientists, and others. The third task is to compute efficiently in the ever-changing world of supercomputing. How efficient the computation is translates to how realistic of a problem is solved, and therefore how useful the results can be to applications. The fourth task is to assess the accuracy of the results in cases where no direct confirmation from physical experiments is possible such as in nanotechnology or in biosystems or in astrophysics, etc. Reliability of the predicted numerical answer is an important issue in the simulation approach as some of the answers may lead to new physics or false physics contained in the discrete model or induced by the algorithm but not derived from the physical problem. Finally, visualizing the simulated phenomenon, in most cases in three-dimensional space and in time, by employing proper computer graphics (a separate specialty on its own) completes the full simulation cycle. The rest of the steps followed are similar to the classical scientific approach.

In classical science we are dealing with matter and therefore *atoms* but in simulation we are dealing with information and therefore *bits*, so it is atoms versus bits! We should, therefore, recognize the simulation scientist as a separate scientist, the same way we recognized just a few decades ago the computer scientist as different than the electrical engineer or the

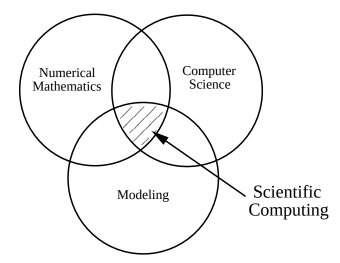


Figure 1.1: Definition of scientific computing as the intersection of numerical mathematics, computer science, and modeling.

applied mathematician. The new scientist is certainly not a computer scientist although she should be computer literate in both software and hardware. She is not a physicist although she needs a sound physics background. She is not an applied mathematician although she needs expertise of mathematical analysis and approximation theory.

With the rapid and simultaneous advances in software and computer technology, especially commodity computing, the so-called *soupercomputing*, every scientist and engineer will have on her desk an advanced simulation kit of tools consisting of a software library and multi-processor computers that will make analysis, product development, and design more optimal and cost-effective. But what the future scientists and engineers will need, first and foremost, is a solid inter-disciplinary education.

Scientific computing is the heart of simulation science, and this is the subject of this book. The emphasis is on a balance between classical and modern elements of numerical mathematics and of computer science, but we have selected the topics based on broad modeling concepts encountered in physico-chemical and biological sciences, or even economics (see figure 1.1).

1.2 A Seamless Approach Path

Our aim in writing this book has been to provide the student, the future simulation scientist, with a seamless approach to numerical algorithms, modern programming techniques, and parallel computing. Often times such concepts and tools are taught serially across different courses and different textbooks, and hence the interconnection between them is not immediately apparent. The necessity of integrating concepts and tools usually comes after such courses are concluded, e.g. during a first job or a thesis project, thus forcing the student to synthesize what is perceived to be three independent subfields into one in order to produce a solution. Although this process is undoubtly valuable, it is time consuming and in many cases it may not lead to an effective combination of concepts and tools. Moreover, from the

pedagogical point of view, the integrated seamless approach can stimulate the student simultaneously through the eyes of multiple disciplines, thus leading to enhanced understanding of subjects in scientific computing.

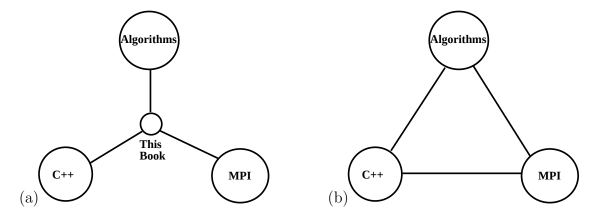


Figure 1.2: Simultaneous integration of concepts shown in (a) in contrast with the classical serial integration shown in (b).

As discussed in the previous section, in the scientific simulation approach there are several successive stages that lead from :

- 1. The real world problem to its mathematical formulation;
- 2. From the mathematical description to the computer implementation and solution, and
- 3. From the numerical solution to visualization and analysis.

In this book, we concentrate on stage (2) which includes not only the mathematics of numerical linear algebra and discretization but also the implementation of these concepts in C++ and MPI.

There are currently several excellent textbooks and monographs on these topics but without the type of integration that we propose. For example, the book by Golub & Ortega [45] introduces pedagogically all the basic parallel concepts, but a gap remains between the parallel formulation and its implementation. Similarly, the books by Trefethen & Bau [88] and Demmel [26] provide rigor and great insight into numerical linear algebra algorithms, but they do not provide sufficient material on discretization and implementation. On the other hand, popular books in C++ (e.g., by Stroustrup [86]) and MPI (e.g., by Pacheco [73]) are references that teach programming using disconnected algorithmic examples, which is useful for acquiring general programming skills but not for parallel scientific computing. Our book treats numerics, parallelism, and programming equally and simultaneously by placing the reader at a vantage point between the three areas as shown in the schematic of figure 1.2(a), and in contrast with the classical approach of connecting the three subjects serially as illustrated in figure 1.2(b).

1.3 The Concept of Programming Language

In studying computer languages, we want to study a new way of interacting with the computer. Most people are familiar with the use of software purchased from your local computer store, software ranging from word processors and spreadsheets to interactive games. But have you ever wondered how these things are created? How do you actually "write" software? Throughout this book we will be teaching through both lecture and example how to create computer software that solves scientific problems. Our purpose is not to teach you how to write computer games and the like, but the knowledge gained here can be used to devise your own software endeavors.

It has been stated by some that the computer is a pretty dumb device, in that it only understands two things — on and off. Like sending Morse code over a telegraph wire with signals of dots and dashes, the computer uses sequences of zeros and ones as its language. The zeros and ones may seem inefficient, but it is not just the data used, but the rules applied to the data which make it powerful. This concept, in theory, is no different than human language. If we were to set before you a collection of symbols, say a b c d ... z, and indicate to you that you can use these to express even the most complex thoughts and emotions of the human mind and heart, you would think we were crazy. Just 26 little symbols? How can this be? We know, however, that it is not merely the symbols that are important, but the rules used to combine the symbols. If you adhere to the rules defined by the English language, then books like this can be written using merely combinations of the 26 characters! How is this similar to the computer? The computer is a complex device for executing instructions. These instructions are articulated by using our two-base characters, 0 and 1, along with a collection of rules for combining them together. This brings us to our first axiom:

Axiom I: Computers are machines which execute instructions. If someone is not telling the computer what to do, it does nothing.

Most people have had some experience with computers, and immediately they will read this statement and say: "Hey, I have had my computer do all kinds of things that I didn't want!". Ah, but read the axiom carefully. The key to this axiom is the use of the term someone. The one thing to keep in mind is that some human, or collection of humans, developed software to tell the computer what to do. At a relatively low level, this would be the people who wrote the operating system used by the computer. At a higher level, this would be the people that developed the word processor or game that you were using. In both cases, however, someone determined how the computer would act and react to your input. We want you, after reading this book and understanding the concepts herein, to be able to be in the driver's seat. This leads us to our second axiom:

Axiom II: Computer programming languages allow humans a simplified means of giving the computer instructions.

We tell you that we want you to be in the driver's seat, and you tell me "I don't want to learn how to communicate in zeros and ones ... learning English was hard enough!" You can imagine how slowly the computer age would have progressed if every programming class consisted of the following lecture scheme. Imagine the first day of class. On the first day,

the instructor tells you that you will be learning the two basic components of the computer language today: 0 and 1. He may force you to say zero and one a few times, and then write zero and one many times on a piece of paper for practice, but then, what else would there be to learn concerning your character set? Class dismissed. Then, for the rest of the semester, you would spent your time learning how to combine zeros and ones to get the computer to do what you want. Your first assignment might be to add two numbers a and b, and to store the result in c (i.e., c = a + b). You end up with something that looks like:

Seems like a longwinded way of saying

$$c = a + b$$
?

But this is what the computer understands, so this is how we must communicate with it. However, we do not communicate in this fashion. Human language and thought use a higher abstraction than this. How can we make a bridge for this gap? We bridge this gap via programming languages, see figure 1.3.

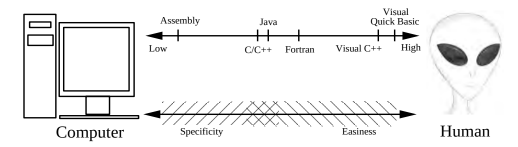


Figure 1.3: Programming languages provide us a means of bridging the gap between the computer and the human.

The first programming language we will mention is assembly. The unique property of assembly is that for each instruction, there is a one-to-one correspondence between a command in assembly and a computer understandable command (in zeros and ones). For instance, instead of writing

01011001010001000111010101000100

as a command, you could write 'load a \$1'. This tells the computer to load the contents of the memory location denoted by 'a' into register \$1 in the computer's CPU (Central Processing Unit). This is much better than before. Obviously, this sequence of commands is much easier for the human to understand. This was a good start, but assembly is still considered a "low-level language". By low-level we mean that one instruction in assembly is

equal to one computer instruction. But as we said earlier, we want to be able to think on a higher level. Hence, there was the introduction of "higher level" languages. Higher level languages are those in which one instruction in the higher-level language equals one or more computer level instructions. We want a computer language where we can say 'c = a + b', and this would be equivalent to saying:

```
load a $1
load b $2
add $1 $2 $3
save $3 c
```

One high-level instruction was equivalent to four lower level instructions (here written in pseudo-assembly so that you can follow what is going on). This is preferable for many reasons. For one thing, we as humans would like to spend our time thinking about how to solve the problem, not just trying to remember (and write) all the assembly code! Secondly, by writing in a higher-level language, we can write code that can work on multiple computers, because the translation of the higher level code can be done by a compiler into the assembly code of the processor on which we are running.

As you read through this book and do the exercises found herein, always be mindful that our goal is to utilize the computer for accomplishing scientific tasks encountered in simulation science. At a high-level, there is a science or engineering problem to solve, and we want to use the computer as a tool for solving the problem. The means by which we will use the computer is through the writing and execution of programs written using the computing language C++ and the parallel message passing libraries of MPI.

1.4 Why C++ and What is MPI?

The algorithms we present in the book can certainly be implemented in other languages, e.g. FORTRAN or Java, as well as other communication libraries, e.g. PVM (Parallel Virtual Machine). However, we commit to a specific language and parallel library in order to provide the student with the immediate ability to experiment with the concepts presented. To this end, we have chosen C++ as our programming language for a multitude of reasons: First, it provides an object-oriented infrastructure that accommodates a natural breakdown of the problem into a collection of data structures and operations on those structures. Secondly, the use of C++ transcends many disciplines beyond engineering where traditionally FORTRAN has been the prevailing language. Thirdly, C++ is a language naturally compatible with the basic algorithmic concepts of

- partitioning,
- recursive function calling,
- dynamic memory allocation, and
- encapsulation.

Similarly, we commit to MPI (Message Passing Interface) as a message passing library because it accommodates a natural and easy partitioning of the problem, it provides portability and efficiency, and it has received wide acceptance by academia and industry.

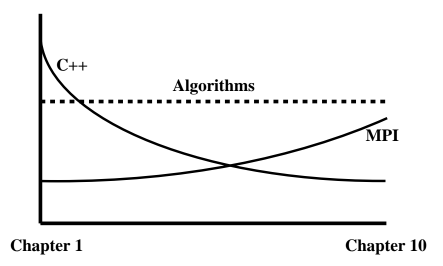


Figure 1.4: Progression of new material throughout the book in the three areas shown in figure (1.2).

The simultaneous integration we propose in this book will be accomplished by carefully presenting related concepts from all three sub-areas. Moving from one chapter to the next requires different dosages of new material in algorithms and tools. This is explained graphically in figure 1.4, which shows that while new algorithms are introduced at an approximately constant rate, the introduction of new C++ and MPI material vary inversely. We begin with an emphasis on the basics of the language, which allows the student to immediately work on the simple algorithms introduced initially, while as the book progresses and the computational complexity of algorithms increases the use of parallel constructs and libraries is emphasized.

More specifically, to help facilitate the student's immersion into object-oriented thinking, we provide a library of *classes* and *functions* for use throughout the book. The classes contained in this library are used from the very beginning of the book as a natural, user-defined, extension of C++. As the book progresses, the underlying logic and programming implementation of these classes are explained, bringing the student to a deeper understanding of the development of C++ classes. We will denote all classes used within the book and not inherent to C++ with the letters \mathbf{SC} , such as the classes SCVector and SCMatrix.



This is done to clearly distinguish between C++ defined and user-defined data types, and also to accentuate the utility of user-defined types within the C++ programming language. As students become more familiar and confident in their ability to devise and use datatypes, we encourage them to use these facilities provided by the language for more effective programming and problem solving. All the codes of this book and many more examples are included in the *software suite*, which is distributed with this book.

1.5 What About OpenMP?

Due to the recent proliferation of distributed shared-memory (DSM) machines in the scientific computing community, there is much interest in how best to appropriately utilize both the distributed and the shared-memory partitioning of these systems. MPI provides an efficient means of parallel communication among a distributed collection of machines; however, not all MPI implementations take advantage of shared-memory when it is available between processors (the basic premise being that two processors, which share common memory, can communicate with each other faster through the use of the shared medium than through other communication means).

OpenMP (<u>Open Multi Processing</u>) was introduced to provide a means of implementing shared-memory parallelism in FORTRAN and C/C++ programs. Specifically, OpenMP specifies a set of environment variables, compiler directives, and library routines to be used for shared-memory parallelization. OpenMP was specifically designed to exploit certain characteristics of shared-memory architectures such as the ability to directly access memory throughout the system with low latency and very fast shared-memory locks. To learn more about OpenMP, visit www.openmp.org.

A new parallel programming paradigm is emerging in which both the MPI and OpenMP are used for parallelization. In a distributed shared-memory architecture, OpenMP would be used for intra-node communication (i.e., between a collection of processors which share the same memory subsystem) and MPI would be used for inter-node communication (i.e., between distributed distributed collections of processors). The combination of these two parallelization methodologies may provide the most effective means of fully exploiting modern distributed shared-memory (DSM) systems.

1.6 Algorithms and Top Ten List

The Greeks and Romans invented many scientific and engineering algorithms, but it is believed that the term 'algorithm' stems from the name of the ninth-century Arab mathematician al-Khwarizmi, who wrote the book al-jabr wa'l muqabalach which eventually evolved into today's high school algebra textbooks. He was perhaps the first to stress systematic procedures for solving mathematical problems. Since then, some truly ingenious algorithms have been invented, but the algorithms that have formed the foundations of the scientific computing as a separate discipline were developed in the second part of the twentieth century. Dongarra & Sullivan put together a list of the top ten algorithms of the twentieth century [33]. According to these authors, these algorithms had the greatest influence on science and engineering in the past. They are in chronological order:

- 1. **1946:** The Monte Carlo method for modeling probabilistic phenomena.
- 2. 1947: The Simplex method for linear optimization problems.
- 3. 1950: The Krylov subspace iteration method for fast linear solvers and eigensolvers.
- 4. **1951:** The Householder matrix decomposition to express a matrix as a product of simpler matrices.

- 5. **1957:** The FORTRAN compiler that liberated scientists and engineers from programming in assembly.
- 6. **1959-1961:** The QR algorithm to compute many eigenvalues.
- 7. **1962:** The Quicksort algorithm to put things in numerical or alphabetical order fast.
- 8. **1965:** The Fast Fourier Transform to reduce operation count in Fourier series representation.
- 9. **1977:** The Integer relation detection algorithm, which is useful for bifurcations and in quantum field theory.
- 10. **1987:** The Fast multipole algorithm for N-body problems.

Although there is some debate as to the relative importance of these algorithms or the absence of other important methods in the list, e.g. finite differences and finite elements, this selection by Dongarra & Sullivan reflects some of the thrusts in scientific computing in the past. The appearance of the FORTRAN compiler, for example, represents the historic transition from assembly language to higher level languages, as discussed earlier. In fact, the first FORTRAN compiler was written in 23,500 assembly language instructions! FORTRAN has been used extensively in the past, especially in the engineering community, but most of the recent scientific computing software has been re-written in C++, e.g. the Numerical Recipes [75].

In this book we will cover in detail the algorithms (3), (4), (6) and (8) from the above list, including many more recent versions, which provide more robustness with respect to round-off errors and efficiency in the context of parallel computing. We will also present discretizations of ordinary and partial differential equations using several finite difference formulations.

Many new algorithms will probably be invented in the twenty-first century, hopefully some of them from the readers of this book! As Dongarra & Sullivan noted "This century will not be very restful for us, but is not going to be dull either!"

Chapter 2

Basic Concepts and Tools

In this chapter we introduce the main themes that we will cover in this book and provide an introduction for each of them. We begin with a brief overview of C++ and define the two basic concepts of functions and classes as well as other syntactic elements of the language. We then introduce basic mathematical concepts that include elements of linear algebra, vector orthogonalization, and corresponding codes and software. Finally, we introduce parallel programming and review some generic parallel architectures as well as standard parallel algorithms for basic operations, e.g., the fan-in algorithm for recursive doubling. We also provide a brief overview of the main MPI commands.

2.1 Introduction to C++

An ancient proverb states that the beginning of a thousand mile journey begins with a single step. For us, this single step will be a brief overview of the C++ programming language. This introduction is not designed to be all-inclusive, but rather it should provide the scaffolding from which we will build concepts throughout this book. Admittedly, what you will read now may seem daunting in its scope, but as you become more familiar with the concepts found herein, you will be able to use the C++ language as a tool for furthering your understanding of deeper mathematical and algorithmic concepts presented later in the book. With this in mind, let us begin our thousand mile journey with this first step.

Any programming language can be broken down into two high level concepts:

- Data, and
- Operations on data.

Though this may seem like a trivial statement, quite often in science and engineering problems the real work that needs to be done is identifying what is the relevant data, and what operations need to be executed on that data to obtain the desired results. From the programming point of view, we will assume that you already have a clear concept of what data is needed to solve the problem, and what algorithms will be acting on the data; we will focus on translating these needs into the programming language.

We have chosen to present this material in a top-down manner; that is, we will start from the high level of the program and work our way down toward lower and lower levels of detail. At the end of this section, we will recapitulate what we have learned, and show how all the pieces do indeed fit together though an example. We start with the idea of a program, or 'code' as it is sometimes referred to within the scientific computing community. A program is a sequence of instructions acting on a collection of data. Just as this chapter had a starting point, every program must have a starting point, and in C++, the starting point of execution is the "main" function, which is called **main**. This tells the computer where to start execution of your program. The simplest C++ code that can be written is the following:

```
int main(int argc, char ** argv){
}
```

This piece of code will compile and execute, but it will do absolutely nothing. Though it may contain data inputed through the arguments argc and argv, it contains no operations on that data. It merely provides the computer with a starting point of execution, and then immediately terminates because all executable commands have been executed (which in this case is none!).



This is your first C++ program. In keeping with programming tradition, your first non-trivial C++ program should be the following:

```
#include<iostream.h>
int main(int argc, char ** argv){
   cout << "Hello World" << endl;
}</pre>
```

At this stage, you should type in the program above, compile it using your native C++ compiler, and execute this program. The result of this program should be that the statement "Hello World" is printed to your screen. If you have problems with this exercise, see Appendix A.1.

In theory, you now have your first C++ program. You have written the code, compiled and linked the code, and are able to execute the code on your native machine. Now that this first step is behind us, let us jump into discussing some of the basic concepts, one of which we have just gained some experience, i.e., the concept of a function.

2.1.1 Two Basic Concepts in C++

There are two basic concepts used throughout the C++ programming language: the concepts of

- Function, and of
- Class.

The C programming language, upon its inception, had at least one self-defining feature: modularity. The C language was designed to be modular, and this modularity was accomplished through the employment of functions. Almost everything in C is a function, and some have said that "... all C really is a big function calling a bunch of other functions". Well, this is almost right. The C language basically consists of two components, a core language specification which contains basic data types and constructs (such as **if** statements, **for** statements, etc., some of which we discuss later on in this chapter), and a collection of libraries, each of which contains many pre-defined functions. C++ built on this philosophy, and introduced the "class" as a second fundamental building block of the language. Within C++, functions and classes are intertwined to create the desired program.

We begin by defining what we mean by a function and a class. Functions and classes can be distinguished by their fundamental premises. The primary premise of a function revolves around what the function does, whereas the fundamental premise of a class revolves around the data that the class contains. Functions are designed to be abstractions of algorithms; Classes (at least as presented in this book) are an abstraction of data and operations on that data. We will clarify this distinction by examining the two concepts in more detail.

Functions

Functions are abstractions which encapsulate a concept or algorithm. The concept of a function is probably not new to you. In mathematics, we see functions all the time. We

define functions so that we can abstract a particular operation or collection of operations into one statement. In mathematics, we note functions in a manner like

$$f(x) = x^3 - x^2 + 2.$$

We understand that if we evaluate the function at the point x = 2, denoted f(2), this is equivalent to substituting the number 2 into the expression $x^3 - x^2 + 2$, yielding $2^3 - 2^2 + 2 = 6$. We hence would say that f(2) = 6. In mathematical parlance, we would add rigor to all that we have done so far, and state this collection of operations as follows:

Given x as a real number, define f(x) as a function returning a real number, where the definition of f(x) is given by the expression $f(x) = x^3 - x^2 + 2$.

This example demonstrates the three major components of a function:

• input, output, and contract (or algorithm).

We specified the valid range of parameters that can be inputed into this function (mathematically referred to as the domain of the function); we specified the range in which the output would lie (mathematically referred to as the range of the function); and finally we specified what, given a particular input, the function will do. The same holds true for C++. For a function, we need to specify the input, output, and contract.

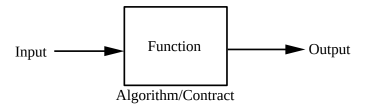


Figure 2.1: A schematic of a function in C++.

In C++, the process of specifying the input, output, and contract is done in two stages, see figure 2.1 ¹. These two stages are specifying the following for each function:

- Function declaration, and
- Function definition.

A function's declaration accomplishes several things in one step. It declares the name of the function and the data types of the input and output of the function. The function definition specifies what the function will accomplish given particular input data. Hence, the definition of a function is the algorithmic explanation of the contract of the function.

A schematic for the syntax used for the declaration of a C++ function is given in figure 2.2. Using this as our guide, let us attempt to formulate our mathematical function into a

¹According to the language standard, it is possible to combine both of these items into one statement satisfying the requirement for both simultaneously. For pedagogical clarity, we will always keep the two stages separate.

C++ function. For the purposes of this demonstration, let us assume that we have a data type called *float*, which is the floating point representation of a real number. The function declaration of our function "f" is given by:

float f(float x);

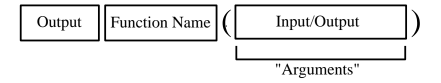


Figure 2.2: Schematic of the syntax of a C++ function.

Examining our schematic given in 2.2, we can dissect this code to understand what is going on. Let us see if we have met the three components of a function declaration. First, we specified the name of the function, "f". Next, we specified that the valid input to our function is a floating point value. Finally, we specified that our function returns a floating point value. Hence, we now have declared our function! What does declaration really mean though? Declaration of a function allows us to use this function throughout our code with the assumption that this function will act upon the contract of the function (later specified by the definition of the function). A key thing to realize is that:

• A function must be declared before it can be used.

Because of this fact, many programmers place all their function declarations at the beginning of their C++ program so that all functions are accessible everywhere. You will notice throughout this book that we either place our function declarations within a header file (the files with a .h extension) which are included at the beginning of a program, or we directly insert the function declarations after the include files and prior to the main function definition. An example template of this type of file setup is shown at the end of this section.

Another important fact is that within the *argument list* (the list of inputs and outputs given as arguments to the function), the names specified are irrelevant. The compiler is only concerned about the data type. Hence, it would be perfectly valid to write

float f(float);

Why have a variable name there, if it is just to be ignored? The most practical reason we put variable names in function declarations is that we normally cut-and-paste the function declaration from the function definition. The function definition does require variable names to be used. We will now give the function definition so that this becomes more apparent.

```
float f(float x){
   float y;

   y = x*x*x - x*x + 2;

   return y;
}
```

Notice that the function definition has a very similar beginning to the function declaration. As before, you specify the function name and the input and output data types. The difference, however, is that the function definition is the implementation of our contract. For the function definition, including specific variable names within the argument list is essential because we will use these variable names throughout the definition to refer to that data that was inputed. In C++, when data is inputed into a function, the information is passed by value. This means that when the function is called, a copy of the information is created for the function to use, not the original data. This is an important and yet subtle point about C++. We will discuss this in more detail later (see section 3.1.2). For now, the thing to remember is that the function takes from its argument list the information passed to it, and it stores a copy of the information in a variable specified by the name given in the argument list.

In this example, we declare a variable y in which we temporarily store the value of our function, given by the expression x*x*x - x*x + 2, and then we return the value of the variable y. The return statement designates the variable from which a value is to be returned from the function back to the caller. If we were to examine a code snippet, we could use our function just as we did mathematically by writing:

```
float w;
w = f(2);
```

If were to print the value of w, we would see that returned value is the floating point value 6.000. Some of this will become more clear after we have discussed basic data types. The key items to remember from this discussion are:

- Every function must have a function declaration and definition.
- Function declarations specify the name of the function and the data types of the inputs and outputs.
- Function definitions specify the implementation of the algorithm used to carry out the contract of the function.
- Variable names in function declarations do not matter.

- Variable names in function definitions do matter because they specify how the data is to be referred to in the implementation of the function.
- Variables passed to C++ functions are passed by *value* unless otherwise specified.



Recall our little main function we wrote, compiled, and ran at the beginning of this section. Let us now combine that code with our new function.

If you were to compile and run this code, you would obtain a statement on your screen that says: The value of w is: 6.00.

In the program above, we use an object named cout, the declaration of which is found in the system header file iostream.h. The object cout is used for printing to standard output, which in many cases is the screen. For now, it is sufficient to know that the << symbols delineate expressions to be printed. In the code above, the first statement to be printed is the string "The value of w is:," then the value associated with the variable w, and then the end-of-line character denoted by the term endl. We will speak more about cout later in this chapter.

Classes

Classes are abstractions which encapsulate data and operations on that data. In C++, the concept of classes is used to simplify through encapsulation very complex data structures.

A class consists of two parts: data and methods operating on the data. What you will find is that methods are merely functions which are "attached" to classes. The concept of a method is analogous to that of a function, with the primary focus of a method being to act on the data of the class, and not on arbitrary inputed data.

For example, in the Gram-Schmidt routines in section 2.2.9, we utilize several user-defined classes; among those classes is the class SCVector. Vector does not exist as a basic data type in C++; however, the language allows us to define a new type, called SCVector, which consists of a collection of basic data types and operations on that data. The declaration for the class SCVector is given below. We will not go over every detail now, but will defer explicit explanation until later in this book (see section 3.1.8). However, we call your attention to the two basic concepts of classes in C++:

- 1. encapsulated data, and
- 2. methods acting on that data.

In this class, the variables *dimension* and *data* are encapsulated by the class, and all the remaining methods (in the section marked 'public') act on this basic data.



We now present the class declaration of the SCVector class:

```
class SCVector{
private:
                     // dimension of the vector
        dimension;
  double *data; // pointer to array containing vector components
public:
  SCVector(int dim); // default constructor
  SCVector(const SCVector& v); // copy constructor
  SCVector(int col, const SCMatrix &A); //secondary constructor
  ~SCVector(); //destructor
         Dimension() const; //dimension of the vector
  int
                       // Euclidean norm of the vector
  double Length();
  void
         Normalize();
                       // normalize vector
  double Norm_l1();
  double Norm_12();
  double Norm_linf();
  //***********
  // User Defined Operators
```

```
//****************
int operator==(const SCVector& v) const;
int operator!=(const SCVector& v) const;
SCVector & operator=(const SCVector& v);

double operator()(const int i) const;
double& operator()(const int i);

void Print() const;
};
```

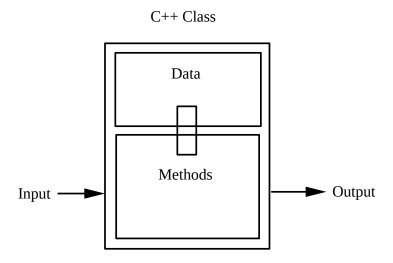


Figure 2.3: A C++ class encapsulates data and methods acting on that data.

We will explain classes more thoroughly later (section 3.1.8), but let us take this opportunity to point out a few features of classes:

- In the class above, there are several "constructors". A **constructor** is the first method which is called when an object is instantiated. These methods can be used to initialize data, set up information within the class, etc.
- A **destructor** is the method called prior to an object being deleted. The operating system will call this method (if it is available) to allow the object to "clean up for itself" prior to the operating system (OS) finishing the job by freeing the memory to which the class was allocated.
- Notice that some methods of this class modify the data contained within the object, while others merely compute things based upon the data within the object. For instance, the function *Normalize* does exactly that it normalizes the vector data contained with the object to have norm one. The function *Norm_l2*, however, does not modify the data contained with the object, but merely computes the Euclidean norm of the vector based upon the information within the object.

Type	Description
short	short integer
int	integer
long	long integer

Table 2.1: Integer data types.

• Classes allow us to define what are referred to as **overloaded operators**. In the declaration given above, we have listed these as "user defined operators". In addition to defining new data types, we can also define (or re-define) how common unary and binary operators act on those objects (such as defining what '+' means when two newly defined objects are involved).

2.1.2 Learning the Syntax and Other Basic Commands

Getting Past ";" and "{ }"

As you may have already noticed from the small amount of code that we have presented to you, the symbols ";" and "{ } " are integral to C++. We will briefly describe the purpose of these symbols here.

In general, the ";" is used to terminate an executable statement, hence why you see it at the conclusion of the commands listed above. Having such a symbol denote the end of an executable statement allows the compiler easily delineate between statements.

The { } brackets (called curly brackets) are used to denote *scope*. We will not go into all the nuances of scope right now other than to tell you that the scope of a variable or a function is the area of code in which that variable or function can be used.

Basic Data Types

In C++, variables are used to store information. In this section we go over some (not all, see the Appendix A.2 for more information) of the basic data types available in C++. Just like in mathematics, a variable is a symbol used to denote a particular value, either numerical or character. One way of thinking of a variable is that it is a box in which information can be stored. In C++, all variables must have a *type*. The *type* tells the computer what kind of box to create, the dimension of the box, the shape, etc. The syntax for creating a variable is:

Some basic data types are listed in the tables 2.1, 2.2 and 2.3. Given the convention above, we see that to declare a variable called x of type int, we would write:

int x;

Type	Description
float	single precision
double	double precision

Table 2.2: Floating point data types.

Type	Description
char	character

Table 2.3: Character data type.

This allocates a block of memory of the size of an integer, and would assign the symbol x to refer to that location in memory. Hence, from now on, if we act on the variable x, we are acting on the content of the memory assigned to x. This may sound odd at first, but this is one of the subtle differences between computer programming and mathematics. One thing that most people do not realize is that the computer does not have integer memory, floating point memory, character memory, etc 2 . As far as the computer is concerned, memory is memory. The data type that is used tells the computer how to interpret memory. A particular set of four bytes can be used in one program to represent an integer, and in another program to represent a float. The computer does not care. What is important is that the computer needs to know how to interpret the bit pattern of the four bytes. Does it interpret the collection of bits as an integer or a float, or some other variable type? Coming to grips with this notion will be very important when we start discussing the idea of addresses, the subject of pointers, etc. So, there are two key points to remember about data types. The data type you specify tells the computer two things:

- The number of bytes to use to hold the data.
- How to interpret the bit pattern specified in those bytes.

Now that you know that variables exist, and you know some of the basic types, here are some rules to remember when using variables:

- Variables must be declared before they are used.
- Variable names can be of arbitrary length.
- Variable names are case sensitive.
- Variables are to be assumed to be uninitialized. *Do not* rely on the operating system to initialize/zero values for you.

²Computers do in fact have specialized memory within the processor, called *registers*, which are either integer or float/double.

Symbol	Interpretation
+	addition
-	subtraction
*	multiplication
/	division

Table 2.4: Binary arithmetic operations.

- Variable lists may be used. If we wish to allocate three integer variables a,b and c, we may do so as follows: int a,b,c;. All three variables will be declared as integers. It is also perfectly valid to declare each one with a separate statement.
- Variables may be initialized by either constants or expressions within the declaration statement. Hence, if we wanted to initialize the integer variable a to zero, we could do the following: int a=0;. You can also do this in the context of variable lists. Suppose you want to initialize the integer variable b to one, but not the other variables. You can do this as follows: int a,b=1,c;

Basic Operations

Now that we have some concept of variables, the natural question to ask is: "What can we do with them?" C++ provides a collection of basic operations, some of which are listed in Table 2.4.

The operations presented look very much like the operations you would expect in mathematics. We must make two special notes, however. The first note is concerning the assignment operator, and the second note is concerning order of precedence and associativity. In C++, the symbol "=," which we, in English, pronounce "equals," should be interpreted as "is assigned the value of,", e.g. x=2 is to be read x "is assigned the value of" 2. Take the following C++ code example:

```
int x,y,z;
x = 2;
y = 4;
z = x + y;
```

The C++ interpretation of this code is as follows: First, declare three variables, x, y, and z as integers. Next, assign x the value 2, then assign y the value 4. We then add the values of x and y, and assign to z the newly computed value. The ordering is important. In mathematics, we may say p = q or q = p, and both statements mean that p is equal to q. However, in C++, p = q says that the variable p is assigned the same value as the variable q, whereas q = p says that the variable q is assigned the value of p. As you can see, these two statements are not equivalent.

The second item to note is that operators have both precedence and associativity. The C++ operator precedence and associativity are provided in table 2.5. What does operator precedence and associativity really mean to the programmer? Examine the following example: Assume that we want to add up six numbers: 1, 2, 3, 4, 5, 6. Mathematically, we write this operation as 1 + 2 + 3 + 4 + 5 + 6. However, if we implement this expression, keeping in mind that we are dealing with a binary operator "+," then we would write the following for summing: 1 + 2 = 3, 3 + 3 = 6, 6 + 4 = 10, 10 + 5 = 15, 15 + 6 = 21. We begin by adding the first two numbers together, and then we accumulate as we come to each new value. The same is true for the computer. When the computer is faced with the expression:

```
int x,y,z,w;

x = 2.0;
y = 3.0;
z = 4.0;

w = x + y + z;

it interprets this as being equivalent to:
   int x,y,z,w;

x = 2.0;
y = 3.0;
z = 4.0;

w = x + y;
w = w + z;
```

Notice that in the second expression, each evaluation involves only one binary expression. Hence associativity is left to right in this case. Now suppose we had the following expression:

```
int x,y,z,w;

x = 2.0;
y = 3.0;
z = 4.0;

w = x + y * z;

The computer interprets this to be the following:
int x,y,z,w;

x = 2.0;
y = 3.0;
```

```
z = 4.0;
w = y * z;
w = w + x;
```

Why multiplication prior to addition? The multiplication operator has precedence over the addition operator, and hence all multiplications are done first. This is a very important concept to realize:

Key Concept

• Order of operations is important. Precedence can make all the difference!

OPERATIONS	ASSOCIATIVITY
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	left to right
! \sim ++ + - * & (type) size of	right to left
* / %	left to right
+ -	left to right
<< >>	left to right
< <= > >=	left to right
== !=	left to right

Table 2.5: Unitary +, -, and * have higher precedence than the binary forms.

One thing we should speak to is the use of () in expressions. Notice in the precedence table that () are at the top of the list. This is for a reason. The use of () gives the programmer the right to specify the order of precedence by explicitly placing () within the expression. For instance, in the following piece of code

```
x = 2.0;
y = 3.0;
z = 4.0;
w = (x + y) * z;
the computer interpret this to be the following:
int x,y,z,w;
```

int x,y,z,w;

Symbol	Interpretation
!	NOT

Table 2.6: Unitary Boolean Operations.

```
x = 2.0;
y = 3.0;
z = 4.0;
w = x + y;
w = w * z;
```

We have in effect told the computer the precedence order that we, the programmer, want by explicitly specifying that the addition is to be done first, and then the multiplication. This brings us to a good, sound coding rule:

Key Concept

Use () to explicitly denote the order or precedence that is desired.
 () cost you nothing in terms of computational time, yet they can save you hours of debugging time trying to find an order of precedence error.

The Boolean Expression

One of the most fundamental concepts used in computer science is the concept of a boolean expression. A boolean expression returns a value of either true or false. In C++, true and false are valid values of the enumerated (variable) type bool. For now, we will merely concern ourselves with the fact that true may be converted to the integer value '1' and false may be converted to the integer value '0'. As you will see in three fundamental structures presented below, boolean expressions are used to determine the flow of control. Flow of control is, in layman's terms, which C++ statements should be executed in a particular situation. Both the unary and binary boolean operators are presented in tables 2.6 and 2.7.



There are several key facts to know about these operators. First, they are binary operators just like + and - (addition and subtraction, respectively). Thus, you can assign a variable the value obtained by using them. For example, the following code is perfectly legitimate:

Symbol	Interpretation
&&	AND
	OR
>	greater than
>=	greater than or equal to
<	less than
<=	less than or equal to
==	equal to

Table 2.7: Binary boolean operations.

```
#include <iostream.h>
int main(int argc, char ** argv){
   int a,b,c;

   a = 3;
   b = 5;

   c = a < b;

   cout << "The value of c = " << c << endl;
}</pre>
```

If we were to print the value of c immediately following the assignment statement (as we have in the code above through the use of the *cout* statement), we would find that the value of c is 1, because it is true that the number 3 < 5. The ability to assign a variable the value of a boolean expression holds true for all boolean binary operators.

Two other operators that may not be as familiar to most readers are the boolean AND (&&), and boolean OR (\parallel). These two operators are commonly used to simplify logical expressions so that several cases can be considered in one statement. The boolean values of these two expressions are given below:

$\mathrm{OR}(\parallel)$			Al	$\mathrm{AND}(\&\&)$		
	0	1		0	1	
0	0	1	0	0	0	
1	1	1	1	0	1	

The two above tables should be interpreted in the following manner. Suppose we have two variables a and b. The boolean value of variable a is denoted by the values on the left of the table, and the boolean value of the variable b is denoted by the values on the top of the table. If we were to execute the operation a operator> b (where operator> is either OR or AND), then the result of this operation is given by the value in the square given by the

respective row and column given by the values of a and b. Hence, if a = 1 and b = 0, a | | b yields the value 1 (true) while a&&b yields the value 0 (false). These logical relationships are very important and hence should be memorized.



You may be wondering what happens if you use these boolean operations on regular values. Let us suppose that you had the following piece of code. What happens to the variable 'c'?

```
#include <iostream.h>
int main(int argc, char ** argv){
   int a,b,c;

   a = 3;
   b = 5;

   c = a && b;

   cout << "The value of c = " << c << endl;
}</pre>
```

Under such circumstances, a and b are implicitly cast to boolean expressions before the AND operation is performed. Implicit casting does not affect the actual value of a or b, but merely internally converts these values so that the computer can do the appropriate boolean operation that you have requested. In such cases the following implicit casting rule is applied:

• Any number not equal to zero (either positive or negative) denotes true (the logical value 1), and zero denotes false (the logical value 0).

If we now try to answer the question of what will the computer do, we see that the computer will first implicitly cast a and b to their appropriate values, which in this case are both logical true since both are non-zero, and then the AND operation will be carried out. Looking at the table above, we see that (true && true) equals true, and hence the value of c is true (cast to the value 1).

An Example: The Collatz Problem

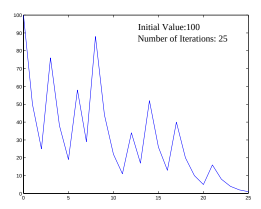
We will now proceed to explain three fundamental flow of control structures in C++: the **if** statement, the **while** statement, and the **for** statement. We will motivate our discussion of these three constructs with the following problem, known as the *Collatz Problem*. The problem itself is given by a very simple algorithm:

• Start with any integer greater than zero; if the number is even, divide it by two, otherwise multiply it by three and add one to it. Iterate this process until the number you reach is the number one.

Hence, if you start with the value 10, the sequence of numbers what you will obtain from this algorithm is the sequence 10,5,16,8,4,2,1. In Figure 2.4 we plot the iterate value versus iteration for two different initial guesses, 100 and 1000. The description of the algorithm is quite simple, and the patterns given by plotting the iterated solution versus iteration number are intriguing, however the problem that has stumped mathematicians for decades is the following proposition attributed to Collatz:

• Given any integer greater than one, the algorithm described above will terminate (i.e., the value will reach one) in a finite number of iterations.

Since the algorithm is fairly simple to implement, many explicit numerical tests have been done which demonstrate that for extremely large numbers this proposition holds true. However, at the time of this writing no theoretical proof exists for Collatz's proposition.



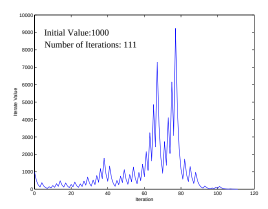


Figure 2.4: Iteration history of the Collatz algorithm for two different initial guesses, n=100 and n=1000.

We will present two pieces of code which implement the algorithm described above. The first algorithm, denoted Collatz-A, makes the assumption that Collatz is right; the other, Collatz-B, is a little less confident that the proposition holds true!



• Algorithm Collatz-A

#include<iostream.h>

```
xn = initial_guess;
i = 0;

while(xn != 1){
   cout << i << " " << xn << endl;

   if(xn%2==0) // use of integer modulus operator
        xn = xn/2;
   else
        xn = 3*xn+1;

   i=i+1;
}</pre>
```

• Algorithm Collatz-B

```
#include <iostream.h>
int main(int argc, char ** argv){
  int i,xn;
  int max_iterations = 1000; // declaration and initialization
  int initial_guess = 100;  // declaration and initialization
  xn = initial_guess;
  for(i=0;i<max_iterations;i=i+1){</pre>
    cout << i << " " << xn << endl;
    if(xn == 1)
                 // use of break statement to exit for loop
      break;
                 // when the condition xn==1 is true
    if(xn\%2==0)
      xn = xn/2;
    else
      xn = 3*xn+1;
  }
}
```

The fundamental difference between these two implementations is the choice of whether to use the *while* statement or the *for* statement. In the Collatz-A code, the *while* statement will terminate only if our iterate value reaches one; if it never reaches one, our program will run forever (or until the system administrator kills your process thinking you accidently left some mindless process running). Collatz-B, however, places a limit on the maximum number of iterations that you will allow before giving up on Collatz. Both algorithms utilize the *if* statement, so we will focus there first, and then move on to the description of both the *for* and *while* statements.

The IF Statement

The *if* statement is truly at the core of any programming language. Almost all computer architectures implement a form of the *if* statement as part of their fundamental set of computer instructions. This is done because when someone sets up a computer problem, the programmer is quite often confronted with algorithmic decisions in which if something is true, then a particular piece of code should be done, and if it is not true, then another piece of code should be done. Such logical decisions are not unique to programming, but are fundamental to the way in which we, as humans, think. For this particular algorithm, the "if" decision is based on whether the iterate value is odd or even. The way in which we implemented this was as follows:

```
if(xn%2==0)
    xn = xn/2;
else
    xn = 3*xn+1;
```

Here, we check if the xn modulus 2 is zero (i.e., if the remainder from dividing xn by 2 is zero); if it is, this implies that xn is even, and thus we should divide xn by two. Otherwise, xn is odd, and thus we should multiply it by three and add one. From this you can see the basic structure of the if statement. The basic structure is expressed in the following two logical examples: if A then B, and if A then B else C.

Several examples are provided below so that you can see the syntactic structure of these statements. There are a few things to point out: First, notice that if no {} brackets are used, only the next statement immediately following the **if** statement is executed as part of the **if**. This is a common programming mistake – people think that their **if** statement encompasses many statements, but because they forgot to put {} brackets, only the first statement is executed as part of the **if**. The other thing to notice is that you are not limited to only one statement in the conditional. You can have logical statements such as if A then B,C,D,E,... . Just remember, you need to use the {} brackets to denote this collection of commands!

Examples:

```
if( boolean statement )
    statement 1;
```

```
if( boolean statement )
   statement 1;
else
   statement 2;
if( boolean statement ){
   statement 1;
   statement 2;
}
if( boolean statement ){
   statement 1;
   statement 2;
}
else{
   statement 3;
   statement 4;
}
if( boolean statement ){
   statement 1;
   statement 2;
}else if{
   statement 3;
   statement 4;
}
```

The WHILE Statement

The while statement is commonly used for executing a set of instructions while something is true. For Collatz-A, in which we presume that Collatz was right, we continue to iterate until our iterate xn reaches the value of one.

Two example *while* statements are presented below. Note that the rules concerning {} discussed for *if* statements hold true here also. Multiple statements can be executed as part of the *while* if {} are used to denote the extent of the statements.

Examples:

```
while( boolean expression )
    statement 1;
while( boolean expression ){
```

```
statement 1;
statement 2;
...
}
```

The FOR Statement

For scientific programming in general, one of the most common statements used is the *for* loop. For loops are used to denote a finite number of times a particular set of instructions is to be executed. In Collatz-B, we only allow finite number of iterations to occur, in this case 1000. If the iterate value has not reached the value one after 1000 iterations, the loop terminates. The *for* statement used for Collatz-B is given below:

```
for(i=0;i<max_iterations;i=i+1){
    // ... statements ...
}</pre>
```

We begin by initializing a variable i to the value zero. The second part of the statement states that as long as i is less than $max_iterations$, the statements contained within the $\{\}$ should be executed. The third component of the statement says to increment the value of i by one (i.e., i = i + 1) at the conclusion of each iteration. Hence, in block form, the expression becomes:

Hence if $max_iterations$ is equal to zero, the loop will never execute. For any integer value of $max_iterations$ greater than zero, the statements inside the for statement will execute $max_iteration$ times.

There are many variations of the *for* statement, all of which follow the concept illustrated in the example above. We list here several instanciations of the *for* statement.

Examples:

```
for( statement 1; boolean expression; statement 2)
   statement 3;
```

```
for( statement 1; boolean expression; statement 2){
   statement 3;
   statement 4;
}
```

In the place of statement 1, we normally have the initialization of the looping variable. Then, we have a boolean expression which tells us when the loop is to end. Then finally, we have an increment statement which updates the looping variable. Although in this case we have used solely integers, the *for* statement does not require it. If we were marching across an evenly spaced grid on [0,1] with grid spacing h, we could equally have used the following *for* loop:

2.1.3 Learning to Print

The printing routines to standard output (i.e., the screen) are handled by the class *cout*. The declaration of *cout* can be found in the *iostream.h* header file. You will notice that we included it in every program we have written so far which has used output to the screen. This file must be included in any program using the *cout* statement.

Recall our Collatz example described above. In that example, we wanted to print the value of i at each iteration, and the value of xn for that iteration, so we used the following statement:

```
cout << i << " " << xn << endl;
```

The printing to standard output is initiated by cout. Following the cout class, the symbols '<<' are used to delineate between items to print. In this case, we want to print the value of i, followed by a space (here denoted by the string "") followed by the value of xn. The symbol 'endl' denotes printing the end of line character. If we wanted to print the final iteration count only, we could execute the following statement after the while or for statement:

```
cout << "The final iteration number is: " << i << endl;</pre>
```

The general rule for the *cout* statement is that strings (denoted by quotation marks) are printed literally; that is, exactly what is placed between quotes is printed. When variables are placed in the *cout* statement, it is tacitly assumed that the contents of the variable are to be printed, and not the variable name itself.

We now present some general remarks concerning the *cout* statement:

- The << used with *cout* is actually an *overloaded* operator. The class *cout* encapsulates into an object the information necessary to accomplish printing to standard output. The << operator is defined for all pre-defined variable types (int, float, double, etc.) and allows us to "feed' *cout* with the data that we wish to have printed to the screen.
- Instead of using 'endl', you can also use the end of line character '\n' called newline. For example, instead of

```
cout << "The final answer is yes" << endl;
you could equal use the following:
cout << "The final answer is yes\n";</pre>
```

Then ' \n' character is considered one character, and hence can be stored in a single variable of type char.

2.1.4 Learning to Read

Printing to the screen is important, but what about inputing information into the computer? To read from standard input (i.e., the keyboard) we use the object *cin*. The declaration of *cin* can be found in the *iostream.h* header file. This file must be included in any program using the *cin* statement.

The *cin* statement allows us to read data from the keyboard. In the Collatz programs that we presented earlier, every time that the user wanted to produce the pattern for a different number, the user would have to change the value of the initial guess variable and then recompile the program. Suppose in our Collatz problem that we wanted the user to input the initial value from the keyboard so that the program could be compiled just once and use the inputed information each time it ran. To accomplish this, we can use the *cin* statement as follows:

```
#include <iostream.h>
int main(int argc, char ** argv){
  int i,xn;
  int max_iterations = 1000; // declaration and initialization
```

```
int initial_guess = 100;  // declaration and initialization

cout << "Input a new value: ";

cin >> initial_guess;

xn = initial_guess;

for(i=0;i<max_iterations;i=i+1){
    // .... remainder of the program given previously</pre>
```

In the code above, we query the user for a new value (using *cout* to print a message to the screen), and then we use *cin* to retrieve the user's keyboard input and place it into the variable *initial_guess*. Using the *cin* statement as above, we can now compile this program once and rerun it allowing the user to input new values for which to obtain a Collatz sequence.

We now present some general remarks concerning the *cin* statement:

- Just as << was on overloaded operator for *cout*, >> is an overloaded operator for *cin*. The operator >> is used to delineate between inputed items.
- *cin* ignores white space when reading from standard input. Suppose that we want to input two integers into our program; we write the following:

```
int A,B;
cout << "Input two integers:";
cin >> A >> B;
```

The user could then type two integers separated by any number of white spaces (which cin would ignore), and cin would put the two integers into the variables A and B.

• *cin* reads successively through the inputed stream of information. For example, suppose that you wanted to input an integer and a floating point number into your program. We write the following:

```
int A;
float B;

cout << "Input two integers:";
cin >> A >> B;
```

If the user inputs an integer and a float separated by white space, all will work as expected. If the user, however, accidentally enters the floating point number first, the program will read up to the decimal point to obtain the integer (suppose 10.3 is

entered, cin will read the value 10 into the variable A) and then will continue from that point to read the floating point value (which for this case cin will read the value 0.3 into the variable B).

For a more comprehensive description of the *cin* operator, we suggest that the reader consult [86].

2.1.5 How to Program in Style

Though C++ is quite tolerant of programming styles, humans are not. Someone can write a perfectly legitimate C++ code (syntactically and semantically), and yet it would be virtually incomprehensible to anyone else but the programmer (and in some cases, it may even baffle the programmer if she has not looked at it for a while). In general, it is good practice to follow a few basic stylistic rules when programming:

- Comment the non-obvious. Here, non-obvious refers both to algorithmic tricks used and programming tricks used.
- Space lines of code appropriately. White space and blank lines are ignored by the compiler, hence you should use them to your advantage. A properly spaced out code is much easier to read than a compacted, mangled piece of code.
- Use indentation to denote scope. When programming for, if, while, etc., you should use indentation to denote scope. For example, examine the differences between the two codes:

```
for(i=0;i<N;i=i+1){
  y[i] = 0.0;
  for(j=0;j<N;j=j+1){
  y[i] = y[i] + A[i][j]*x[j];
}

versus

for(i=0;i<N;i=i+1){
   y[i] = 0.0;
   for(j=0;j<N;j=j+1){
      y[i] = y[i] + A[i][j]*x[j];
   }
}</pre>
```

As you can see, the indentation of the statements leads to immediate recognition of the nesting of the statements. This type of 'spot-check' ability is very important when searching for either algorithmic or syntactic bugs.

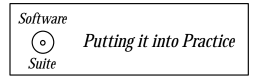
How To Comment Code

C++ provides two means of commenting code: the single line comment and the block comment. The single line comment is denoted by //. When the compiler reaches the // statement, it ignores all characters which follow after up to the line return. Thus, we can use this comment symbol to comment our coding example above as follows:

The second means of commenting is by using the /**/ syntax. The C++ compiler will ignore everything between /* and */, even if this spans across several lines. Hence, if we wanted to comment out the loops from above, we could do the following:

```
/*
for(i=0;i<N;i=i+1){
   y[i] = 0.0;
   for(j=0;j<N;j=j+1){
      y[i] = y[i] + A[i][j]*x[j];
   }
} */</pre>
```

By placing these symbols before and after this block of code, we have commented out the entire block of code. Obviously, this is not the only (nor the primary) use of comment blocks. The purpose of this form of commenting is to allow the user to place more detailed descriptions of algorithmic components of the code, which may take up more than one or two single lines.



We now introduce a full C++ program using the previously discussed coding examples. The program below contains all the essential items necessary to compile and execute our Collatz Algorithm (in this particular case, we are looking at *Collatz-A*).

```
#include <iostream.h>
#include <iomanip.h>
int main(int argc, char ** argv){
  int i,xn;
  int initial_guess = 100;
  xn = initial_guess;
  i = 0;
  while(xn != 1){
    cout << i << " " << xn << endl;
    if(xn == 1)
      break;
    if(xn\%2==0)
      xn = xn/2;
    else
      xn = 3*xn+1;
    i=i+1;
  }
}
```

The general format of our code is as follows: At the top of the file we place our "include" statements. These statements are called "pre-compiled directives." This means that they are instructions to be carried out before the compilation is done. In this case, the "#include" statement tells the compiler to insert the variable and function declarations found in the header file "iostream.h" and "iomanip.h". Since we have no other functions than our *main* function, we have no other functions to declare. Our program begins with a *main* function, which tells the computer where to start executing. Inside this function we place the C++ description of the algorithm we want executed.

To recapitulate, the general format of our C++ codes will be:

```
/***************************
/* Include Statements */
/*********************
#include <iostream.h> //Input/Output Header File
#include <iomanip.h> //Input/Output Manipulation Header File
#include <fstream.h> //File Input/Output Header File
#include <string.h> //String Manipulation Header File
#include <math.h> //Math Library Header File
// .... etc ....
```

```
/* User-Defined Variable Declarations
// Items such as Class declarations, etc.
Function Declarations
/*********************************
// User-defined function declarations.
/*******************************
       Main Program
/***********************************
int main(int argc, char ** argv){
 // ... Algorithm ...
}
/************/
     Function Definitions
/*********************************
```

Quite often throughout this book we will omit repeating this basic structure; we will focus merely on providing algorithm and function definitions. Algorithms described throughout this book can be inserted in the C++ programming shell above, compiled, and executed. When deviations from the above style are needed, they will be explicitly mentioned in the text.

2.2 Mathematical and Computational Concepts

2.2.1 Notation

We will denote a vector by a bold letter, so the transpose of a vector \mathbf{x} of length n is $\mathbf{x}^T = (x_1, x_2, x_3, \dots, x_n)$. We will denote a matrix of size $m \times n$ by a capital bold letter, say matrix \mathbf{A} , which has entries $(a_{ij}; i = 1, \dots, m; j = 1, \dots, n)$. We will often write the matrix \mathbf{A} in terms of its columns \mathbf{a}_i , each one having m entries. We will also use the symbol $\mathcal{O}(n^p)$ (read as "order-of" n^p) to denote either asymptotic computational complexity or convergence rate.

2.2.2 Binary Numbers and Round-off

Appreciation of the finite arithmetic in scientific computing is very important and sloppy handling of arithmetic precision often leads to erroneous results or even disasters in large scientific computations. It has been reported, for example, that the Patriot missile failure in Dharan, Saudi Arabia, on February 25, 1991 which resulted in 28 deaths, is ultimately attributable to poor handling of rounding errors. Similarly, the explosion of an Ariane 5 rocket just after lift-off on its maiden voyage off French Guinea, on June 4, 1996, was ultimately the consequence of simple overflow (the conversion from a 64-bit floating point value to a 16-bit signed integer value).

While we are familiar and more comfortable with the base 10 arithmetic system, a computer is restricted to a binary numbering system. The number 126, for example, has the representation

$$126 = 1 \times 10^2 + 2 \times 10^1 + 6 \times 10^0$$

in the base-10 system, or equivalently

$$011111110_2 = 0 \times 2^7 + 1 \times 2^6 + 1 \times 2^5 + 1 \times 2^4 + 1 \times 2^3 + 1 \times 2^2 + 1 \times 2^1 + 0 \times 2^0$$

in the base-2 system. This is the **floatic point** representation.

In computing we call each place in a binary number a digit or a **bit**, and we call a group of 8 bits a **byte**. Similarly, we call 1,024 bytes a Kilo-byte (1 KB) and 1,048,576 bytes a Megabyte (1 MB), and so on. An equivalent way to write the number 126 in scientific notation is:

$$+ .126 \times 10^{3}$$

sign fraction exponent

Therefore, in the computer we need to store the sign, the fraction and the exponent separately. To this end, there is a standard notation adopted by IEEE (Institute of Electrical and Electronic Engineers) for binary arithmetic, which is used in most computers (the old Cray computers did not follow this convention). There are two types of floating point numbers, depending on the number of binary digits (bits) we store: Specifically, in the **single precision** (float type in C++) we have 8 bits for the exponent and 23 bits in the fraction

whereas in the **double precision** (double type in C++) we have 11 bits for the exponent and 52 bits for the fraction. In both cases we need to also reserve one bit for the sign. What this means simply is that there is a lower and an upper bound on the size of numbers we can deal with in the computer. In the single precision type this range extends from 2^{-126} to 2^{128} and in the double precision from 2^{-1022} to 2^{1024} , so clearly the latter allows great flexibility in dealing with very small or very large numbers. The lower limit in this range determines an underflow while the upper limit determines an overflow. What value a variable takes on when it overflows/underflows depends both on the variable type and the computing architecture on which you are running. Even this large range, however, may not be sufficient in applications, and one may need to extend it by using the so-called **double extended precision** (long double in C++) which can store up to total of 128 bits. In practice, it is more efficient to use adaptive arithmetic only when it is needed, for example in refining the mesh down to very small length scales to resolve small vortices in a flow simulation.

The finite arithmetic in computing implies that the *effective zero* in the computer is about 6×10^{-8} for single precision and 10^{-16} for double precision. We can determine the value of machine epsilon by finding the value of $\frac{1}{2P}$ such that to the computer:

$$1.0 + \frac{1}{2p} = 1.0.$$



This is accomplished by increasing the value of p incrementally, and monitoring the point at which the computer cannot distinguish between the value 1 and the value $1 + \frac{1}{2^p}$. This procedure is implemented for both floating point and double precision variables in the following two functions:

```
float FloatMachineEps(){
   float fmachine_e, ftest;
   fmachine_e = 1.0;

   ftest = 1.0 + fmachine_e;
   while(1.0 != ftest){
      fmachine_e = fmachine_e/2.0;
      ftest = 1.0 + fmachine_e;
   }

   return fmachine_e;
}

double DoubleMachineEps(){
   double dmachine_e, dtest;
   dmachine_e = 1.0;
```

```
dtest = 1.0 + dmachine_e;
while(1.0 != dtest){
   dmachine_e = dmachine_e/2.0;
   dtest = 1.0 + dmachine_e;
}
return dmachine_e;
}
```

Now, a natural question is "How do I use these functions?" For starters, we would write the following program which uses both functions:

```
#include <iostream.h>

float FloatMachineEps();
double DoubleMachineEps();
int main(int * argc, char ** argv[]){
   float fep;
   double dep;

   fep = FloatMachineEps();
   dep = DoubleMachineEps();
   cout << "Machine epsilon for single precision is: " << fep << endl;;
   cout << "Machine epsilon for double precision is: " << dep << endl;
}</pre>
```

The machine zero values obtained by running the program above on a Pentium-4 processor are given in table 2.8.

Key Concept

- Notice the structure of this code:
 - 1. Function declarations
 - 2. "main" Function
 - 3. Function definitions

Variable Type	Machine Zero
float	5.96046e-08
double	1.11022e-16

Table 2.8: Machine zero for *float* and *double* precision for a Pentium-4 processor.

This code example demonstrates two important concepts. First, it demonstrates that in computing, it is important to understand how arithmetic works on your machine. Secondly, this example demonstrates that with very little programming, a user can investigate the machine upon which she is running. We must be mindful that no computer can accomplish infinite precision arithmetic; it is limited to finite precision. Finite precision arithmetic is explained as follows: when the exact value of a basic operation, e.g. addition of two numbers, is not represented with a sufficient number of digits, it is then approximated with the closest floating point number. The approximation error incurred is referred to as the *round-off* error. It is for this reason that such a fundamental property of addition as the associative property is not always satisfied in the computer. For example

$$-1.0 + (1.0 + \epsilon) \neq (-1.0 + 1.0) + \epsilon$$

as on the left-hand-side a very small number is added to a large number and that change may not be represented exactly (due to round-off) in the computer.

2.2.3 Condition Number

The condition number is a very useful measure of the sensitivity of the numerical solution to a slight change in the input. This number is proportional to the magnitude of the first derivative of the solution. This can be formally shown by considering a solution $\phi(x)$ and re-computing it for a slightly perturbed input, i.e., $x + \delta x$, where δx is the perturbation. Using the Taylor series expansion and keeping the first term only, we obtain

$$\phi(x + \delta x) \approx \phi(x) + \phi'(x)\delta x.$$

Thus, the change in the function values is

$$\frac{|\phi(x+\delta x) - \phi(x)|}{|\phi(x)|} \approx \frac{|\phi'(x)||x|}{|\phi(x)|} \times \frac{|\delta x|}{|x|}$$

Using the above equation we define as *condition number* the first term in the product of the right-hand-side. It represents the relative error in the solution (response) given a small change in the input expressed by the independent variable x.

2.2.4 Vector and Matrix Norms

We define the most important norms that we will use to measure errors in this book. We need norms both for vectors as well as for matrices. The norm of a vector $\mathbf{x}^T = (x_1, x_2, x_3, \dots, x_n)$ of length n is a scalar which obeys the following rules:

- $\parallel \mathbf{x} \parallel \geq 0$.
- $\parallel \mathbf{x} \parallel = 0 \Leftrightarrow \mathbf{x} = 0$.
- $\| \alpha \mathbf{x} \| = |\alpha| \| \mathbf{x} \|$, where α is a scalar.
- $\bullet \parallel \mathbf{x} + \mathbf{y} \parallel \leq \parallel \mathbf{x} \parallel + \parallel \mathbf{y} \parallel.$

Some of the most commonly used norms are:

- Discrete L_{∞} norm defined as: $\|\mathbf{x}\|_{\infty} = \max_{i} |x_{i}|$.
- Discrete L_2 norm defined as: $\|\mathbf{x}\|_2 = \left(\sum_{i=1}^n x_i^2\right)^{\frac{1}{2}}$.
- Discrete L_1 norm defined as: $\|\mathbf{x}\|_1 = \sum_{i=1}^n |x_i|$.
- L_p norm defined as: $\|\mathbf{x}\|_p = (\sum_{i=1}^n |x_i|^p)^{1/p}$.

There is a theorem of equivalence of vector norms, and by this we mean that: $\|\mathbf{x}\|_p$ is equivalent to $\|\mathbf{x}\|_q$ if there exist numbers A(p,q,n) and B(p,q,n) so that

$$\|\mathbf{x}\|_{p} \leq A \|\mathbf{x}\|_{q}$$

 $\|\mathbf{x}\|_{q} \leq B \|\mathbf{x}\|_{p}$.

Theorem: All L_p norms are equivalent for $p \ge 1$ and the size of the vector is n finite.

Remark: For p > q > 1, then $||x||_p \le ||x||_q \le n ||x||_p$, where n is the vector length.

The above theorem is the reason why sometimes we omit the sub-index in the norm notation. As an example, the L_1 norm and the L_2 norms are equivalent but not equal. In practice, this means that either one can be used to measure errors and the convergence result we obtain in our analysis would not depend on the particular norm.

The Cauchy-Schwarz Inequality is very useful in numerical analysis. For any two vectors \mathbf{x} and \mathbf{y} we have:

$$|(x,y)| \le ||x||_2 ||y||_2$$

where (\mathbf{x}, \mathbf{y}) is the inner product of the two vectors (see section 2.2.7) defined as $(x, y) = \sum_{i=1}^{n} x_i y_i$.

Matrix Norms: The matrix norm generated by the vector norm $||x||_p$ is defined by

$$\parallel \mathbf{A} \parallel_p = \max_{x \neq \mathbf{0}} \frac{\parallel \mathbf{A} \mathbf{x} \parallel_p}{\parallel \mathbf{x} \parallel_p}.$$

Similarly:

• The L_{∞} norm generates $\|\mathbf{A}\|_{\infty} = \max_{i} \sum_{j=1}^{n} |a_{ij}|$, which is the maximum row sum.

- The L_1 norm generates $\|\mathbf{A}\|_1 = \max_j \sum_{i=1}^n |a_{ij}|$, which is the maximum column sum.
- The L_2 generates the average matrix norm, i.e., $\|\mathbf{A}\|_2 = \sqrt{\max \lambda(\mathbf{A}^*\mathbf{A})}$, where $\mathbf{A}^*\mathbf{A}$ is a positive symmetric matrix with positive real eigenvalues. Here \mathbf{A}^* denotes the complex conjugate matrix of \mathbf{A} , which in the case of real matrix entries it is the transpose of \mathbf{A} , i.e., $\mathbf{A}^* = \mathbf{A}^T$.

There are also two useful inequalities involving matrix norms:

$$\parallel \mathbf{A} + \mathbf{B} \parallel \leq \parallel \mathbf{A} \parallel + \parallel \mathbf{B} \parallel$$

 $\parallel \mathbf{A} \mathbf{x} \parallel \leq \parallel \mathbf{A} \parallel \parallel \mathbf{x} \parallel$.

In the above inequalities we assume that a specific norm, as defined above, is chosen.

2.2.5 Eigenvalues and Eigenvectors

The eigenvalues and corresponding eigenvectors of a matrix **A** are determined by solving

$$\mathbf{A}\mathbf{x} = \lambda\mathbf{x} \Rightarrow (\mathbf{A} - \lambda\mathbf{I})\mathbf{x} = 0,$$

where

$$\mathbf{I} = \begin{bmatrix} 1 & & & & \\ & 1 & & & O \\ & & 1 & & \\ & & \ddots & & \\ O & & & 1 \end{bmatrix}$$

is the identity matrix. The eigenvectors \mathbf{x} are non-zero if the determinant is zero, i.e.,

$$\det(\mathbf{A} - \lambda \mathbf{I}) = 0.$$

and this equation determines the eigenvalues λ . It can be re-written as

$$\det(\mathbf{A} - \lambda \mathbf{I}) = \prod_{i=1}^{n} (a_{ii} - \lambda_i) + p_{n-1}(\lambda) = 0,$$

where a_{ii} are the diagonal elements of **A** and $p_{n-1}(\lambda)$ is a $(n-1)^{th}$ -order polynomial. Therefore, an $n \times n$ matrix has exactly n eigenvalues, i.e., the roots of the above n^{th} -order polynomial, which may be real or complex and simple or multiple. However, the above approach is rarely used in practice to compute the eigenvalues because it is computationally very expensive and also unstable. This is illustrated with the *Wilkinson matrix*

$$\mathbf{W} = \begin{bmatrix} 1 & & & \mathcal{O}(\epsilon) \\ & 2 & & & \\ & & 3 & & \\ & & & \ddots & \\ & & & 19 & \\ \mathcal{O}(\epsilon) & & & 20 \end{bmatrix}$$

where by $\mathcal{O}(\epsilon)$ we denote possible round-off error. In the absence of round-off the eigenvalues are determined by

$$(1 - \lambda)(2 - \lambda)(3 - \lambda)\dots(19 - \lambda)(20 - \lambda) = 0,$$

and thus: $\lambda_i = i$. However, for $\epsilon \neq 0$ the *characteristic* polynomial is

$$\lambda^{20} - 210\lambda^{19} + \ldots + 20! + \epsilon\lambda^{19}$$
.

Let us assume that $\epsilon = 10^{-11}$, then we obtain

$$\lambda_i = 1, 2, \dots, 8, 9, 10.01, 11.3, 12.5 \pm 0.5i, 14.5 \pm 0.5i, \dots, 20.$$

Therefore, the presence of even slight noise in the data results in several eigenvalues being wrong, even complex in this case!

Similarity Transformation

Next, we provide some basic background on linear algebra. First, we define the *similarity transformation*. Specifically, we say that the matrix $\bf A$ is *similar* to matrix $\bf B$ if $\bf A$ and $\bf B$ have the same eigenvalues (i.e., the same eigenspectrum) but not necessarily the same eigenvectors. Therefore, the transformation

$$A \longrightarrow PAP^{-1}$$

where \mathbf{P} is a non-singular matrix, leads to a matrix $\mathbf{B} = \mathbf{P}\mathbf{A}\mathbf{P}^{-1}$ which is similar to \mathbf{A} . This can be proved based on the definitions. Let us assume

$$Ax = \lambda x$$

and

$$\mathbf{PAP}^{-1}\mathbf{y} = \mu\mathbf{y}.$$

Then

$$\mathbf{A}\mathbf{P}^{-1}\mathbf{y} = \mathbf{P}^{-1}\mu\mathbf{y} = \mu\mathbf{P}^{-1}\mathbf{y}$$

and by defining $\mathbf{x} = \mathbf{P}^{-1}\mathbf{y}$, we have $\mu = \lambda$, so all n eigenvalues are the same for both \mathbf{A} and $\mathbf{P}\mathbf{A}\mathbf{P}^{-1}$. We also note that if \mathbf{P} is an orthogonal matrix then $\mathbf{P}^{-1} = \mathbf{P}^{T}$ and then $\mathbf{P}\mathbf{A}\mathbf{P}^{T}$ is similar to \mathbf{A} . In theory, any non-singular matrix \mathbf{P} can be used in the similarity transformation, but the best choice in practice is to use an orthonormal matrix. The reason is that with finite arithmetic ill-conditioned matrices amplify the round-off error and may lead to erroneous results unlike orthonormal matrices.

Remark 1: The transpose matrix \mathbf{A}^T is similar to matrix \mathbf{A} since they have the same characteristic polynomial. However, they do not have the same eigenvectors. In contrast, the inverse matrix \mathbf{A}^{-1} has the same eigenvectors with \mathbf{A} but inverse eigenvalues, λ_i^{-1} . This is true because

$$\mathbf{A}\mathbf{x} = \lambda \mathbf{x} \Rightarrow \mathbf{x} = \mathbf{A}^{-1}\lambda \mathbf{x} \Rightarrow \lambda^{-1}\mathbf{x} = \mathbf{A}^{-1}\mathbf{x}.$$

Remark 2: The matrix \mathbf{A}^k , where k is a positive integer, has eigenvalues λ^k , where λ are the eigenvalues of \mathbf{A} . However, \mathbf{A}^k and \mathbf{A} have the same eigenvectors. This can be extended further and it is easy to show that if we construct the *polynomial matrix*

$$p(\mathbf{A}) \equiv \alpha_0 \mathbf{A}^0 + \alpha_1 \mathbf{A} + \alpha_2 A^2 + \ldots + \alpha_k \mathbf{A}^k,$$

then

$$p(\lambda_1), p(\lambda_2), p(\lambda_3) \dots p(\lambda_n)$$

are the eigenvalues of $p(\mathbf{A})$. Correspondingly, the eigenvectors of \mathbf{A} are also eigenvectors of $p(\mathbf{A})$. As an example, the eigenvalues of $p_1(\mathbf{A}) = \mathbf{A} + \sigma \mathbf{I}$ are $(\lambda_i + \sigma)$.

We have already seen that computing the eigenvalues accurately from the determinant may not always be possible, although the Newton-Raphson method of chapter 4 is an accurate method of computing the roots of polynomials, but it may be inefficient. In chapter 10 we present several methods to compute iteratively and selectively the maximum and minimum eigenvalues and corresponding eigenvectors.

2.2.6 Memory Management

Before we present **BLAS** (Basic Linear Algebra Subroutines) in the next section, we will give you some preliminary information concerning memory management which will help to make the next discussion more relevant.

In this section, we will discuss two issues:

- Memory layout for matrices, and
- cache blocking.

Our discussion will be directed toward understanding how memory layout in main memory and cache (see figure 2.11) affect performance.

Memory Layout for Matrices

Computer memory consists of a linearly addressable space as illustrated in figure 2.5. In this illustration, we denote memory as a one-dimensional partitioned strip. To the right of the strip, the labels addr denote the address of the parcel of memory. By linearly addressable we mean that addr2 = addr1 + addrset where addrset is the memory offset between two contiguous blocks addressable memory.³ Single variables and one-dimensional arrays fit quite nicely into this concept since a single integer variable needs only one parcel of memory, and the array needs only one parcel of memory per element of the array.

How are two-dimensional arrays stored in memory? Since memory is linearly addressable, in order to store a two-dimensional array we must decide how to decompose the matrix into one-dimensional units. The two obvious means of doing this is decomposing the matrix into

³We have remained general because different architectures allow different addressable sets. Some architectures are bit addressable, some byte addressable, and some only word addressable. We will not delve further into this matter, but the interested reader should consult a computer architecture book for more details.

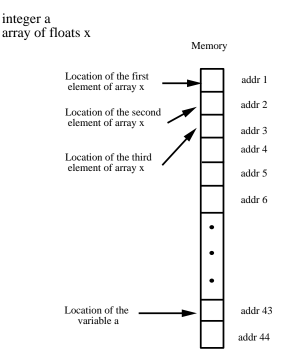


Figure 2.5: Schematic showing the memory layout for an integer variable and an array of floating point values. The partitioned strip denotes the memory of the computer, and the labels addr to the right denote the address of the parcel of memory.

a collection of rows or decomposing the matrix into a collection of columns. The first is referred to as "row-major order", and the later is referred to as "column-major order". This concept is illustrated in figure 2.6.

After examining figure 2.6, we draw your attention to the following statements:

- The amount of memory used by both ordering is the same. Nine units of memory are used in both cases.
- The linear ordering is different. Although both orderings start with the same entry (S_{00}) , the next addressable block in memory contains a different entry of \mathbf{S} (S_{01} for row-major order versus S_{10} for column-major order). This is an important observation which we will discuss further in just a moment.
- If **S** is a symmetric matrix (meaning $S_{ij} = S_{ji}$), then row-major ordering and column-major ordering appear identical.
- C++ uses row-major ordering while FORTRAN uses column-major ordering. In section 5.1.4 we discuss how multi-dimensional arrays are allocated in C++, and how row-major ordering comes into play.

In addition to the comments above, one other item needs to be mentioned before we can conclude this discussion. As shown in figure 2.11, most modern computing architectures have several layers of memory between what we have referred to as "main memory" and the central processing unit (CPU). Main memory is normally much slower than the CPU,

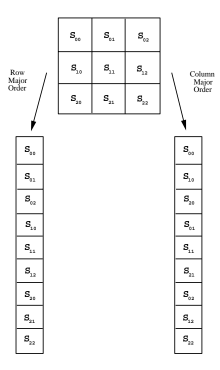


Figure 2.6: The 3×3 matrix **S** is decomposed by row-major ordering on the left and by column-major ordering on the right.

and hence engineers decided to insert smaller but faster memory between main memory and the CPU (in this discussion, we will lump register memory in with the CPU and will refer to Level 1 (L_1) and Level 2 (L_2) cache simply as "cache"). If all the items necessary to accomplish a computation could fit into cache, then the total time to execute the instructions would be reduced because the time cost of *loads* and *stores* to memory would be accelerated.

In general, even for programs in which not all the data can fit into cache, the use of cache decreases the total time cost by using cached data as much as possible. This is referred to as **cache reuse**. The goal is that once some piece of information has been loaded into cache, it should be reused as much as possible (since access to it is much faster than accessing the same piece of data from main memory). Whenever a piece of needed information is found in cache, we have what is referred to as a **cache hit**. Whenever a piece of information cannot be found in cache, and hence it must be obtained from main memory (by loading it into cache), we have what is referred to as a **cache miss**.

Items from main memory are loaded in blocks of size equal to the size of a **cache line**, a term which comes from the size of the lines going from main memory modules to the cache memory modules. Hence, if you want to access one element of an array, and it is not in cache (a miss), an entire cache line's worth of information will be loaded into cache, which may include many contiguous elements of the array. We draw your attention to figure 2.7. In this illustration, we are examining the memory layout during a matrix-vector multiplication. The 3×3 matrix **A** is stored in row-major order followed by a 3×1 vector **x**. In this simplified example, our cache consists of nine units which are loaded/stored in blocks of three units (our cache line is of size three units).

At macro time t_0 , the first element of **A** is needed, so an entire cache line's worth of

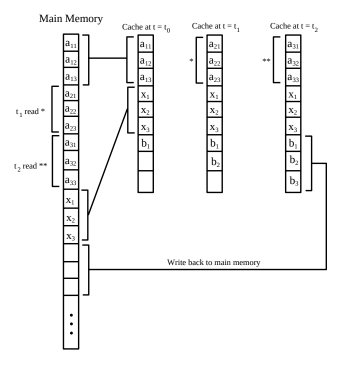


Figure 2.7: Main memory and cache layout for a matrix-vector multiplication.

information is loaded, which in this case consists of three units. Hence, the entire first row of \mathbf{A} is loaded into cache. The first element of the vector \mathbf{x} is needed, and hence an entire cache line's worth of information is loaded, which is equal to all of \mathbf{x} . To accomplish the dot product of the first row of \mathbf{A} with the vector \mathbf{x} , the other entries already in cache are needed. Thus, we have several cache hits as other computations are accomplished in order to obtain the end result of the dot product, b_1 . At macro time t_1 , the first item in the second row of \mathbf{A} is needed, and hence there is a cache miss. An entire cache line of information is loaded, which in this case is the entire second row. Again, many computations can be accomplished until another cache miss happens, in which case new information must be loaded to cache.

Although this is a very simplied example, it demonstrates how memory layout can be very important to *cache reuse*. What would have happened if we had stored the matrix \mathbf{A} in column-major order, but had implemented the algorithm above? Because of the particular cache size and cache line size, we would have had a cache miss almost every time we needed an entry of \mathbf{A} , and hence the total time cost would have been greatly increased because of the excessive direct use of main memory instead of the fast cache.

Cache Blocking

The concept of *cache blocking* is to structure the data and operations on that data so that maximum cache reuse can be achieved (maximize cache hits). In the example above, we have achieved this by choosing to store the matrix \mathbf{A} in row-major order for the particular algorithm which we implemented.

Keep this concept in mind during the discussion of BLAS in the next section. In several cases, different implementations are given for the same mathematical operation, some de-

pending on whether your matrix is stored in column-major order or in row-major order. In the end, the computational time used to run your program can be significantly altered by paying attention to which algorithm is most efficient with which memory layout. Taking into account cache characteristics when determining the data partition used is very important, and can greatly enhance or deteriorate the performance of your program.

2.2.7 Basic Linear Algebra - BLAS

Basic operations with vectors and matrices dominate scientific computing programs, and thus to achieve high efficiency and clean computer programs an effort has been made in the last few decades to standardize and optimize such operations. Two of the most basic operations with vectors which appear in a code repeatedly are **inner** (dot) **product** and **outer product**. The inner or dot product returns a scalar, however, the outer product returns a matrix. For example:

Inner Product

$$\begin{bmatrix} a_1 a_2 a_3 \dots \end{bmatrix} \begin{bmatrix} b_1 \\ b_2 \\ b_3 \\ \vdots \end{bmatrix} = c,$$

where c is the scalar value of the inner product, and c = 0 implies that the two vectors are orthogonal, a terminology similar to vector calculus.

Outer Product

$$\begin{bmatrix} a_1 \\ a_2 \\ a_3 \\ \vdots \end{bmatrix} [b_1 \, b_2 \, b_3 \dots] = \begin{bmatrix} a_1 b_1 & a_1 b_2 & a_1 b_3 \dots \\ a_2 b_1 & a_2 b_2 & a_2 b_3 \dots \\ a_3 b_1 & a_3 b_2 & a_3 b_3 \dots \\ \vdots & & & \end{bmatrix}.$$

In addition to the inner and outer products, there are several other standard matrix and vector operations that are used repeatedly in scientific computing. A convenient taxonomy is based on their computational complexity, i.e., the number of floating point operations required to complete an operation. In the previous example with the inner product of vectors with length n, we require n multiplications and (n-1) additions or a total of approximately 2n operations. We denote the computational complexity of the *inner product* as $\mathcal{O}(n)$ (read as "order n"). Similarly, we can estimate the computational complexity of the *outer product* to be $\mathcal{O}(n^2)$. We can then define levels of operations as

$$\mathcal{O}(n)$$
, $\mathcal{O}(n^2)$, and $\mathcal{O}(n^3)$,

and this is exactly what has been done with the **BLAS**, a collection of routines that perform specific vector and matrix operations. BLAS stands for **B**asic **L**inear **A**lgebra **S**ubprograms and were first proposed by Lawson et al. [67] and further developed in [27, 29]. BLAS serve as building blocks in many computer codes, and their performance on a certain computer usually reflects the performance of that code on the same computer. This is why most of the

computer vendors optimize BLAS for specific architectures. BLAS provide both efficiency and modularity. The most recent trend has been to develop a new generation of "self-tuning" BLAS libraries targeting the rich but complex memory systems of modern processors.

The adaptive BLAS software is called **ATLAS**, which stands for **A**utomatically **T**uned **L**inear **A**lgebra **S**oftware [93]. ATLAS is an implementation of empirical optimization procedures that allow for many different ways of performing a kernel operation with corresponding timers to determine which approach is best for a particular platform. More specifically, ATLAS uses two techniques: *Multiple implementation* and *code generation*. In the latter, a highly parameterized code is written that generates many different kernel implementations. In the former, different hand-written versions of the same kernel are explicitly provided. The ATLAS framework also allows the user to supply machine-specific implementations for other operations, for example *prefetch* instructions and even different timers in order to affect optimization.

Level "1"

The first level, **BLAS1**, include $\mathcal{O}(n)$ operations, and include scalar-vector multiplication, vector addition, inner (dot) product, vector multiply, and the so-called "saxpy" operation. The latter simply means "scalar alpha x plus y" and serves as a mnemonic rule for adding a vector to another vector multiplied with a scalar.

$$c = dot(\mathbf{x}, \mathbf{y})$$

 $c = c + x(i)y(i); i = 1, n$

and also

$$\mathbf{z} = saxpy(\alpha, \mathbf{x}, \mathbf{y})$$

 $z(i) = \alpha x(i) + y(i); \quad i = 1, n$

In the above expressions the equal sign implies "assignment" rather than equality as is common in computer programming.

A typical performance of the double precision (ddot) inner product on the Intel Pentium-4 (1.7GHz) (see figure 2.12) is shown in figure 2.8. Specifically, the ATLAS version of BLAS is employed with two different options of handling the data: "hot" implies that the vector is in cache and "cold" that is out of cache. This processor has two levels of cache, which are both on the chip (see figure 2.12). The primary (L_1) cache is 16 Kbytes while the secondary (L_2) cache is 256 Kbytes.⁴ We see that the "hot/hot" combination results in the best performance, which, however, is less than half of the maximum possible speed of 1.7 Gflops for a single operation per cycle. On the other hand, when the vector sizes exceed the cache size, at array size approximately 10,000, the performance asymptotes to approximately 250 Mflops, which is only a fraction of the maximum speed.

Level "2"

⁴This is relatively small cashe. The new processors have cache of several MB.

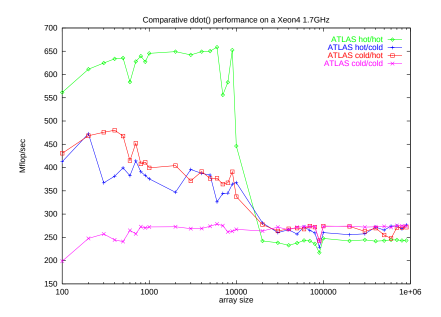


Figure 2.8: Performance of the dot product on the Intel Pentium-4 with speed 1.7 GHz. (Courtesy of C. Evangelinos)

The second level, **BLAS2**, include $\mathcal{O}(n^2)$ operations, and specifically the most important operation in scientific computing, i.e., *matrix-vector* multiplication. ⁵ There are two ways of performing this operation depending on how we access the matrix **A**, either by row or by column.

• $\mathbf{z} = \text{MatVec.}ij(\mathbf{A}, \mathbf{x})$. This is the row version.

Initialize
$$z(i) = 0.0 \text{ for } i = 1, n$$
 Begin Loop
$$i = 1, n$$
 Begin Loop
$$j = 1, n$$

$$z(i) = z(i) + A(i, j)x(j)$$
 End Loop End Loop

• $\mathbf{z} = \text{MatVec.} ji(\mathbf{A}, \mathbf{x})$. This is the *saxpy* version. We give an example of this version as it may be less familiar to the reader compared to the inner product version.

$$\left[\begin{array}{cc} 1 & 2 \\ 3 & 4 \end{array}\right] \left[\begin{array}{c} 5 \\ 6 \end{array}\right] = 5 \left[\begin{array}{c} 1 \\ 3 \end{array}\right] + 6 \left[\begin{array}{c} 2 \\ 4 \end{array}\right].$$

The saxpy version is basically a linear combination of the columns of the matrix \mathbf{A} with weights given by the entries of the vector. We will also refer to this version as the FORTRAN programming version since the matrix \mathbf{A} is accessed by columns, which is the way that matrices are stored in FORTRAN language. The loop that implements

 $^{^{5}}$ The matrix-vector BLAS routine has the name dgemv (double precision) but also the name mxv is used.

the column version is shown below. It is different than the row version only in the order of the loops (i then j instead of the j then i presented below)..

Initialize
$$z(i) = 0.0$$
 for $i = 1, n$
Begin Loop $j = 1, n$
Begin Loop $i = 1, n$
 $z(i) = z(i) + A(i, j)x(j)$
End Loop
End Loop

An operation that involves matrix-vector products, which is the generalization of saxpy, is the Gaxpy:

• Gaxpy (General **A x** plus **y**).

$$\mathbf{z} = \mathbf{y} + \mathbf{A}\mathbf{x}; \quad \mathbf{z} = gaxpy \quad (\mathbf{A}, \quad \mathbf{x}, \quad \mathbf{y})$$

$$\downarrow \quad \downarrow \quad \downarrow$$

$$m \times n \quad n \quad m$$

To compute a gaxpy we use the saxpy operation in the following loop.

$$\mathbf{z}=\mathbf{y}$$
Begin Loop
$$j=1,n$$

$$z=z+x(j)\underbrace{A(.,j)}_{\text{column}}$$
 End Loop

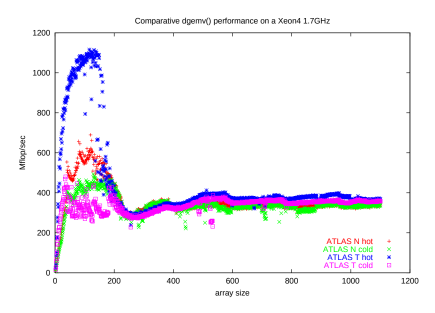


Figure 2.9: Performance of the matrix-vector multiply on the Intel Pentium-4 with speed of 1.7 GHz. (Courtesy of C. Evangelinos)

A typical performance of the double precision (dgemv) matrix-vector operation on the Intel Pentium-4 (1.7GHz) is shown in figure 2.9. Specifically, the ATLAS version of the BLAS2 routine is employed with different ways of handling the matrix, accessing it either with unit stride ("T" stands for transpose) or with larger stride ("N" stands for normal). Similarly to the dot operation in figure 2.8, hot and cold refer to in-cache or out-of-cache operations, respectively. Here we see that the maximum achievable speed is larger than the BLAS1 level operation of figure 2.8, but again in the above matrices with a rank of 200 the out-of-cache operations result in inferior performance.

Level "3"

In this level we have the *matrix-matrix* multiplication, which is an $\mathcal{O}(N^3)$ operation. It can be accomplished in six different ways, based on the basic operations of lower computational complexity of level 1 and level 2. ⁶ These six different loop arrangements were first proposed by Dongarra, Gustavson and Karp [31]. The corresponding BLAS are **BLAS3**. We present here the inner (dot) product version, the middle product or *saxpy* version, and the outer product version. The other three ways are similar to the three versions that we present here and form the so-called "dual" algorithms. In all cases the basic loop is

$$c_{ij} = c_{ij} + a_{ik}b_{kj}$$

for the matrix multiplication $\mathbf{C} = \mathbf{A} \mathbf{B}$, so it is the order of indexing which is different. To make ideas more clear we present a specific example below with a 2×2 matrix-matrix multiplication.

• Inner (dot) product version:

$$\begin{bmatrix} 1 & 2 \\ 3 & 4 \end{bmatrix} \begin{bmatrix} 5 & 6 \\ 7 & 8 \end{bmatrix} = \begin{bmatrix} 1 \cdot 5 + 2 \cdot 7 & 1 \cdot 6 + 2 \cdot 8 \\ 3 \cdot 5 + 4 \cdot 7 & 3 \cdot 6 + 4 \cdot 8 \end{bmatrix}$$

• Middle (saxpy) product version:

$$\begin{bmatrix} 1 & 2 \\ 3 & 4 \end{bmatrix} \begin{bmatrix} 5 & 6 \\ 7 & 8 \end{bmatrix} = \begin{bmatrix} 5 \begin{bmatrix} 1 \\ 3 \end{bmatrix} + 7 \begin{bmatrix} 2 \\ 4 \end{bmatrix} \quad 6 \begin{bmatrix} 1 \\ 3 \end{bmatrix} + 8 \begin{bmatrix} 2 \\ 4 \end{bmatrix} \end{bmatrix}$$

• Outer product version:

$$\left[\begin{array}{cc} 1 & 2 \\ 3 & 4 \end{array}\right] \left[\begin{array}{cc} 5 & 6 \\ 7 & 8 \end{array}\right] = \left[\begin{array}{c} 1 \\ 3 \end{array}\right] \left[\begin{array}{cc} 5 & 6 \end{array}\right] + \left[\begin{array}{c} 2 \\ 4 \end{array}\right] \left[\begin{array}{cc} 7 & 8 \end{array}\right].$$

Although mathematically equivalent, the different versions can have very different levels of computer performance because of the different ways that they access memory.

Next we write the general loops for the three versions presented above for a matrix $A(m \times r)$; $B(r \times n)$; C = AB. The asymptotic operation count for this operation is $\mathcal{O}(2mnr)$ flops.

 $^{^6{}m The~matrix\text{-}matrix}$ BLAS routine has the name dgemm (double precision) but also the name mxm is used.

• Inner (Dot) product version or **MatMat.ijk** Algorithm. Matrix **A** is accessed by rows and matrix **B** by columns, while the matrix **C** is constructed row-by-row:

```
Initialize C(i,j) = 0.0 \ for \ i = 1, m; j = 1, n Begin Loop i = 1, m Begin Loop j = 1, n Begin Loop k = 1, r C(i,j) = C(i,j) + A(i,k)B(k,j) End Loop C(i,j) = C(i,j) + A(i,k)B(k,j) End Loop C(i,j) = C(i,j) + A(i,k)B(k,j)
```

• Dual inner (Dot) product version or **MatMat.jik** Algorithm. Matrix **A** is accessed by rows and matrix **B** by columns, while the matrix **C** is constructed column-by-column (note that the ordering of the *i* and *j* loops has changed):

```
Initialize C(i,j)=0.0\, \text{for}\, i=1,m; j=1,n Begin Loop j=1,n Begin Loop i=1,m Begin Loop k=1,r C(i,j)=C(i,j)+A(i,k)B(k,j) End Loop End Loop End Loop
```

• Middle (*Gaxpy*) product version or **MatMat.jki** Algorithm. Matrix **A** and matrix **B** are both stored by columns, so we have repeated matrix-vector multiplies of matrix **A** with columns of **B**, and this is done using the *linear combination* version of BLAS2:

```
Initialize C(i,j) = 0.0 \text{ for } i = 1, m; j = 1, n Begin Loop j = 1, n Begin Loop k = 1, r Begin Loop i = 1, m C(i,j) = C(i,j) + A(i,k)B(k,j) End Loop C(i,j) = C(i,j) + A(i,k)B(k,j) End Loop C(i,j) = C(i,j) + A(i,k)B(k,j)
```

Notice that the above operation can be accomplished by using the *gaxpy* operation as follows

$$C(.,j) = gaxpy(A, B(.,j), C(.,j))$$

The dual operation of the outer product is MatMat.ikj, and it is implemented with a similar loop as above where we need to exchange i and j.

• Outer product version or **MatMat.kji** version Algorithm. Here matrix **A** is accessed by columns and matrix **B** is accessed by rows in order to form outer products.

Begin Loop
$$k=1,r$$
 Begin Loop
$$j=1,n$$
 Begin Loop
$$i=1,m$$

$$C(i,j)=\underbrace{C(i,j)+A(i,k)B(k,j)}_{saxpy}$$
 End Loop End Loop End Loop

The dual operation for the outer product version is MatMat.kij, and it is implemented with a similar loop as above where we need to exchange i for j.

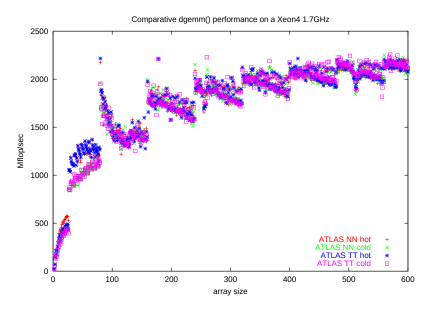


Figure 2.10: Performance of the matrix-matrix multiply on the Intel Pentium-4 with speed 1.7 GHz. (Courtesy of C. Evangelinos)

A typical performance of the double precision (dgemm) matrix-matrix operation on the Intel Pentium-4 (1.7GHz) is shown in figure 2.10. Specifically, the ATLAS version of the BLAS3 routine is employed with different ways of handling the matrix, accessing it either T or N, hot or cold, as before. Here ATLAS employs cache blocking by carefully partitioning the matrices into blocks and making effective reuse of the data. This results in a performance which is approximately the same for all different ways of accessing the data. If no such fine tuning were performed, the asymptotic performance would drop dramatically for matrices of rank above 100. It is interesting to note here that the asymptotic performance is well above 1.7 Gflops that corresponds to a single operation per cycle. This is because one of the two available floating point units of Pentium-4 (the SSE2) is capable of executing any pairwise operation of eight available registers every cycle for a maximum performance of four single or two double precision flops per cycle at IEEE 754 precision.

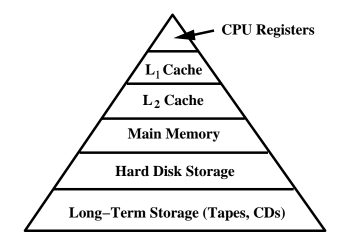


Figure 2.11: Memory hierarchies in typical computer.

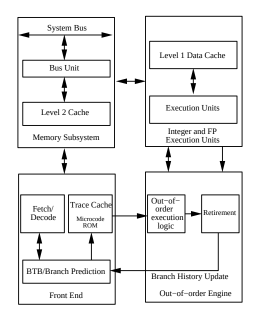


Figure 2.12: Block diagram of the Intel Pentium-4 .

BLAS and Memory Access

The practical difference in the various ways of implementing matrix-vector and matrix-matrix operations lies in the way we access memory and the type of memory, i.e., main memory or cache memory, as the latter is typically *ten times* faster than the former. To appreciate this difference we sketch in figure 2.11 a pyramid of hierarchies (see [30] for more details) in today's typical computer that shows the relative distance between the central processing unit (CPU) and the various levels of memory. The larger the size of memory, the longer it takes to access it. A layout of the Intel Pentium-4 is shown in figure 2.12.

The cost associated with the BLAS programs can be computed by taking into account the total number of floating operations but also including the cost associated with *memory* access to load the operands. The following time estimate (T) is a conservative one, as it assumes that there is no overlap between computation and loading of data:

$$T = n_f \times \delta t + n_m \times \tau = n_f \times \delta t \left(1 + \frac{n_m}{n_f} \times \frac{\tau}{\delta t}\right),$$

where n_f is the number of floating point operations, n_m is the number of memory references, and δt and τ are the times to execute a floating point operation and the time to load an operand, respectively, and both are fixed for a given computer model.

From the above equation, we can see that the ratio of n_m/n_f plays an important role in minimizing the total time. For each BLAS routine we can estimate this ratio for the three levels of operation. For example:

- For BLAS1 such as the saxpy operation, we have that $n_f = 2n, n_m = 3n + 1$ and thus $n_m/n_f \to 3/2$ for large n.
- For BLAS2 (matrix-vector multiply) we have that $n_f = 2n^2$, $n_m = n^2 + 3n$ and thus $n_m/n_f \to \frac{1}{2}$ for large n.
- Finally, for BLAS3 (matrix-matrix multiply) we have that $n_f = 2n^4$, $n_m = 4n^3$ and thus $n_m/n_f \to 2/n$ for large n.

From the three levels of operations, it is clear that the matrix-matrix multiply is the most efficient one as the number of memory references per flop decreases for larger size matrices. This is the reason that the asymptotic performance measured in the figures 2.8, 2.9 and 2.10 is maximum for the matrix-matrix operation. It is, therefore, a good practice in writing code to involve more matrix-matrix multiplications and use BLAS2 or BLAS1 less often, if possible.

Remark 1: There are also several other routines in all levels of BLAS for different operations. For example, generation of plane rotations in BLAS1, solvers of triangular equations in BLAS2, multiplications of a general matrix by a triangular matrix in BLAS3, etc. The reader can find all this information on the web using the URL for the *netlib* freeware libraries:

http://www.netlib.org/blas and also http://www.netlib.org/atlas

Remark 2: BLAS accommodate both single and double precision corresponding to prefixes s and d, respectively, in the names of subroutines. For a subset of routines (dot products and matrix-vector products) extended-precision is also available for very-high accuracy computations (prefix e). BLAS also accommodate complex arithmetic.

2.2.8 Exploiting the Structure of Sparse Matrices

The asymptotic limit of computational complexity in obtaining

$$\mathbf{C}(m \times n) = \mathbf{A}(m \times r)\mathbf{B}(r \times n)$$

is $\mathcal{O}(2nmr)$, but this limit overpredicts the count of operations for small size matrices. In practice, we often encounter sparse matrices, and thus we have to be able to obtain a better estimate for the operation count taking into account that sparsity. One such example is the multiplication of two upper triangular square matrices **A** and **B** $(n \times n)$, which contain a_{ij} and b_{ij} as entries. The resulting matrix **C** is also upper triangular, as is shown below for the case of n = 3.

$$C = \begin{bmatrix} a_{11}b_{11} & a_{11}b_{12} + a_{12}b_{22} & a_{11}b_{13} + a_{12}b_{23} + a_{13}b_{33} \\ 0 & a_{22}b_{22} & a_{22}b_{23} + a_{23}b_{33} \\ 0 & 0 & a_{33}b_{33} \end{bmatrix}$$

To implement the general case of multiplication of upper triangular matrices, we write the following loop:

Initialize
$$C(i,j) = 0.0 \text{ for } i = 1, n; j = 1, n$$
 Begin Loop
$$i = 1, n$$
 Begin Loop
$$j = i, n$$

$$k = i, j$$

$$C(i,j) = C(i,j) + A(i,k)B(k,j)$$
 End Loop
$$C(i,j) = C(i,j) + A(i,k)B(k,j)$$
 End Loop
$$C(i,j) = C(i,j) + A(i,k)B(k,j)$$
 End Loop

Notice that this is the inner product version MatMat.ijk but with the lower limits in the index j and k modified so that only entries in the upper triangular matrix are involved in the computation. Clearly, we want to avoid multiplying zero entries! In order to evaluate the flop count, we sum the amount of arithmetic associated with the most deeply nested statements in the algorithm. To this end, we obtain

$$\sum_{i=1}^{n} \sum_{j=i}^{n} 2(j-i+1) = \sum_{i=1}^{n} \sum_{j=1}^{n-i+1} 2j \approx \sum_{i=1}^{n} 2\frac{(n-i+1)^2}{2} = \sum_{i=1}^{n} i^2 \approx \frac{n^3}{3}$$
 (2.1)

Note that we used the following sums in the above derivation:

$$\sum_{n=1}^{n} p = \frac{n(n+1)}{2} \approx \frac{n^2}{2}$$

and also

$$\sum_{p=1}^{n} p^2 = \frac{n^3}{3} + \frac{n^2}{2} + \frac{n}{6} \approx \frac{n^3}{3}.$$

The corresponding cost for computing these matrices as full matrices is $2n^3$, and thus the savings with the above algorithm is six-fold.

In the same spirit, we could exploit the structure of other sparse matrices both in computing and in storing them. A particular structure that is often encountered in solving numerical partial differential equations is that of **banded matrices** of the form shown below, where we have a tridiagonal matrix [a(i), b(i), c(i)] with b(i) on the diagonal.

$$\mathbf{A} = \begin{pmatrix} b_1 & a_1 & & & 0 \\ c_1 & b_2 & a_2 & & \\ & \ddots & \ddots & \ddots & \\ 0 & & c_{n-1} & b_n & a_{n-1} \end{pmatrix}$$

First, it is advantageous to store such matrices in terms of their diagonals [a,b,c]. Next, we can easily verify that

$$\mathbf{A}\mathbf{x} = \mathbf{b}\mathbf{x} \oplus^u \mathbf{a}\mathbf{x}_1 \oplus^l \mathbf{c}\mathbf{x}_2$$

where $\mathbf{x}^T = (x_1, \dots, x_n)$ is a vector of length n but $x_1^T = (x_2, x_3, \dots, x_n)$ and $x_2^T = (x_1, x_2, \dots, x_{n-1})$ are vectors of length (n-1). Because of this mismatching in vector length, the summation \oplus in the above equation should be interpreted appropriately, i.e., we should add (\oplus^u) the diagonal \mathbf{a} above the main diagonal \mathbf{b} (here the second term in the sum) to the first component of \mathbf{bx} and also add (\oplus^l) the diagonal \mathbf{c} below the main diagonal to the last component of \mathbf{bx} . Also, the multiplications between all vectors should be performed in an element-wise fashion.

2.2.9 Gram-Schmidt Vector Orthogonalization

Important operations in scientific computing are vector orthogonalization and normalization. The Gram-Schmidt process starts with n linearly independent vectors \mathbf{x}_i and ends with n orthonormal vectors \mathbf{q}_i , i.e., vectors which are orthogonal to each other, and also their L_2 -norm is unity. Let us consider the vectors \mathbf{x}_i , $i = 0, \ldots, n-1$ each of length M. We want to produce vectors \mathbf{q}_i , $i = 0, \ldots, n-1$ where the first vector \mathbf{q}_0 is the normalized \mathbf{x}_0 , the vector \mathbf{q}_1 is orthogonal to \mathbf{q}_0 and normalized, the vector \mathbf{q}_2 is orthogonal to \mathbf{q}_0 and \mathbf{q}_1 , and so on. The idea is to produce a vector

$$\mathbf{y}_i = \mathbf{x}_i - (\mathbf{q}_{i-1}^T \mathbf{x}_i) \mathbf{q}_{i-1} - \ldots - (\mathbf{q}_0^T \mathbf{x}_i) \mathbf{q}_0$$

which subtracts the projection of \mathbf{x}_i onto each vector \mathbf{q}_j for $j = 0, \dots, i-1$. Having obtained \mathbf{y}_i we can then normalize it to obtain the corresponding orthonormal vector, i.e.,

$$\mathbf{q}_i = rac{\mathbf{y}_i}{\parallel \mathbf{y}_i \parallel_2}$$
 .

We can summarize this algorithm as follows:

- Initialize: Compute $r_{00} = \parallel \mathbf{x}_0 \parallel_2$. If $r_{00} = 0$ STOP, else $\mathbf{q}_0 = \mathbf{x}_0/r_{00}$.
- Begin Loop: For j = 0, ..., n-1 Do:
 - 1. Compute $r_{ij} = \mathbf{q}_i^T \mathbf{x}_j$, $i = 0, \dots, j-1$
 - 2. $\mathbf{y}_j = \mathbf{x}_j \sum_{i=0}^{j-1} r_{ij} \mathbf{q}_i$
 - 3. $r_{ij} = ||\mathbf{y}_i||_2$
 - 4. If $r_{jj} = 0$ STOP else $\mathbf{q}_j = \mathbf{y}_j/r_{jj}$
- End Loop.

Example: Let us assume that we want to orthonormalize the vectors

$$\mathbf{x}_0 = \begin{bmatrix} 1 \\ 0 \\ 2 \end{bmatrix} \text{ and } \mathbf{x}_1 = \begin{bmatrix} 2 \\ 3 \\ 0 \end{bmatrix}$$

thus, following the above algorithm, we obtain

•
$$r_{00} = (1^2 + 0^2 + 2^2)^{1/2} = 2.2367$$
 and $\mathbf{q}_0 = \frac{1}{2.2367} (1 \ 0 \ 2)^T$

SO

$$\mathbf{q}_0 = \left[\begin{array}{c} 0.4472 \\ 0 \\ 0.8942 \end{array} \right]$$

•
$$r_{01} = q_0^T \mathbf{x}_1 = 0.8942$$

and

$$\mathbf{y}_1 = \mathbf{x}_1 - r_{01}\mathbf{q}_0 = \begin{bmatrix} 2\\3\\0 \end{bmatrix} - 0.8942 \begin{bmatrix} 0.4472\\0\\0.8942 \end{bmatrix} = \begin{bmatrix} 1.6001\\3\\-0.7996 \end{bmatrix}$$

SO

$$r_{11} = ||y_1||_2 = [(1.6001)^2 + 3^2 + (-0.7996)^2]^{1/2} = 1.7887$$

and

$$\mathbf{q}_1 = \frac{\mathbf{y}_1}{r_{11}} = \begin{bmatrix} 0.89456\\ 1.6722\\ 0.4470 \end{bmatrix}$$

Notice that we can write

$$\begin{bmatrix} 1 & 2 \\ 0 & 3 \\ 2 & 0 \end{bmatrix} = \begin{bmatrix} 0.4472 & 0.89456 \\ 0 & 1.6772 \\ 0.8942 & 0.4470 \end{bmatrix} \begin{bmatrix} 2.2367 & 0.8942 \\ 0 & 1.7887 \end{bmatrix}$$

The algorithm above is presented using common mathematical abstractions, such as vectors and matrices. The beauty of C++ is that these mathematical abstractions can be implemented in C++ as "user-defined" data types, in particular for this case, as *classes*. We now present the implementation of the above algorithm in C++, utilizing some predefined user-defined classes that we have created. Explanation of the class syntax of this function will be given later in section 3.1.8, along with details as to how to create your own user-defined data types.



Gram-Schmidt Code

In the coding example below, we are using the *SCVector class* that we previously defined (section 2.1.1). Because we have defined the SCVector class to have mathematical properties just like what we would expect, we see that we can translate the algorithm given above directly into the code. Admittedly, some of the C++ syntax in the function below goes beyond what you have been taught thus far; the details of the class implementation of this code will be given later on in this book (see section 3.1.8). What you should notice, however, is that classes, such as SCVector, allow you to more closely model the mathematical definitions used in the algorithmic description of the solution of the problem.

```
SCstatus GramSchmidt(SCVector * x, SCVector * q){
  int dim = x[0].Dimension();
  SCVector y(dim);
  SCMatrix r(dim);
 r(0,0) = x[0].Norm_12();
  if(r(0,0)==0.0)
   return(FAIL);
  else
   q[0] = x[0]/r(0,0);
  for(j=1; j<dim; j++){</pre>
                               // corresponds to Begin Loop
    for(i=0;i<=j-1;i++)
      r(i,j) = dot(q[i],x[j]); // corresponds to 1
   y = x[j];
    for(i=0;i<=j-1;i++)
      y = y - r(i,j)*q[i]; // corresponds to 2
   r(j,j) = y.Norm_12(); // corresponds to 3
```

Observe in the code above that we allocate within this function an SCMatrix r which we use throughout the function, and which is discarded when the function returns to its calling function. We may want to retain r, however. In this case, we can create a function which has an identical name as the previous function but contains an additional variable within the argument list. The name and the argument list are used to distinguish which function we are referring to when we call the function. (this concept will be discussed further in section 4.1.4).



In the function below, we pass into the function GramSchmidt a SCMatrix r which it populates over the course of the computation.

```
SCstatus GramSchmidt(SCVector * x, SCVector * q, SCMatrix &r){
  int i,j;
  int dim = x[0].Dimension();
  SCVector y(dim);
 r(0,0) = x[0].Norm_12();
  if(r(0,0)==0.0)
   return(FAIL);
  else
   q[0] = x[0]/r(0,0);
  for(j=1; j<dim; j++){</pre>
                               // corresponds to Begin Loop
    for(i=0;i<=j-1;i++)
      r(i,j) = dot(q[i],x[j]); // corresponds to 1
   y = x[j];
    for(i=0;i<=j-1;i++)
     y = y - r(i,j)*q[i]; // corresponds to 2
   r(j,j) = y.Norm_12(); // corresponds to 3
```

Key Concept

• Classes can help you more closely mimic the natural data structures of the problem. We are not confined to working with only the low level concepts of integers, floats, and characters.

QR Factorization and Code

Another important point which we will often use in this book is a special **matrix factorization**. In particular, if the vectors \mathbf{x}_i , i = 0, ..., n-1 form the columns of a matrix \mathbf{X} of size $m \times n$, also \mathbf{q}_i , i = 0, ..., n-1 form the columns of matrix \mathbf{Q} , and r_{ij} are the entries of a square $n \times n$ matrix \mathbf{R} (which turns out to be upper triangular) the following equation is valid

$$X = QR$$

which is known as QR decomposition (or factorization) of the matrix \mathbf{X} , and it has important implications in obtaining eigenvalues and solutions of linear systems.



We now present a C++ function which accomplishes the QR decomposition of a matrix.

Just as was stated above, we input a matrix X to be decomposed into the matrices Q and R. We begin by creating two arrays of vectors q and v, which will serve as input to our original Gram-Schmidt routine. As you will see, this routine contains only two basic components:

1. A data management component, which is going from matrices to a collection of vectors and back, and

2. A call to the Gram-Schmidt routine that we wrote previously (and now you understand why we may have wanted to be able to retrieve the value of the SCMatrix r).

This routine demonstrates one important issue in scientific computing, i.e., the compromise between computational time and programmer's time. In this case, one may argue that if we were to write a routine specifically for QR decomposition, then we could reduce some of the cost of the data management section, and thus have a "more optimal code". However, this consideration must be balanced by considering how much computational time is used for data manipulation versus the time to properly write and debug an entirely new function. In this particular case, in theory, we have already written and tested our GramSchmidt(v,q,R) function, and hence we are confident that if we give the Gram-Schmidt function proper inputs, it will return the correct solution. Hence, we can focus our programming and debugging on the extension of the concept, rather than on the details of optimization. Optimization is certainly important if we were to be calling this routine many times in a particular simulation; however, optimization-savy individuals, as the old saying goes, often miss the forest for the trees!

```
SCstatus QRDecomposition(SCMatrix X, SCMatrix &Q, SCMatrix &R){
  int i,j;
  int num_vecs = X.Rows();
  int dim = X.Columns();
  SCstatus scflag;
  Vector *q = new SCVector[num_vecs](dim),
         *v = new Vector[num_vecs](dim);
  for(i=0;i<num_vecs;i++){</pre>
    for(j=0;j<dim;j++)</pre>
      v[i](j) = X(j,i);
  }
  scflag = GramSchmidt(v,q,R);
  for(i=0;i<num_vecs;i++)</pre>
    for(j=0;j<dim;j++)</pre>
      Q(j,i) = q[i](j);
  return scflag;
}
```

Modified Gram-Schmidt Algorithm and Code

Notice that the Gram-Schmidt method breaks down at the k^{th} stage if \mathbf{x}_k is linearly dependent on the previous vectors \mathbf{x}_j , $j=0,\ldots,k-2$ because $\parallel \mathbf{x}_k \parallel_2 = 0$. It has also been observed that, in practice, even if there are no actual linear dependencies, orthogonality may be lost because of finite arithmetic and round-off problems, as discussed earlier. To this

end, a **modified Gram-Schmidt** process has been proposed which is almost always used in computations. Specifically, an intermediate result is obtained,

$$\mathbf{y}_{i}^{0} = \mathbf{q}_{i} - (\mathbf{q}_{0}^{T} \mathbf{x}_{j}) \mathbf{q}_{0},$$

which we project onto \mathbf{q}_0 (instead of the original \mathbf{x}_j), as follows:

$$\mathbf{y}_{i}^{1} = \mathbf{y}_{i}^{0} - (\mathbf{q}_{1}^{T} \mathbf{y}_{i}^{0}) \mathbf{q}_{1},$$

and so on. This process then involves successive one-dimensional projections. In the following, we present a row-oriented version of the modified Gram-Schmidt algorithm.

- Initialize: Set $\mathbf{q}_i = \mathbf{x}_i$, $i = 0, \dots, n-1$.
- Begin Loop: For i = 0, ..., n-1 Do:

$$r_{ii} = ||\mathbf{q}_i||_2$$

 $\mathbf{q}_i = \mathbf{q}_i/r_{ii}$
For $j = i+1, \dots, n-1$ Do:
 $r_{ij} = \mathbf{q}_i^T \mathbf{q}_j$
 $\mathbf{q}_j = \mathbf{q}_j - r_{ij} \mathbf{q}_i$
End Loop

• End Loop.



We present a C++ implementation of the modified Gram-Schmidt algorithm below. With the exception of the commented block of code, the remaining code is identical to the original code provided above.

```
SCstatus ModifiedGramSchmidt(Vector * x, Vector * q, Matrix &r){
  int i,j;
  int dim = x[0].Dimension();
  Vector y(dim);

  r(0,0) = x[0].Norm_12();

  if(r(0,0)==0)
    return(FAIL);
  else
   q[0] = x[0]/r(0,0);
```

}

```
for(j=1;j<dim;j++){
 /* We replace the following block of lines from the
                                         */
 /* original Gram-Schmidt algorithm presented above,
                                         */
 /* for(i=0;i<=j-1;i++)
                                         */
      r(i,j) = dot(q[i],x[j]);
 /*
                                         */
 /*
                                         */
 /*
      y = x[j];
                                         */
      for(i=0;i<=j-1;i++)
                                         */
        y = y - r(i,j)*q[i];
 /*
                                         */
 /*
                                         */
 /* with the modification described above.
                                         */
 /* following lines implement that modification.
                                         */
 y = x[j];
 for(i=0;i<=j-1;i++){
  r(i,j) = dot(q[i],y);
  y = y - r(i,j)*q[i];
 End of Modification
 r(j,j) = y.Norm_12();
 if(r(j,j) == 0)
  return(FAIL);
  q[j] = y/r(j,j);
}
return(SUCCESS);
```

Remark 1: The computational complexity of the Gram-Schmidt process is $\mathcal{O}(mn^2)$ irrespective of which version is used. This is evident by comparing the comment block inserted into the Modified Gram-Schmidt code. If you carefully examine the deleted code versus the newly inserted code, you will see that the number of operations that is performed is identical. It is often the case in scientific computing that although two algorithms may be

identical mathematically (i.e., in infinite precision), one algorithm is inherently better than the other when implemented numerically. Furthermore, in this case, we see that we achieve an additional benefit from the modified algorithm at no additional cost.

Remark 2: The loss of orthogonality of \mathbf{Q} in the modified Gram-Schmidt method depends on the condition number $\kappa(\mathbf{A})$ of the matrix \mathbf{A} obtained by using the specified vectors as columns [8]. In general, the orthogonality of \mathbf{Q} can be completely lost with the classical Gram-Schmidt method while the orthogonality property may not be lost with the modified Gram-Schmidt method but it may not be acceptable when the matrix \mathbf{A} is ill-conditioned. A better approach is to employ the Householder method discussed in section 9.3, which is more accurate and also computationally less expensive. For example, the cost for Gram-Schmidt is $\mathcal{O}(mn^2)$ while for the Householder method is $\mathcal{O}(mn^2 - n^3/3)$.

2.3 Parallel Computing

• Imagine a large hall like a theater, except that the circles and galleries go right round through the space usually occupied by the stage. The walls of this chamber are painted to form a map of the globe...A myriad of computers are at work upon the weather of the part of the map where each sits, but each computer attends only to one equation or part of an equation. The work of each region is coordinated by an official of higher rank...From the floor of the pit a tall pillar rises to half the height of the hall. It carries a large pulpit on its top. In this sits the man in charge of the whole theater; he is surrounded by several assistants and messengers. One of his duties is to maintain a uniform speed of progress in all parts of the globe. In this respect he is like the conductor of an orchestra in which the instruments are slide rules and calculating machines. But instead of waving a baton he turns a beam of blue light upon those who are behindhand.

Lewis F. Richardson, "Weather Prediction By Numerical Process" (1922)

This prophetic quote describes quite accurately the many hardware and software ingredients of a modern parallel computer. It refers to a multiple instruction/multiple data type and involves domain decomposition as the mode of partitioning the work load. The concepts of master node that synchronizes the processes as well as of load balancing are also included in the statement.

In the following, we briefly review some parallel computer architectures and introduce parallel concepts and tools.

2.3.1 From Supercomputing to Soupercomputing

A supercomputer is the fastest computer of its time; today's supercomputer is tomorrow's desktop or laptop computer. One of the first supercomputers of historical significance was the Cray-1. It was used quite successfully in many applications involving large-scale simulation in the early 1980s. The Cray-1 was not a parallel computer, however, but it employed a powerful (at the time) vector processor with many vector registers attached to the main memory (see figure 2.13). Today, all supercomputers are parallel computers. Some are based

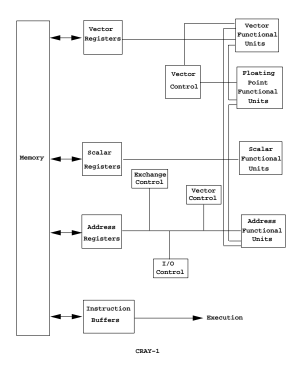


Figure 2.13: Schematic of the first Cray computer, the Cray-1.

on specialized processors and networks, but the majority are based on commodity hardware and open source operating system and applications software. In this section, we will review briefly some of the history and the recent trends.

Types of Parallel Computers

A popular taxonomy for parallel computers is the description introduced by Michael Flynn in the mid 1960s [36] of the programming model as single instruction/ multiple data stream (SIMD) or multiple instruction/ multiple data stream (MIMD). In a SIMD computer, such as the Thinking Machines CM-2 or the NCUBE Inc. computers of the 1980s, each processor performs the same arithmetic operation (or stays idle) during each computer clock, as controlled by a central control unit (see figure 2.14). In this model (also referred to as a data parallel program) high-level languages (e.g., CM Fortran, C^* , and Lisp) are used, and computation and communication among processors are synchronized implicitly at every clock period.

On a MIMD computer (see figure 2.15) each of the parallel processing units executes operations independently of each other, subject to synchronization through appropriate message passing at specified time intervals. Both parallel data distribution as well as the message passing and synchronization are under user control. Examples of MIMD systems include the Intel Gamma and Delta Touchstone computers and, with fewer but more powerful processors, the Cray C-90, and the first generation of IBM SP2 (all made in the 1990s).

While it is often easier to design compilers and programs for SIMD multiprocessors because of the uniformity among processors such systems may be subject to great computational inefficiencies. This is due to their inflexibility when stages of a computation are

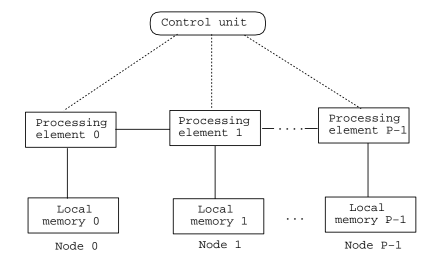


Figure 2.14: Schematic of SIMD parallel computer.

encountered in which there is not a large number of identical operations. There has been a natural evolution of multiprocessor systems towards the more flexible MIMD model, especially the merged programming model in which there is a single program (perhaps executing distinct instructions) on each node. This merged programming model is a hybrid between the data parallel model and the message passing model and was successfully exemplified in the Connection Machine CM-5. In this SPMD (single program multiple data) model, data parallel programs can enable or disable the message passing mode, and thus one can take advantage of the best features of both models.

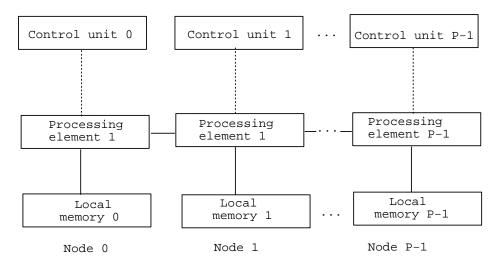


Figure 2.15: Schematic of MIMD parallel computer.

MIMD computers can have either shared memory as the SGI Origin 2000 or distributed memories as in the IBM SP system. The issue of shared memory requires further clarification as it is different from the centralized memory. Shared memory means that a single address space can be accessed by every processor through a synchronized procedure. In non-shared memory systems explicit communication procedures are required. The prevailing paradigm

in parallel computing today is one where the physical memory is distributed, but the address space is shared as this is a more flexible and easier as a programming environment.

PC Clusters

The most popular and cost-effective approach to parallel computing is cluster computing, based for example, on PCs running the Linux operating system (hereafter referred to merely as Linux). The effectiveness of this approach depends on the communication network connecting the PCs together, which may vary from fast Ethernet to Myrinet that can broadcast messages at a rate of several Ggabits per second (Gbs).

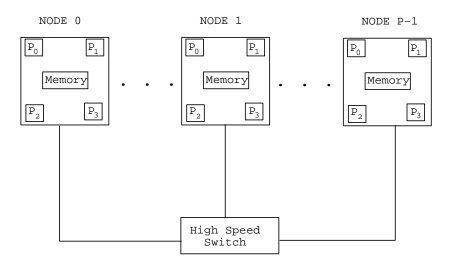


Figure 2.16: Schematic of Generic Parallel Computer (GPC).

Issues of computer design, balancing memory, network speed, and processing speed can be addressed by examining the Generic Parallel Computer (GPC) depicted in figure 2.16. The key components of the GPC are an interconnecting set of P processing elements (PE) with distributed local memories, a shared global memory, and a fast disk system (DS). The GPC serves as a prototype of most PC-based clusters that have dominated supercomputing in the last decade both on the scientific as well as the commercial front.

The first PC cluster was designed in 1994 at NASA Goddard Space Flight Center to achieve one Gigaflop. Specifically, 16 PCs were connected together using a standard Ethernet network. Each PC had an Intel 486 microprocessor with sustained performance of about 70 Megaflops. This first PC cluster was built for only \$40,000 compared to \$1 million, which was the cost for a commercial equivalent supercomputer at that time. It was named Beowulf after the lean hero of medieval times who defeated the giant Grendel. In 1997 researchers at the Oak Ridge national laboratory built a Beowulf cluster from many obsolete PCs of various types; for example, in one version it included 75 PCs with Intel 486 microprocessors, 53 Intel Pentium PCs and five fast Alpha workstations. Dubbed the stone soupercomputer because it was built at almost no cost, this PC heterogeneous cluster was able to perform important simulations producing detailed national maps of ecoregions based on almost 100 million degrees of freedom [54]. A picture of this first soupercomputer is shown in figure 2.17.



Figure 2.17: Soupercomputer of the Oak Ridge national laboratory. (Courtesy of F. Hoffman)

Building upon the success of the first such system, the BEOWULF project [7, 81], several high performance systems have been built that utilize commodity microprocessors with fast interconnects exceeding one Gigabits per second in bandwidth. *Moore's law* (an empirical statement made in 1965 by the Intel co-founder Gordon Moore) suggests that the performance of a commodity microprocessor doubles every 18 months, which implies that, even without fundamental changes in the fabrication technology, processors with a speed of several tens of Gigaflops can become available. Nanotechnology can help in prolonging the validity of this statement, which has been true for at least four decades. New developments include the TeraHertz transistor and the packaging of more than one billion transistors on a single chip will hopefully keep Moore's law alive. Intel's Pentium-4 (see figure 2.12) has about 42 million transistors).

In addition to enhancements in the speed of individual processors, there have been several key developments that have enabled commodity supercomputing:

- The development and maturization of the free operating system Linux, which is now available for all computer platforms. The freely distributable system and the open source software movement has established Linux as the operating system of choice, so almost all PC clusters are Linux based.
- The MPI standard that has made parallel coding portable and easy. There are several implementations such as MPICH, SCore, etc. but they all share the same core commands which we present in this book.
- The rapid advances in interconnect and fast switches with small latencies, which are now widely available unlike the early days of proprietory and expensive systems available only by a few big vendors.

Grid Supercomputing

The computational grid is a new distributed computing paradigm, similar in spirit to the electric power grid. It provides scalable high-performance mechanisms for discovering and

negotiating access to geographically remote resources. It came about by the internet and world wide web advances and the fact that similarly to Moore's law for computer speed, the speed of networks *doubles* every about nine months. This is twice the rate of Moore's law, and it implies that the performance of a wide area network (WAN) increases by two orders of magnitude every five years!

Computing on remote platforms involves several steps, to first identify the available sites, to negotiate fast access to them, and configure the local hardware and software to access them. The Grid provides the hardware and software infrastructure that allows us to do this. The community-based opensource Globus toolkit is the most popular software infrastructure [38], see also

http://www.globus.org

It implements protocols for secure identification, allocation and release of resources from a globally federated pool of supercomputers, i.e., the Grid.

The Grid also allows the implementation of network-enabled solvers for scientific computing, such as the package NetSolve [14]. NetSolve searches for available computational resources within the Grid and chooses the best available resource based upon some sort of match-making procedure. It consists of three parts: a client, an agent, and a server. Client is the user issueing a request that is received by the agent. The latter allocates the best server or servers which perform the computation and return the results to the client. The server is a daemon process, which is on the alert awaiting requests from the client.

Performance Measurements and Top 500

As regards performance of parallel computers, there is no universal yardstick to measure it, and in fact the use of a single number to characterize performance such as the peak performance quoted by the manufacturer is often misleading. It is common to evaluate performance in terms of benchmark runs consisting of kernels, algorithms, and applications so that different aspects of the computer system are measured. This approach, however, is still dependent on the quality of software rather than just hardware characteristics. The controversy over performance evaluation methods has been recognized by the computer science community and there have been several recent attempts to provide more objective performance metrics for parallel computers [57]. A discussion of some of the most popular benchmarks, the BLAS routines, was presented in 2.2.7, and more information can be found on the web at:

http://www.netlib.org/benchmark

A good basis for performance evaluation of supercomputers is also provided in the Top500 list, see:

URL: http://www.top500.org/

This was created by Dongarra in the early 1990s and it is updated twice a year. This list reports the sites around the world with the 500 most powerful supercomputers. Performance on a LINPACK benchmark [28] is the measure used to rank the computers. This is a code that solves a system of linear equations, see chapter 9, using the best software for each

platform. Based on the data collected so far and the current Teraflop sustained speeds achieved, it is predicted that the first PETAFlop/s (10^{15} floating point operations per second) supercomputer would be available around 2010 or perhaps sooner.

2.3.2 Mathematical Parallelism and Recursive-Doubling

We now review briefly the mathematics of parallelism. There are many basic mathematical operations that have a high degree of parallelism, and by this we mean that they can be performed simultaneously and independently of each other. Consider, for example, the element-wise multiplication of two vectors \mathbf{x}, \mathbf{y} to produce another vector \mathbf{c} , i.e.,

$$c_i = x_i y_i, \quad i = 1, \dots, N.$$

Clearly, in this case all N products can be obtained simultaneously, and thus we can imagine that each term in this product being evaluated by a different computer. In this particular example there is of course no need to engage N different processors to do such a simple operation, but the point we want to make is that for this operation there are no dependencies among the different pairs of data. This is an example of perfect mathematical parallelism. This form of parallelism also applies to the task of finding the maximum in each pair of a set of N pairs of numbers, i.e., $max(x_i, y_i)$, i = 1, ..., N. This operation is also perfectly parallel, and we will refer to such problems as EP (Embarassingly Parallel).

Notice, however, that if we attempt to find the absolute maximum number in the above pairs we introduce inter-dependencies among the data, and such an operation is not perfectly parallel anymore. The same is true for simple BLAS operations, for example the evaluation of an inner (dot) product, i.e.,

$$c = \sum_{i=1}^{N} x_i y_i,$$

where c is a scalar. This requires the summation of all N product pairs (x_iy_i) , which is clearly a *serial* operation as it involves accumulation of the sum of a next pair to the previous accumulation, and so on. Another such example is the evaluation of a polynomial p(x), e.g.

$$p(x) = a_0 + a_1 x + a_2 x^2 + a_3 x^3 + \ldots + a_N x^N,$$

at a specific point x_0 , which is also an accumulation operation. For this example, a straightforward computation would require recomputing the powers of x or else we would require extra storage. This last issue can be avoided by resorting to *Horner's rule* and alternating multiplications and additions appropriately. For example, the polynomial

$$p(x) = 2 + 3x + 7x^2 + 4x^3 + x^4,$$

can be computed from the equivalent equation

$$p(x) = 2 + (3 + (7 + (4 + x)x)x)x,$$

which can be computed recursively with (N-1) multiplication and N additions. This is an important point: observe that the mathematical result of the two forms is the same, however

the number of operations that must be accomplished to obtain the result is different. This type of operation rearrangement for optimization is common (and powerful) in scientific computing. This particular operation, however, is still *serial* because in order to proceed with the next iteration we need the results from the previous one.

Let us now revisit the computation of the inner product mentioned above. To be able to perform the addition of the terms (x_iy_i) faster than in the straightforward serial fashion from left to right, we can break the sum into two smaller sums (assuming for convenience that N is an even number). For example, we have that the two sets are

$$\sum_{i=1}^{N/2} x_i y_i \quad \text{and} \quad \sum_{i=N/2+1}^{N} x_i y_i.$$

We can compute the two sums separately and then collect the two results and perform another sum to obtain the final result. The total number of additions is, of course, the same, i.e., (N-1) but assuming that we can execute the two big sums simultaneously the wall clock time is almost half of what it was before. We can estimate more precisely the wall clock time by assuming that it takes the computer a time δt to perform an addition. This time δt is related to the processor speed, and it is in the range of several nanoseconds for a relatively fast computer. The total time required for the straightforward approach is $T_1 = (N-1)\delta t$. Then, the total time required after we break up the problem into two subproblems is $T_2 = (N/2-1)\delta t + \delta t + C$, where C represents the time required to collect the two results from the sub-sums. We can measuse the speed-up of this data partionining method from the ratio

$$S_2 = \frac{T_1}{T_2} = \frac{N-1}{N/2 + C/\delta t}$$
.

For efficiency we want S_2 to be larger than one, which will be true if the relative communication-to-computation time $C/\delta t$ is small. If this cost is negligible then for N very large we obtain $S_2 \approx 2$, which is the theoretical maximum speed-up.

This simple exercise illustrates how we can extract parallelism from an operation that is seemingly serial at a first glance. Moreover, this data partitioning approach can be continued recursively so that the sub-sums are smaller and smaller, to a single pair of numbers in this particular approach. This **divide-and-conquer** approach is fundamental to parallel processing thinking. It is also known by other names, e.g. the fan-in algorithm or the recursive-doubling algorithm. Another advantage of recursive-doubling is that it leads to enhanced numerical stability because when we sum up a large set of numbers in a serial fashion significant accumulation of errors can occur. However, the recursive doubling algorithm and corresponding pair-wise summation is assured to be more stable.

We can now generalize the example above by assuming that we have P computer processors available, and that P = N, with $N = 2^q$, so that we can reduce the evaluation of a dot product into a summation of two numbers on each processor. The total number of stages or branches in this tree, which is illustrated in figure 2.18 for the case of N = 8, is q; here q = 3. Using the aforementioned recursive thinking, we can estimate the speed-up factor to be

$$S_P = \frac{T_1}{T_P} = \frac{(N-1)\delta t}{q\delta t + qC},$$

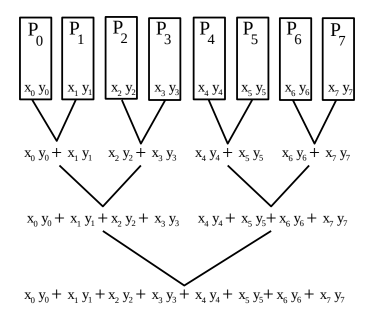


Figure 2.18: Inner (dot) product operation following the fan-in algorithm for N = P = 8.

where we include a total communication cost of qC assuming that in each stage the communication time penalty to collect terms and send them to other processors is C; let us denote the relative time by $\alpha = C/\delta t$. We can rewrite the speed-up factor in terms of the problem size N or the number of processors P

$$S_P = \frac{N-1}{(1+\alpha)\log_2 N} = \frac{P-1}{(1+\alpha)\log_2 P}.$$

We see from the last equation that even for zero communications ($\alpha = 0$), the theoretical maximum speed-up factor is

$$S_P = \frac{P-1}{\log_2 P} < P$$
 and $\eta_P = \frac{P-1}{P \log_2 P}$,

where the last equation defines the parallel efficiency η_P . For an EP (embarassingly parallel) problem we have that $\eta_P = 1$ or 100%. It is clear that in this recursive algorithm the parallel efficiency is less than perfect even if we ignore communications among the processors. At successive stages of the computations less and less processors are involved in the computation, and in fact in the last stage only two processors are involved while (P-2) are staying idle. The parallel efficiency, therefore, should reflect this load imbalance amongst the P processors.

If communication is taken into account in the above simple model, which assumes that $C = \alpha \delta t$, then we have that

$$S_P(\alpha = 1) = \frac{1}{2} S_P(\alpha = 0).$$

Therefore, we are computing at only 50% parallel efficiency. In practice, there are other factors which may limit the parallel efficiency even more. For example, the number of processors may not match exactly with the size of the problem, the total problem or subproblems may not fit in the memory, or a startup time penalty known as *latency* may slow

down the transferring of data. In this latter case, a better model for the communication cost is

$$C = L + \beta l$$

where L is the latency and l is the message length. Also β^{-1} is the bandwidth, which reflects the maximum rate at which messages can be exchanged. Latency is significant when we transfer small parcels of data often, but for longer size messages it is a subdominant time cost. Typical ranges for L are from a few nanoseconds to a few microseconds, and it is usually inversely proportional to the cost of the computer multiprocessor.

2.3.3 Amdahl's Law

A more general model for the speed-up factor was proposed by Gene Amdahl (1967), which is often referred to as Amdahl's law [2]. In its simplest form it assumes that some percentage, say ξ , of the program or code cannot be parallelized, and that the remaining $(1 - \xi)$ is perfectly parallel. Neglecting any other communication delays such as memory contention, latencies, etc. Amdahl's model for speed-up states that

$$S_P = \frac{T_1}{[\xi + (1 - \xi)/P]T_1} = \frac{1}{\xi + \frac{1 - \xi}{P}},$$
(2.2)

and thus an upper bound in the limit of $P \to \infty$ is $S_P \le 1/\xi$. This implies that even if $\xi = 1\%$ (meaning only 1% of the program is not parallelizable) for P = 100 we have that $S_{100} = 50$, so we operate at half the maximum efficiency; see also figure 2.19.

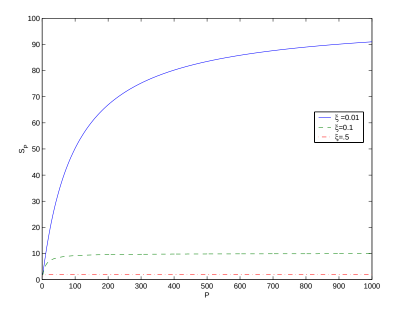


Figure 2.19: Speed-up factor versus number of processors for three different degrees of parallelizable code.

Although useful, this performance measure S_P can sometimes be misleading since it may favor inefficient but highly parallelizable algorithms instead of more efficient algorithms that may be more difficult to map onto a parallel multiprocessor computer. Also, the derivation

of Amdahl's law relies on the assumption that the serial work ξ is independent of the size of the problem size N. In practice, it has been observed that ξ decreases as a function of problem size. Therefore, the upper bound on the speed-up factor S_P usually increases as a function of problem size. Another anomaly is the so-called *superlinear speed-up*, which means that the speed-up factor has been measured to be more than P. This may happen because of memory access and cache mismanagement or because the serial implementation on a single processor is suboptimal.

There are several industry standard benchmark programs such as Whetstone, ScaLA-PACK and LINPACK benchmarks. While these benchmarks have been used extensively in all advanced computer system evaluations, specific benchmarks have been developed to evaluate shared, distributed, and hybrid memory parallel computers. They vary from simple parallel loops to measure the ability of parallelizing compilers to the PERFECT benchmark which consists of thirteen programs including several fluid dynamics programs, to the MIMD benchmarks, e.g. Genesis consisting of FFTs, partial differential equations, molecular dynamics, and linear algebra. A particularly popular set of benchmarks was developed at NASA Ames, the NAS parallel benchmarks, a suite of eight programs with three versions for serial, machine-dependent, and MPI-based implementations:

(see URL: http://www.nas.nasa.gov/Software/NPB).

Measures of performance based on Amdahl's law are particularly effective for small programs that do not require extensive and intensive use of computer memory. Most computer benchmarks are of this sort, but they do not in fact represent many of the requirements for the solution of large-scale simulations.

2.3.4 MPI - \underline{M} essage \underline{P} assing \underline{I} nterface

Parallel computing, in its conceptual form, appears to be a very reasonable concept. Many of the concepts found in parallel computing have analogous concepts in social areas such as business management. The idea of many people working together toward one goal is similar to many processes working together toward one solution. The idea of wanting to partition the work so that all processors are busy and none remain idle is similar to wanting to keep your team busy, with noone having to sit around waiting on someone else for information. From this perspective, we see that parallel computing is a natural extension of the concept of divide-and-conquer; that is, we first begin with a problem which we want to solve, we then access the available resources we can use toward solving the problem (which in the case of computing will be the number of processors that can be used), and we then attempt to partition the problem into manageable pieces which can be done concurrently by each person on the team.

Key Concept

• Parallel Computing is a Divide-and-Conquer Strategy.

The most common difficulty that people have with this concept is not the divide-and-conquer component; most often, we are quite comfortable with the *concept of partitioning* a problem into workable subproblems. However, it may be difficult to see how a specific problem can be partitioned efficiently to be solved in parallel, and secondly to understand how you can get computers to work in parallel. Both of these issues, at varying levels, will be addressed in this book.

Throughout this book, we will be discussing how to take specific numerical algorithms and partition them among multiple processors. Hence, at this stage of your reading, we will focus on the second of the two major difficulties:

• Understanding and implementing parallel programming on a computer.

The first issue that we will draw your attention to is that the goal of this book is not to give detailed descriptions of the hardware and software mechanisms used for implementing parallel computers. Our goal in this book, and the goal of most simulation scientists, is to use the parallel computing tools already designed and created for them. Hence, we focus on the problem we are trying to solve, and not on the intricate details of getting computers to communicate with each other. To this end, we have chosen to employ MPI, a Message-Passing Interface, for handling our parallel computing communication needs.

There are many good books on MPI itself, e.g. [51] and [73], both of which give some of the history and the developmental rational behind a message passing interface. We will not cover this information other than to mention that the basic premise behind MPI is that multiple parallel processes work concurrently towards a common goal using "messages" as their means of communicating with each other. This idea is illustrated in figure 2.20. Multiple MPI processes may run on different processors, and these processes communicate through the infrastructure provided by MPI. As users, we need not know the implementation of this infrastructure, we need only to know how to take advantage of it! To this end, almost everything in MPI can be summed up in the single idea of "Message Sent - Message Received."

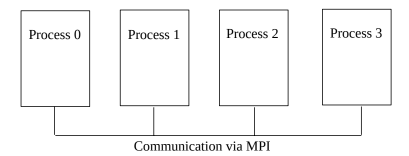


Figure 2.20: Schematic of MPI processes working together.

Key Concept

• Almost everything in MPI can be summed up in the single idea of "Message Sent - Message Received."

We will be discussing the mechanics of this process, i.e., the MPI routines to use in order to accomplish this, throughout this book. Let us begin our parallel computing experience by understanding from a high-level the differences between our previous "serial" codes (those programs that we have been addressing previous to this point) and our new "parallel" codes. Up to now, every program that we have discussed has had one single thread of execution. A simplistic breakdown is the following: the program started, data was allocated, work was done on that data, and an answer was produced. All the steps were presented in a serial fashion: A then B then C, etc. Think of this model as the "you working alone" model. You, as a single person, have a goal to accomplish, and you set out with a specific collection of tasks that need to be done in order to get the job done. You decide an order in which these need to be accomplished, and then you plow through them one-by-one. Now, imagine that you are working with someone else toward a common goal. You must now ask yourself the following question:

• How can I partition the problem so that we can utilize our resources and accomplish the goal in the least amount of time (or with the least effort)? This is a parallelization question!

This parallelization question can be broken down into the following two questions:

- What work can be accomplished concurrently?
- When is communication between concurrent processes necessary?

Take the following example: Suppose that you are taking a class in which this book is being used as the text. The goal (hopefully) is for everyone to read the text. If there is only one copy of the text, then one possible strategy is for the instructor to prepare an ordered list of all those in the class, and for each person to read the text when it is his turn. This is thus a serial process; you as a student must wait until your turn in order to read the book, and must remain idle (with respect to the reading the book) until you have obtained a copy. However, if the goal is for everyone to read the book, then a more optimal solution with respect to time is for everyone to have a copy of the book simultaneously! Then each person can be reading the book concurrently. This is an example of an **Embarrassingly Parallel Algorithm**.

Now, how would such an algorithm be implemented in parallel using MPI? We will begin with a modification of our "Hello World" program. We must first specify our goal: We want each process to print to the screen "Hello World!" Since this does not require any communication, this should be able to be done concurrently by all the processes involved.



Our first MPI program to accomplish this is the following program:

```
#include <iostream.h>
#include <mpi.h>
int main(int argc, char ** argv){
   MPI_Init(&argc,&argv);
   cout << "Hello World!" << endl;
   MPI_Finalize();
}</pre>
```

Compilation and execution details for MPI programs is provided in the Appendix B.1. What is the first thing that you should notice about MPI? The programming interface to MPI is a collection of functions! MPI is a library of functions designed to handle all the nitty-gritty details of message passing on the architecture on which you want to run. We notice several things immediately about the program above:

- We included mpi.h. This provides us with the function declarations for all MPI functions.
- We must have a beginning and an ending. The beginning is in the form of an $MPI_Init()$ call, which indicates to the operating system that this is an MPI program and allows the OS to do any necessary initialization. The ending is in the form of an $MPI_Finalize()$ call, which indicates to the OS that "clean-up" with respect to MPI can commence.
- If the program is embarrassingly parallel, then the operations done between the MPI initialization and finalization involve no communication.

When you compile and run this program, you obtain a collection of "Hello World!" messages printed to your screen. The number of messages is equal to the number of processes on which you ran the program ⁷. This is your first parallel program!

The two MPI functions that you used in the previous program have the following form:

• MPI_Init

⁷Though the printing by the program is done concurrently, the computer must serialize the output in order to write it to the screen. We still consider this to be an embarrassingly parallel process.

• MPI_Finalize

```
int MPI_Finalize(void)
```

In this book, we will take for granted the information that is passed to MPI_Init. Just as in Figure 2.20, our MPI program started up a number of processes, each of which printed its message. The most natural question to ask is:

• How does a process know which number he is?

There are two important commands very commonly used in MPI:

• MPI_Comm_rank

```
int MPI_Comm_rank(

MPI_Comm comm /* in */,
int* result /* out */)
```

• MPI_Comm_size

```
int MPI_Comm_size(

MPI_Comm comm /* in */,
int* size /* out */)
```

The first of these two functions, MPI_Comm_rank , provides you with your process identification or rank (which is an integer ranging from 0 to P-1, where P is the number of processes on which are are running), and MPI_Comm_size provides you with the total number of processes that have been allocated. The argument com is called the communicator, and it essentially is a designation for a collection of processes which can communicate with each other. MPI has functionality to allow you to specify varies communicators (differing collections of processes); however, in this book, whenever a communicator must be specified, we will always use MPI_COMM_WORLD , which is predefined within MPI and consists of all the processes initiated when we run our parallel program.

If you were running a P=8 process job, MPI_Comm_size would pass back the number 8 as the total number of processes running, and MPI_Comm_rank would return to you a number 0, 1, ..., 7 denoting which process you currently were executing within. How do you use this information?



Let us modify our code above to not only have each MPI process print out "Hello World!" but also to tell us which process the message is coming from, and how many total processes with which it is joined.

```
#include <iostream.h>
#include <mpi.h>
int main(int argc, char ** argv){
  int mynode, totalnodes;
```

```
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

cout << "Hello world from process " << mynode;
cout << " of " << totalnodes << endl;

MPI_Finalize();
}</pre>
```

When run four processes, the screen output may look like:

```
Hello world from process 0 of 4
Hello world from process 3 of 4
Hello world from process 2 of 4
Hello world from process 1 of 4
```

Note, as we pointed out, the output to the screen may not be ordered correctly since all processes are trying to write to the screen at the same time, and the operating system has to decide on an ordering. However, the thing to notice is that each process called out with its process identification number and the total number of MPI processes of which it was a part.

It is at this point that we want to make a critical observation: when running with MPI, all processes use the same compiled binary, and hence all processes are running the exact same code. What in an MPI distinguishes a parallel program running on P processors from the serial version of the code running on P processors? Two things distinguish the parallel program:

- 1. Each process uses its *process rank* to determine what part of the algorithm instructions are meant for it.
- 2. Processes communicate with each other in order to accomplish the final task.

Even though each process receives an identical copy of the instructions to be executed, this does not imply that all processes will execute the same instructions. Because each process is able to obtain its process rank (using MPI_Comm_rank), it can determine which part of the code it is supposed to run. This is accomplished through the use of **if** statements. Code that is meant to be run by one particular process should be enclosed within an **if** statement, which verifies the process identification number of the process. If the code is not placed with **if** statements specific to a particular id, then the code will be executed by all processes (like in the case of the code above). Shortly we will show you a parallel program in which this fact is illustrated.

We, as of yet, have not actually accomplished the second point, communicating between processes (from the programmers point of view); we have merely relied on the operating system and other software system layers to handle the initiation and termination of our MPI processes across multiple processors. Recall that we said that MPI can be summed up in the concept of sending and receiving messages. Sending and receiving is done with

the following two functions: MPI_Send and MPI_Recv . Below we present the function syntax, argument list description, a usage example and some remarks for the MPI_Send and MPI_Recv commands.

Function Call Syntax

```
int MPI_Send(
         void*
                         message
                                        in
         int
                         count
                                        in
         MPI_Datatype
                         datatype
                                        in
                         dest
                                        in
         int
         int
                         tag
                                        in
         MPI_Comm
                                        in
                         comm
int MPI_Recv(
         void*
                         message
                                        out
        int
                         count
                                        in
         MPI_Datatype
                         datatype
                                        in
                         source
         int
                                        in
         int
                         tag
                                        in
         MPI_Comm
                                        in
                         comm
         MPI\_Status*
                                        out
                         status
```

Understanding the Argument Lists

- message starting address of the send/recv buffer.
- count number of elements in the send/recv buffer.
- datatype data type of the elements in the send buffer.
- *source* process rank to send the data.
- dest process rank to receive the data.
- taq message tag.
- *comm* communicator.
- status status object.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be sent/recv
int sender; // process number of the sending process
```

```
int receiver; // process number of the receiving process
              // integer message tag
int tag;
                     // variable to contain status information
MPI_Status status;
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
// Determine datasize
double * databuffer = new double[datasize];
// Fill in sender, receiver, tag on sender/receiver processes,
// and fill in databuffer on the sender process.
if(mynode==sender)
 MPI_Send(databuffer,datasize,MPI_DOUBLE,receiver,
           tag,MPI_COMM_WORLD);
if(mynode==receiver)
 MPI_Recv(databuffer,datasize,MPI_DOUBLE,sender,tag,
            MPI_COMM_WORLD,&status);
// Send/Recv complete
```

Remarks

- In general, the *message* array for both the sender and receiver should be of the same type and both of size at least *datasize*.
- In most cases the *sendtype* and *recvtype* are identical.
- The tag can be any integer between 0-32767.
- MPI_Recv may use for the tag the wildcard MPI_ANY_TAG . This allows an MPI_Recv to receive from a send using any tag.
- MPI_Send cannot use the wildcard MPI_ANY_TAG. A specific tag must be specified.
- MPI_Recv may use for the source the wildcard MPI_ANY_SOURCE. This allows an MPI_Recv to receive from a send from any source.

• MPLSend must specify the process rank of the destination. No wildcard exists.



To get you started, we will begin with a small numerical example. Imagine that we want to sum up all the numbers from 1 to 1000. We could implement this as the following *serial code*:

#include<iostream.h>

```
int main(int argc, char ** argv){
  int sum;

sum = 0;

for(int i=1;i<=1000;i=i+1)
    sum = sum + i;

cout << "The sum from 1 to 1000 is: " << sum << endl;
}</pre>
```

Instead, let us use multiple processes to do the job. Now admittedly, multiple processes are not needed for this job, but the point of this exercise is for you to see how we partition the problem. The first thing we realize is that to effectively sum up all the numbers from 1 to 1000 using multiple processes, we want to partition the sums across the processes. Suppose that we use only two processes, then we want that process 0 sums from 1 to 500, and process 1 sums from 501 to 1000, and then at the end, the two values are added together to obtain the total sum of all numbers from 1 to 1000. A schematic of this is provided in figure 2.21. Given P processes, the problem of summing is partitioned into P subproblems, and then at the end all processes send their information to process 0 for the final accumulation of the results.

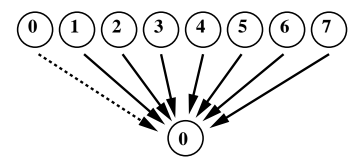


Figure 2.21: Gathering of all information to one process using sends and receives.

The first question we must ask ourselves is how do we partition the processing? Recall from our discussion above that every process can find out how many total processes are

being used, and which process it is (by using MPI_Comm_size and MPI_Comm_rank, respectively). Let mynode be the variable storing the result of the call to MPI_Comm_rank, and let totalnodes be the variable storing the result of the call to MPI_Comm_size. Then, the formula for partitioning the sums across the processes is given by the following code:

```
startval = 1000*mynode/totalnodes+1;
endval = 1000*(mynode+1)/totalnodes;
```

If you use only one process, then totalnodes = 1 and mynode = 0, and hence startval = 1 and endval = 1000. If you are using two processes, then totalnodes = 2, and mynode is either 0 or 1. For mynode = 0, startval = 1 and endval = 500, and for mynode = 1, startval = 501 and endval = 1000. You can continue this procedure until you are using 1000 processes, at which point each process is only summing one value (i.e., not summing at all!), and all the values are sent to process zero for accumulation.

Once we have the starting value and ending value of our sum, each process can execute a for loop which sums up the values between its startval and its endval. Then, once the local accumulation is done, each process (other than process 0) sends its sum to process 0.



The code below is a C++/MPI program which accomplishes this:

```
#include<iostream.h>
#include<mpi.h>
int main(int argc, char ** argv){
  int mynode, totalnodes;
  int sum,startval,endval,accum;
  MPI_Status status;
  MPI_Init(argc,argv);
  MPI_Comm_size(MPI_COMM_WORLD, &totalnodes); // get totalnodes
  MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
                                               // get mynode
  sum = 0; // zero sum for accumulation
  startval = 1000*mynode/totalnodes+1;
             1000*(mynode+1)/totalnodes;
  endval =
  for(int i=startval;i<=endval;i=i+1)</pre>
     sum = sum + i;
  if (mynode!=0)
   MPI_Send(&sum,1,MPI_INT,0,1,MPI_COMM_WORLD);
  else
```

```
for(int j=1;j<totalnodes;j=j+1){
    MPI_Recv(&accum,1,MPI_INT,j,1,MPI_COMM_WORLD, &status);
    sum = sum + accum;
}

if(mynode == 0)
    cout << "The sum from 1 to 1000 is: " << sum << endl;

MPI_Finalize();
}</pre>
```

We will present more detailed information about the *MPI_Send* and *MPI_Recv* commands later in this book (see section 3.4). Note however, the general structure of the message passing. First, observe that there is an *if* statement, which distinguishes between whether you are process 0 or any other process. Why? Because recall, all processes other than process zero are **sending**, whereas process 0 is **receiving**. We should decide our programs so that for each message sent using an the command *MPI_Send*, there is some receiving process.

Hence, whereas each process other than 0 has one MPI_Send call, process 0 has (P-1) MPI_Recv calls (where P is the total number of processes used). This is an important concept to understand. Often times an MPI program has been sitting idle because one process was sending, and there were no process waiting to receive!

This portion of the text was not meant to be all inclusive, nor are you expected to be able to go out and write MPI codes using sends and receives with blinding efficiency (especially since we have not yet explained the argument lists for MPI_Send and MPI_Recv). However, as you go through this book, you will slowly but surely accumulate MPI knowledge and experience, and at the end, you will hopefully be writing C++/MPI code with confidence!

2.4 Homework Problems

- 1. Prove that the condition number in the L_2 -norm of an othogonal matrix is 1, and that the condition number (in any norm) of a matrix is greater or equal one.
- 2. Use the classical Gram-Schmidt and the modified Gram-Schmidt algorithms to orthonormalize the vectors

$$\mathbf{x}_0 = \begin{pmatrix} 1\\10^{-4}\\0 \end{pmatrix} \text{ and } \mathbf{x}_1 = \begin{pmatrix} 1\\0\\10^{-4} \end{pmatrix}$$

Compare the two results. What do you observe?

- 3. Find the eigenvalues of an $n \times n$ matrix with all entries equal to 1.
- 4. Modify Collatz-B so that instead of using the expression $x_{n+1} = 3x_n + 1$ you use $x_{n+1} = 5x_n + 1$. Use the number 100 as an initial guess and with the maximum number of iterations set to 10,000. What do you observe? Postulate as to why this happens.

(*Hint*: Have you ever played an arcade game in which you eventually were scoring a negative number?)

5. The Fibonacci sequence is a sequence of integers created by the following inductive process: given $f_0 = 0$ and $f_1 = 1$, the next number in the sequence is equal to the sum of the previous two numbers, i.e.,

$$f_n = f_{n-1} + f_{n-2}.$$

This process can be continued indefinitely and produces the Fibonacci sequence: 0, 1, 1, 2, 3, 5, 8, 13

- (a) Write a program which allows the user to input a desired number of terms N of the Fibonacci sequence, and outputs the sequence f_0, f_1, \ldots, f_N to the screen.
- (b) Modify your program so that you keep track of the ratio $\frac{f_n}{f_{n-1}}$. How many terms do you need so that the difference between $\frac{f_n}{f_{n-1}}$ and $\frac{f_{n-1}}{f_{n-2}}$ is less than 1.0×10^{-5} ?
- (c) What is the value of this ratio? The number you have converged to is called the *Golden Mean*.
- 6. The harmonic series $\sum_{k=1}^{\infty} 1/k$ diverges, i.e., it grows without bound as we include more terms. Consider the truncation that includes the first n terms, which we call partial sum S_n and can be computed recursively from $S_n = S_{n-1} + 1/n$ with $S_1 = 1$. What is the largest S_n that can be obtained in your computer in single precision?
- 7. The Pythagorean theorem states that the sum of the squares of the sides of a right triangle is equal to the square of the hypotenuse. Thus, if x and y are the lengths of the two sides of a right triangle, and z is the length of the hypotenuse, then $x^2 + y^2 = z^2$. Fermat, a lawyer and amateur mathematician during the 1600s, postulated that there

exists no other integer m (other than m=2) such that $x^m+y^m=z^m$ for the sides of a right triangle as described above.

- (a) Write a function *power* which takes as input a double precision number x and an integer m and returns as output a double precision number which is equal to the value of x raised to the power m (i.e., x^m).
- (b) Write a function *pythagoreus* which takes as input two double precision numbers x and y and an integer m and returns as output a double precision number which is equal to the value of $x^m + y^m$. Use the function *power* written above.
- (c) Write a program which queries the user for three values: two double precision number which equal the length of two of the sides of a right triangle, and an integer N. Your program should first use the function pythagoreus to determine the value of the square of the hypotenuse. Then you should write a loop which checks to see if there exists any integer $2 < m \le N$ such that $z^m = x^m + y^m$. If you find such a value of m, print the result and break out of the loop. If you do not find a value m such that the above expression is true, print a message which states that no value can be found for the value N provided by the user.
- 8. Change the stride of the summing example given in section 2.3 (make the stride of the additions equal to the number of processors). This will require devising new formulae for the variables *startval* and *endval*, and changing the i = i + 1 used within the summing loop to some other increment.
- 9. Modify the summing example as follows:
 - (a) At the beginning of the main function, add an integer variable *master*, and initialize this value to some number between zero and the number of processors minus one.
 - (b) Modify the MPI_Send/MPI_Recv sequence such that all processes except master send, and process master receives.
 - (*Hint*: From the example, in the MPI_Send , the '0', denotes that you are sending to process zero; in the MPI_Recv , the j denotes the process from which a message is being received. These will need to be modified.)
 - (c) Output the sum from master.
 - (d) Add *cout* statements so that each sending process prints a message stating to whom it is sending, and add *cout* statements so that the receiving process acknowledges from whom it has received.
- 10. Modify the summing example as follows:
 - (a) Instead of summing integers, change the appropriate variables so that you will now sum doubles. You will need to use MPI_DOUBLE instead of MPI_INT within the MPI calls. Verify that you obtain the same answer as the integer case.
 - (b) Change the sum so that you are summing $\frac{1}{i}$ instead of i.

(c) At the top of the program, immediately following the #include<iostream.h> statement, add #include<iomanip.h>. Then prior to calling the cout statement, add the following line:

cout << setprecision(20);</pre>

After making these changes and recompiling, run your program on 2,4, and 8 processes. What differences in the sum of $\frac{1}{i}$ do you see? Postulate as to why this is so.

- 11. Modify the parallel MPI code to do the following:
 - (a) Have process zero query the user for the number of elements over which to sum.
 - (b) From process zero, distribute to all processes the number of elements to sum (using sends and receives) and appropriately calculate the interval over which each process is to sum.
 - (c) Accomplish the summing as is already done in the program.
 - (d) After creating the final answer on process zero, print the result.

Chapter 3

Approximation

Two of the most common tasks in scientific computing are interpolation of discrete data and approximation by known functions of the numerical solution, the source terms, and the boundary or initial conditions. Therefore, we need to perform these tasks both accurately and efficiently. The data are not always nicely distributed on a uniform lattice or grid, and thus we must learn how to manage these situations as well. We often use polynomials to represent discrete data as they are easy to "manipulate," i.e., differentiate and integrate. However, sines and cosines as well as special functions, called wavelets, are very effective means to perform interpolation and approximation, and they have very interesting properties.

In this section, we will study different such representations and their corresponding C++ implementations. We consider cases where the data are just sufficient to determine exactly the representation (deterministic case) as well as cases where the data are more than the information needed (overdetermined case).

Finally, we will present a more detailed discussion of MPI_Send and MPI_Recv, the two fundamental building blocks of MPI.

3.1 Polynomial Representation

In this section we will study different ways of interpolating data on equidistant and more general grids using polynomials. We will discuss both accuracy and efficiency and will introduce C++ arrays and other concepts to effectively implement the algorithms.

3.1.1 Vandermonde and Newton Interpolation

Assuming that we have the data available on the discrete set of points $\{x_0, x_1, \ldots, x_N\}$ with corresponding values $\{f(x_0), f(x_1), \ldots, f(x_N)\}$, then we can construct a function f(x) that passes through the pairs $(x_i, f(x_i))$ by the approximation

$$f(x) \approx p_N(x) = \sum_{k=0}^{N} a_k \phi_k(x) ,$$

where $p_N(x)$ is referred to as the interpolating polynomial, $\phi_k(x)$ are a priori known polynomials, and a_k are the unknown coefficients. We call $\phi_k(x)$ the basis, and its choice is very important in obtaining an efficient approximation. For example, assuming that $\phi_k(x) = x^k$, $k = 0, \ldots, N$, then we have the following representation at the known pair $(x_i, f(x_i))$

$$f(x_i) = a_0 + a_1 x_i + a_2 x_i^2 + \ldots + a_N x_i^N, \quad i = 0, \ldots N.$$

All together we have (N+1) such equations for the (N+1) unknowns a_i , $i=0,\ldots,N$. This system of equations can be recast in matrix form with the vector of unknowns $\mathbf{a}^T = (a_0, a_1, a_2, \ldots, a_N)$ as follows

$$\begin{bmatrix} 1 & x_0 & x_0^2 & \dots & x_0^N \\ 1 & x_1 & x_1^2 & \dots & x_1^N \\ & & & & \\ 1 & x_N & x_N^2 & \dots & x_N^N \end{bmatrix} \begin{bmatrix} a_0 \\ a_1 \\ \vdots \\ a_N \end{bmatrix} = \begin{bmatrix} f(x_0) \\ f(x_1) \\ \vdots \\ f(x_N) \end{bmatrix},$$

or in compact form

$$\mathbf{V}\mathbf{a} = \mathbf{f}$$

where the matrix V is known as the *Vandermonde matrix*. This matrix is non-singular because we assume that all $\{x_0, x_1, \dots x_N\}$ are distinct points, and therefore there exists a unique polynomial of order N that represents this data set. We could obtain the vector of coefficients \mathbf{a} from

$$\mathbf{a} = \mathbf{V}^{-1} \mathbf{f}$$

by inverting the Vandermonde matrix V and subsequently performing matrix-vector multiplications with $f(x_i)$. This, however, is an expensive operation with a cost of $\mathcal{O}(N^3)$ to invert the matrix V (see chapter 9), and it is rarely used in practice.

One approach in reducing the computational complexity is to simply change the basis to

$$\phi_k(x) = \prod_{i=0}^{k-1} (x - x_i),$$

so f(x) is now approximated by

$$f(x) \approx a_0 + a_1(x - x_0) + a_2(x - x_0)(x - x_1) + \dots + a_N(x - x_0)(x - x_1) \dots (x - x_{N-1}).$$
 (3.1)

Notice that we still use a polynomial basis, but we have simply shifted it with respect to the coordinates of the data points. This simple shift turns out to have a dramatic effect since now the *new* unknown coefficients can be computed by inverting the following system

$$\begin{bmatrix} 1 & 0 & \dots & & & & 0 \\ 1 & (x_1 - x_0) & \dots & & & & 0 \\ \vdots & & & & & & & \\ 1 & (x_N - x_0) & \dots & (x_N - x_0) & (x_N - x_1) & \dots & (x_N - x_{N-1}) \end{bmatrix} \begin{bmatrix} a_0 \\ a_1 \\ \vdots \\ a_N \end{bmatrix}$$

$$= \begin{bmatrix} f(x_0) \\ f(x_1) \\ \vdots \\ f(x_N) \end{bmatrix},$$

which is a lower triangular matrix and requires only $\mathcal{O}(N^2)$ operations in order to obtain the vector of unknown coefficients. This is done by simple forward substitution, and can be implemented readily using BLAS2.

Remark: It is instructive to compare this method, which is called *Newton interpolation*, with the Vandermonde interpolation. Assuming that we use Gauss elimination to obtain the vector of unknown coefficients (see chapter 9), we see that the change of basis in the Newton approach takes us directly to the second stage of Gauss elimination, which is the forward substitution, while in the Vandermonde approach we have to essentially perform an LU decomposition of the matrix \mathbf{V} , which is an $\mathcal{O}(N^3)$ operation. However, the Vandermonde matrix is a special one, and its inversion can also be done in $\mathcal{O}(N^2)$ operations (e.g. using FFTs, see section 3.2). Thus, the two approaches discussed here are computationally equivalent.

Newton Interpolation: Recursive Algorithm

There is a nice recursive property that we can deduce from Newton's interpolation method, and which can be used for writing compact C++ code as we shall see in the next section.

Solving for the first few coefficients, we obtain

$$a_{0} = f(x_{0})$$

$$a_{1} = \frac{f(x_{1}) - f(x_{0})}{x_{1} - x_{0}}$$

$$a_{2} = \frac{\frac{f(x_{2}) - f(x_{0})}{x_{2} - x_{0}} - \frac{f(x_{1}) - f(x_{0})}{x_{1} - x_{0}}}{x_{2} - x_{1}}$$

:

so we see that the coefficient

$$a_k = \mathcal{F}(x_0, x_1, \dots, x_k) \,,$$

that is the k^{th} coefficient is a function of the first k function values $f(x_k)$. \mathcal{F} is a function of both the x_k variables and the $f(x_k)$ data (and hence, in the end, since f(x) is a function of x, then really \mathcal{F} is just a function of the x_k 's as given above).

To obtain a recursive relation for the coefficient a_k we need to write the approximation in the grid

$$G_0^k \equiv \{x_i\}, \quad i = 0, \dots k$$

where the subscript denotes the starting index and the superscript denotes the ending index. To this end, we consider the two subsets

$$G_0^{k-1} \equiv \{x_0, x_1, \dots, x_{k-1}\}, \text{ and } G_1^k \equiv \{x_1, x_2, \dots, x_k\},$$

of k grid points each. We also denote the corresponding polynomial approximations by $p_0^k(x)$, $p_0^{k-1}(x)$ and p_1^k formed by using the grids G_0^k , G_0^{k-1} and G_1^k , respectively. We then observe that

$$(x_0 - x_k)p_0^k(x) = (x - x_k)p_0^{k-1}(x) - (x - x_0)p_1^k(x),$$
(3.2)

as the polynomial $p_0^k(x)$ passes through all the pairs

$$(x_i, f(x_i)), \quad i = 0, \dots, k.$$

Next, upon substitution of $p_0^k(x)$, $p_0^{k-1}(x)$ and $p_1^k(x)$ in equation (3.2) by their full expansions, which are

$$p_0^k(x) = a_0 + a_1(x - x_0) + \ldots + a_k(x - x_0) \ldots (x - x_{k-1})$$

$$p_0^{k-1}(x) = a_0 + a_1(x - x_0) + \ldots + a_{k-1}(x - x_0) \ldots (x - x_{k-2})$$

$$p_1^k(x) = b_1 + b_2(x - x_1) + \ldots + b_k(x - x_1) \ldots (x - x_{k-1}).$$

and comparing the coefficients of highest polynomial power, x^k , we obtain:

$$(x_0 - x_k)a_k = a_{k-1} - b_k$$

or

$$(x_0 - x_k)\mathcal{F}(x_0, x_1, \dots x_k) = \mathcal{F}(x_0, x_1, \dots x_{k-1}) - \mathcal{F}(x_1, x_2, \dots x_k)$$

and therefore

$$\mathcal{F}(x_0, x_1, \dots x_k) = \frac{\mathcal{F}(x_0, \dots x_{k-1}) - \mathcal{F}(x_1, \dots x_k)}{x_0 - x_k}.$$
 (3.3)

We thus obtain the higher *divided differences* (i.e., coefficients) from the lower ones from equation (3.3).

We illustrate this procedure on a grid G_0^2 containing three grid points (x_0, x_1, x_2) , so that

$$\mathcal{F}(x_0) = f(x_0); \quad \mathcal{F}(x_1) = f(x_1); \quad \mathcal{F}(x_2) = f(x_2),$$

then at the next level

$$\mathcal{F}(x_0, x_1) = \frac{\mathcal{F}(x_0) - \mathcal{F}(x_1)}{x_0 - x_1}$$

$$\mathcal{F}(x_1, x_2) = \frac{\mathcal{F}(x_1) - \mathcal{F}(x_2)}{x_1 - x_2}$$

and

$$\mathcal{F}(x_0, x_1, x_2) = \frac{\mathcal{F}(x_0, x_1) - \mathcal{F}(x_1, x_2)}{x_0 - x_2},$$

and so on, for grids with more points.

3.1.2 Arrays in C++

So far, when we have discussed variables in C++, we have referred to single variables, such as the variables mynode and totalnode presented in section 2.3.4. Now, mathematically, we just introduced a collection of variables in the form of a sequence: $x_0, x_1, x_2, ...x_N$. If you were to write a program which involved such a sequence of numbers, how would you declare these variables? Of course, to start with, you may use the knowledge you gained from section 2.1.2 to decide how to declare the variables. The variable declaration would look like the following (for N = 5):

double x0, x1, x2, x3, x4, x5;

This does not seem too difficult. However, imagine that you want to use 100 points! Do you want to type x0, x1, ..., x99? And even more annoying, suppose that you want to compare the results of running a program using 50 points compared to 1000 points! Do not be dismayed; C++ has a solution to your problem! The C++ solution to this problem is the concept of arrays. In C++, you can allocate a block of memory locations using the concepts of arrays. There are two means of accomplishing this: static allocation and dynamic allocation. We will discuss both briefly.

Static Allocation of Arrays

The first means by which you can allocate an array is to *statically* allocate the array. For our purposes, we will take this to mean that prior to both compilation and execution, the size of the array is known. In the previous section, we discussed the idea of using a discrete set of points $\{x_0, x_1, \ldots, x_N\}$ for interpolation. For a specific example, let us take N = 99 (so that the total number of points is 100 points), and let us assume that we want our grid points to be evenly spaced in the interval [0, 1].



The following piece of code would statically allocate an array of 100 doubles, and would fill in those variables with their appropriate positions in the interval [0,1]:

```
#include <iostream.h>
int main(int argc, char * argv[]){
  int i;
  double x[100];
  double dx = 1.0/99.0;

  for(i=0;i<100;i++)
    x[i] = i*dx;

  for(i=0;i<100;i++)
    cout << "x[" << i << "] = " << x[i] << endl;
}</pre>
```

Let us now examine in detail the statements in this program. First, notice the syntax used for allocating static arrays:

```
<type> <variable name> [ size ]
```

Here, *size* is the number of memory positions that you want allocated. In our example, we wanted 100 doubles to be allocated. Once the allocation is done, how do we access these variables? C++ uses [] for accessing variables in an array. In the above allocation, x[0] is the first element, x[1] is the second element, etc. There are several key points for you to realize:

- C++ array indexing always begins at 0. Hence, the first position in an array is always the position denoted by [0].
- C++ does not verify that you do not overrun an array. To overrun an array is to attempt to access a memory location which has not been allocated to the array. In the above example, trying to access x[100] would be illegal because we only allocated an array containing 100 elements (indexed 0,...,99). C++ will not complain when compiling, but may cause a segmentation fault (or even far worse, it may run normally but give the wrong results!). You should be very careful not to overrun arrays!
- When *statically* allocating arrays, you cannot use a variable for the size parameter. Hence the following C++ code is *invalid*:

```
int npts = 100;
double x[npts];
```

Your C++ compiler will complain that this is illegal! This is because it is not until the program is actually executed that the value of npts is known to the program (recall that upon execution npts is both allocated in memory, and then intialized to the value 100). This type of operation can be done with dynamic memory allocation, which will also be discussed below.

• We can, however, index the array using variables. In the above example, we are able to iterate through all the values of the array using a *for* loop.

WARNING

Programmer Beware!

- C++ arrays always begin their indexing at 0!
- Array allocations are done by size, not the final index value! Hence if you allocate an array with 100 elements, you would index them from 0,...,99!

Implicit and Explicit Casting

At this stage, let us interject a brief note concerning a common mistake made by programmers first learning C++. Notice in the above example that we have allocated a variable dx, and we have initialized it to 1.0/99.0. What would happen if we were to write 1/99? The answer: the variable would be set to 0.0 instead of 0.01010101 as we would have expected. Why you might ask? All binary operations in C++ are type-specific. Although we use the '+' symbol to add two integers together, and we also use the '+' to add two floats together, the operations which the computer accombishes to execute these two operations are different. What happens when you want to mix-and-match variable types? Suppose that you want to add 10.0+1, where 10.0 is a floating point value, and 1 is an integer? The answer: C++ will implicitly cast the value 1 to 1.0 (i.e., from an integer value to a floating point value), and then will carry out the binary operation of '+' between two floating point values. Casting is the conversion of information stored by a variable from one type to another. Implicit casting implies that the casting operation is done automatically for you (C++ also allows explicit casting, which we also mention below). The order of casting is presented in figure 3.1. In the figure, there is a pictorial example of implicit casting.

Why will 1/99 yield the value 0? Since the value 1 is an integer, and the value 99 is an integer, then the division operator used will be the integer division operator '/'. As an integer operation, 99 divides 1 zero times, and hence the solution is zero! However, if we were to write 1.0/99, we now have a floating point value divided by an integer, and hence the computer must first cast the integer value to a float. Operationally, the computer would first cast the value 99 to the value 99.0, and then would use the floating point division operator '/' to divide 1.0 by 99.0, yielding 0.01010101.



Programmer Beware!

• Incorrect assumptions concerning implicit casting can lead to erroneous answers!

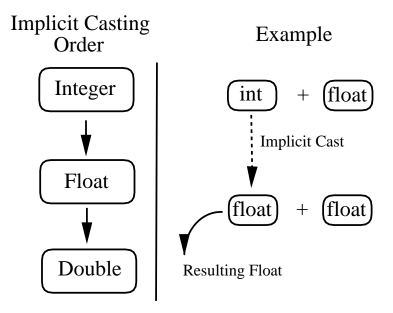


Figure 3.1: Implicit casting precedence.

C++ also allows *explicit casting*; that is, it allows you, the programmer, to specify that you want a value cast to a different type. The syntax used for explicit casting is the following:

```
(type to cast to) <variable name>
```

For example, suppose that we want to explicitly cast the value of an integer variable to a floating point variable. We could write the following:

```
int x = 1;
float y;

y = (float) x;
```

In this example, the value of x is explicitly cast to a floating point value, and then the floating point variable is assigned that value. In the above example, if you were not using the explicit casting operator (float), the variable would have to be implicitly cast. Explicit casting is useful in identifying to both you - the programmer - and to the computer the places that you expect a casting call to occur.

Dynamic Allocation of Arrays

Recall from our static allocation example, we stated that in order for an array to be allocated statically, the size of the array must be known prior to compilation and execution. Suppose that we want to specify the size of an array "on-the-fly;" that is, suppose that we want the user to be able to input the size of an array, and the program, while executing, to allocate the properly sized array. How can this be done? This can be accomplished in C++ using dynamic allocation. Dynamic allocation occurs using the **new** operator.



Let us re-examine our example from above. Suppose that we want the user to be able to input the number of points into which he would like to partition the interval [0,1]. This can be accomplished with the following code:

```
#include <iostream.h>
int main(int argc, char * argv[]){
  int i,npts;
  double *x; //declaration of pointer variable 'x'
  double dx:
  cout << "Enter the number of points in [0,1]: ";</pre>
  cin >> npts;
  x = new double[npts]; // dynamic allocation of npts doubles
  dx = 1.0/(npts-1);
  for(i=0;i<npts;i++)</pre>
    x[i] = i*dx;
  for(i=0;i<npts;i++)</pre>
    cout << "x[" << i << "] = " << x[i] << endl;
  delete[] x; // deallocation of dynamically allocated
                // memory
}
```

We will now analyze the differences between this code and the one presented previously: First, you will notice that we allow the user to input the number of points into the program. This is accomplished through the use of *cin*. This class, like *cout*, is declared in the system header file *iostream.h*. By using *cin*, we can obtain data from standard input, which in this case is from the keyboard. Recall from the previous chapter the following facts:

- cin reads from standard input (which for us is the keyboard)
- Observe that the direction of the >> is opposite of *cout*. This is for a reason. For *cout*, you are taking information from the variables that you provide to the expression and are providing that information to the operating system to be written to the screen. In the case of *cin*, you obtain data from the operating system, and place it into the variables provided in calling statement.
- The program must read one data item for each variable in the *cin* input list. Hence if we have the statement <code>cin</code> >> a >> b; where both a and b are declared as integers,

then cin will expect that there are two distinct integers for it to read from standard input.

- You may place one or more blank (space) characters to delineate numerical values.
- Any blank space prior to the value to be read is ignored.

Immediately following the computer's execution of the *cin* statement, the variable *npts* has a value equal to the number of points that the programmer wants for this discretization. Next, we allocate an array *dynamically* using the **new** operator. This is but one step in a three-step process (to be described further in just a moment). We then do the operations just as we had done before, and we conclude by "freeing" the memory that we allocated by using the **delete/delete**[] operator. To recapitulate, the three steps that occur in dynamic memory allocation are:

- 1. Declaration of a *pointer* variable.
- 2. Allocation of memory using the **new** operator.
- 3. Deallocation of memory using the **delete/delete**[] operator.

We will now discuss each of these three steps individually. We begin with the first step: declaration of a *pointer* variable. Pointers are variables which hold *addresses*. They may hold addresses to integer variables; they may hold addresses to floating point or double precision variables. Pointer variables are type-specific.

Key Concept

• Pointer variables are type-specific.

In the above example, we have obtained from the *cin* statement the number of points that the user wants to use. This is stored in the integer variable *npts*. We now want to allocate an array of size *npts*. First, notice that we have declared a variable x of type double*. The use of the '*' prior to the variable name designates this as a pointer variable (i.e., a variable that holds the address in memory of a double) instead of declaring a double. To recapitulate:

double x;

declares a variable named 'x' of type double (that is, x is a variable valid for containing a double precision number), whereas

double *x;

declares a variable named 'x' of type "double *" ¹, which is a variable valid to contain the address in memory of a double precision variable. For now, you should take this mechanism for granted; this distinction will become more apparant as we use pointers more and more.

One common mistake when declaring multiple pointers is to misplace the '*'. Take for example the following two declarations:

```
double *x1,y1;
double *x2,*y2;
```

The first of these two statements declares a variable named x1 of type double* and then declares a variable named y1 of type double (not double*). However, in the second declaration, both x2 and y2 are declared of type double star.

Once a pointer variable has been declared, we can ask the operating system to allocate for us a block of memory to contain our collection of doubles. This process is accomplished through the use of **new**. The **new** command asks the operating system (OS) for a block of memory of the type and size for which we ask, and returns to us the *address* of that block of memory. If no memory is available (or for whatever other reason the OS may decide not to be cooperative), the **new** command will return *NULL*, which is called the "Null pointer" or "Null address." This implies that we have not successfully obtained our request. In the code above, we ask the OS for *npts* doubles in the following syntactic form:

```
< address > new < type > [ size ]
```

If the OS is successful in giving us the memory we request, we will have a block of memory which we can access just as we did in the static memory allocation case. To access the first double in the array, we access x[0], and so forth.

At the conclusion of our program, or at some stage of the program where the array is no longer needed, we should release the memory used for this array back to the operating system for reuse. This process is called *deallocating* memory. To accomplish this deallocation, we use the **delete** [] command. In our case

```
delete[] x;
```

informs the OS that the memory that was allocated to the pointer variable x is no longer needed, and can be released. There are actually two delete operators, **delete** and **delete** [] which are used for informing the OS that dynamically allocated memory can be reclaimed. The distinction between the aforementioned operators is the following:

• **delete** (with no [] following it) is used when only a single object has been allocated. Suppose we were to allocate space to hold an integer as follows:

```
int * a = new int;
```

Space for only one integer was allocated (i.e., one integer object was allocated), hence to deallocate we would merely use **delete** as follows:

¹Programmers quite often actually say "double star" when pronouncing this variable type.

delete a;

• **delete** [] is used when an array of objects has been allocated. Suppose we were to allocate space to hold 20 integers as follows:

```
int * a = new int[20];
```

Space for 20 integers was allocated (i.e., an array of 20 integer objects was allocated), hence to deallocate we would use the **delete** [] as follows:

delete[] a;

Remark: Both **delete** and **delete**[] take only one argument. A comma separated list is not valid, and although it will compile, it will not give the desired result.

Allocation/Deallocation Rule-of-Thumb: If your allocation statement new requires the use of [], then so does the deallocation statement delete.

WARNING

Programmer Beware!

• Once memory is deallocated, you should not use it!

Passing Arrays to Functions

Finally, we want to draw your attention to passing arrays in C++. Passing arrays in C++ comes down to passing the pointer variable. Whether you declared the array dynamically or statically, you still end up passing a pointer. This is one subtle point which is not apparent when you declare arrays statically: statically declared arrays are really just pointers in disguise. Whether you are dealing with statically declared or dynamically declared arrays, you will pass them to functions in the same fashion.



In the program below, we have encapsulated the generation of the grid into a function called "CreateGrid_EvenlySpaced." This function takes as arguments the size of the array npts and the pointer variable x. With these two pieces of information, the function can successfully fill in the array, as had been done before.

```
int main(int argc, char * argv[]){
  int i,npts;
  double *x;
  double dx;
  cout << "Enter the number of points in [0,1]: ";</pre>
  cin >> npts;
  x = new double[npts];
  CreateGrid_EvenlySpaced(nts, x, 0.0, 1.0);
  for(i=0;i<npts;i++)</pre>
    cout << "x[" << i << "] = " << x[i] << endl;
  delete[] x;
}
// Definition of CreateGrid_EvenlySpaced is in SCchapter3.cpp
void CreateGrid_EvenlySpaced(int npts, double *x,
                                double a, double b){
  double dx = (b-a)/(npts-1.0);
  for(int i=0;i<npts;i++)</pre>
    x[i] = a + i*dx;
  return;
}
```

There are three things that we want to draw your attention to in this example:

- 1. Note that in both the declaration and definition of the function $CreateGrid_EvenlySpaced$ we declare x as a variable of type double*; this is important. The compiler needs to know (and recognize) that it is passing an address.
- 2. Note also that we have to pass *npts* into the function *CreateMesh*. What if we did not? Recall that variables within functions are local to the function. Hence, if we do not explicitly *tell* the function that the size of the array is *npts*, it has no way of knowing it!
- 3. Observe in the example above that for including the standard header file *iostream.h* we use #include<iostream.h>, whereas to include our user-defined header file *SCchapter3.h* we use #include "SCchapter3.h". The angled bracket notation < ... > is to used

when including standard library headers obtained from the standard include directory (files such as *iostream.h*, *iomanip.h*, *math.h*, etc.). To include user-defined header files from the current directory, quotation marks " . . . " are used.

Passing by Value Versus Passing by Reference

Recall in the previous chapter that we mentioned that unless otherwise stated all variables passed to a function are, by default, *passed by value*. We now want to clarify the difference between the ideas of *passing by value*, *passing by reference*, and *passing the address*.

• Passing by Value - When a variable is passed to a function by value, a new memory location is allocated, and a copy of the contents of the variable are placed in the new memory location. The function may then locally modify the contents of the new location. When the function returns, the new memory location is released and its contents lost. Consider the following program:

```
#include <iostream.h>

void func(int a);

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

   b = 4;
   func(b);
   cout << "value of b = " << b << endl;
}

void func(int a){
   cout << "value of a = " << a << endl;
   a = 64;
}</pre>
```

In the program above, the variable b is passed to the function func by value. When the function is executed, a new memory location (distinct from the memory location of the variable b) is allocated on the function stack and assigned to the variable name a locally within the function. The contents of b are copied into the location associated with the variable a. The first cout statement prints that the value of a is b. The local function variable b is then assigned the value b, and the function returns. Upon returning, the local memory is returned to the system. The second b is still b, it was uneffected by the function.

• Passing by Reference - When a variable is passed to a function by reference, no new memory location is allocated; instead, the local variable within the function is assigned to refer to the same memory location as the variable being passed. Consider the following program:

```
#include <iostream.h>

void func(int &a);

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

   b = 4;
   func(b);
   cout << "value of b = " << b << endl;
}

void func(int &a){
   cout << "value of a = " << a << endl;
   a = 64;
}</pre>
```

In the program above, the variable b is passed to the function func by reference. Notice the placement of the & in both function declaration and definition; this syntax informs the compiler that we want the variable passed by reference. When the function is executed, the local variable a is assigned to the same memory location as b. The first cout statement will print that the value of a is 4. The local function variable a is then assigned the value 64, and the function returns. Since the local variable a referred to the same memory location as b, the second cout statement prints that the value of b is 64.

• Passing the Address - Instead of passing by value or by reference, we have the third option of passing (explicitly) the address of the variable. When an address is passed, the address value is stored in a pointer variable of the appropriate type. Consider the following program:

```
#include <iostream.h>

void func(int *a);

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

  b = 4;
  func(&b);
  cout << "value of b = " << b << endl;
}

void func(int *a){
  cout << "value of a = " << *a << endl;</pre>
```

```
*a = 64;
```

In the function func above, we declare the input argument to be a pointer to an integer. When the function func is called from within main, we use the operator &, which stands for "take the address of". Inside of the function func, an integer pointer variable is created, and the address of b is stored there. The contents of the memory location to which a points can be accessed using the operator *, which stands for "the memory location that is pointed to by". The first cout statement will print that the value of *a is 4. The memory to which a points is then assigned the value 64, and the function returns. Since the local variable a pointed to the same memory location as b, the second cout statement prints that the value of b is 64.

Code for Recursive Newton's Algorithm

Let us now try to use the C++ concepts we have just introduced to implement Newton's recursive algorithm. To accomplish this, we first need to look at how to partition the problem. The first thing we observe from the mathematical description of section 3.1.1 is that we will need to generate an array of doubles, which will contain the differencing coefficients a_k from the formulation given previously. First, we need to have:

- 1. The number of interpolation points.
- 2. A grid of points at which the interpolation is to be exact.
- 3. An array containing the function we wish to interpolate evaluated at the interpolating grid.
- 4. An array to store the Newton differencing coefficients a_k .



Let us take the top-down approach. First, assume that we have the four items enumerated above. The following function encapsulates the calculation of the differencing coefficients:

```
void NewtonDiffTable(int npts, double *xpts, double *funcvals,
  double * newton_coeffs){
  int i,j;
  for(i=0;i<npts;i++)
    newton_coeffs[i] = NewtonDiffFunction(0,i, xpts, funcvals);
}</pre>
```

In this function, npts is the number of interpolating points, xpts is an array containing the interpolating grid, funcvals is an array containing the function we wish to interpolate evaluated at the interpolating grid. For each coefficient a_i (contained within the array element $newton_coeffs[i]$), we call the following NewtonDiffFunction:

```
double NewtonDiffFunction(int start_index, int ending_index,
  double * xpts, double * funcvals){
  double val;
  int diff = ending_index-start_index;
  if(diff == 0){
    val = funcvals[start_index];
  }
  else{
    val = (NewtonDiffFunction(start_index,ending_index-1,
                              xpts,funcvals) -
  NewtonDiffFunction(start_index+1,ending_index,
                              xpts,funcvals))/
           (xpts[start_index]-xpts[ending_index]);
 }
  return val;
}
```

As input, this function takes a starting index and an ending index (both of which are assumed to be in between zero and (npts - 1)), the interpolating grid, and the function to be interpolated evaluated on that grid. What is different about this function? It calls itself! This is a powerful concept which can be used in C++, i.e. the concept of recursive function calling. In the function above, notice that we are able to replicate in C++ code the mathematical recursive relation given in equation (3.3).

In order for the recursion to be effective, we must have two things:

- 1. A recursive relationship, and
- 2. A stopping condition.

Key Concept

• Recursive functions require two things: A recursive definition and a stopping condition.

The first item seems quite obvious; recursion requires a recursive definition. The second item, although it sounds trivial, is often the stumbling block – often we do not know when the recursive relationship ends. In the example code above, the recursive definition is given

in equation (3.3) and the stopping condition is determined by the condition that the starting index be the same as the ending index. From our mathematical definitions above, we know that when the starting index and ending index are the same, then the Newton formula gives back the value of the function evaluated at that index value.



Once we have successfully calculated the Newton divided differences, we can now implement our Newton interpolating polynomial as follows:

```
double NewtonInterpolant(double x, int npts, double * xpts,
  double * newton_coeffs){
  int i,j;
  double sum = 0.0, xval;

  for(i=0;i<npts;i++){
    xval = 1.0;
    for(j=0;j<i;j++)
        xval = xval*(x-xpts[j]);
    sum = sum + newton_coeffs[i]*xval;
  }

  return sum;
}</pre>
```

As input, this function takes a value x, which is the value at which we want to know the value of the interpolating polynomial, the number of interpolating points npts, an array of interpolating points xpts, and the array of previously calculated Newton divided differences. As output, this function gives the value of the interpolating polynomial evaluated at the point x. Observe that this code replicates the mathematical definition given in equation (3.1).

We will now use the previously defined functions in a program. Below, we have provided a program which interpolates the *Runge function*

$$f(x) = \frac{1}{1 + 25x^2}, \quad x \in [-1, 1]. \tag{3.4}$$



This code queries the user for the degree of the interpolating polynomial, and prints on the screen the values of the interpolating polynomial evaluated at 1000 evenly spaced points on the interval [-1,1]. This example demonstrates the use of output/input with cout/cin respectively, dynamic memory allocation, and recursive functions (through the calling of the NewtonDiffTable function). In figure 3.2, we plot the results of running the program when the degree of the interpolating polynomial is set to five.

```
#include <iostream.h>
#include "SCchapter3.h"
double func(double x);
int main(int argc, char * argv[]){
  int i;
  int degree, polypnts;
  int npts = 1000;
                        //number of points used for plotting
  double xpt, soln, approx;
  cout << "Enter the degree of the interpolating polynomial: ";</pre>
  cin >> degree;
  polypnts = degree+1; //number of points is
                             equal to 1 + degree
  double * poly_xpts = new double[polypnts];
  double * func_vals = new double[polypnts];
  double * newton_coeffs = new double[polypnts];
  CreateGrid_EvenlySpaced(polypnts, poly_xpts, -1.0, 1.0);
  for(i=0;i<polypnts;i++){</pre>
    func_vals[i] = func(poly_xpts[i]);
  }
  NewtonDiffTable(polypnts, poly_xpts, func_vals,
                  newton_coeffs);
```

```
for(i=0;i<npts;i++){</pre>
    xpt = -1.0 + 2.0*i/(npts-1);
    soln = func(xpt);
    approx = NewtonInterpolant(xpt, polypnts,
                                  poly_xpts, newton_coeffs);
    cout << xpt << " " << soln << " " << approx << endl;</pre>
  }
  delete[] poly_xpts;
  delete[] func_vals;
  delete[] newton_coeffs;
}
double func(double x){
  double y;
  y = 1.0 + 25.0*x*x;
  y = 1.0/y;
  return y;
}
```

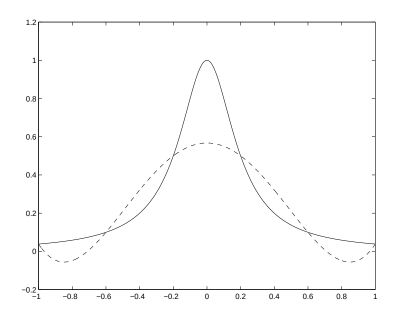


Figure 3.2: Comparison of the solution (solid) and fifth-order interpolation polynomial (dashed) obtained using the Newton differencing program.

We will conclude this section by making some observations concerning the code above.

- Notice in the program above that we have variable declarations after executable statements. This is one major difference between the programming language C (and many other programming languages) and C++. In many languages, all declarations must be made prior to executable statements. This is not true in C++. In C++, variables may be declared at any point within the program. Observe that we declare three pointer variables (double * variables) following the input sequence. Our primary reason for often placing all the variable declarations at the beginning of a function is clarity.
- In chapter 2 we mentioned that in C++ it is possible to initialize a variable with an executable statement and not just a constant. This is exactly what we have done in the program above. Observe that we initialize each new double* variable (with which the previous remark was dealing) with the results of the *new* operator.
- For each item we want to delete, we must execute an individual delete[] statement. You *cannot* combine them like variable declaration in the following manner:

delete[] poly_xpts, func_vals, newton_coeffs;

Though the compiler will not complain about the statement given above, only the last variable will have the valid operation done to it. Both **delete** and **delete**[] take only one argument.

3.1.3 Lagrangian Interpolation

Another basis which is often used in practice to interpolate data primarily on non-equidistant grids is Lagrangian interpolation. Here, the basis $\phi_k(x)$ is equal to the Lagrangian polynomial which is defined by

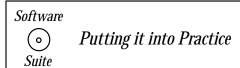
$$h_k(x) = \Pi\left(\frac{(x-x_i)}{(x_k-x_i)}\right), \quad i = 0, 1, \dots N; \quad i \neq k.$$
 (3.5)

This is a polynomial of degree N and satisfies the equation

$$h_i(x_j) = \delta_{ij},$$

where δ_{ij} is the Kronecker delta

$$\delta_{ij} = \left\{ \begin{array}{l} 1, i = j \\ 0, i \neq j \end{array} \right..$$



The following code implements this definition and returns values of the Lagrange polynomial.

```
double LagrangePoly(double x, int pt, int npts, double * xpts){
  int i;
  double h=1.0;

  for(i=0;i<pt;i++)
     h = h * (x - xpts[i])/(xpts[pt]-xpts[i]);

  for(i=pt+1;i<npts;i++)
     h = h * (x - xpts[i])/(xpts[pt]-xpts[i]);

  return h;
}</pre>
```

This code is relatively simple, yet there is one point to which we would like to draw your attention. Notice in the mathematical definition of Lagrange polymonials there is the condition $i \neq j$, which in the above code translates into $i \neq j$. This can be implemented in one of two ways:

- 1. As above, by breaking the sum into two parts, and having a separate for loops for each part.
- 2. Having one for loop ranging from $0 \le i < npts$, in which inside the for loop there is an if statement which checks if the value of the looping variable is the same as the value of pt.

Thus, the alternative function definition is the following:

```
double LagrangePoly(double x, int pt, int npts, double * xpts){
  int i;
  double h=1.0;

  for(i=0;i<npts;i++)
     if(i!=pt)
        h = h * (x - xpts[i])/(xpts[pt]-xpts[i]);

  return h;
}</pre>
```

These two definitions accomplish the same objective; however, their efficiency may be different. The reason for the efficiency difference comes from the repeated use of the *if* statement. The *if* statement is, in general, a very difficult computer instruction to optimize because the value of the boolean expression is not known ahead of time, and hence which branch of the *if* the computer will have to execute is uncertain until the boolean expression

is evaluated. Hence, avoiding unnecessary if statements is, in general, a good policy because it allows the compiler to accomplish more code optimization.²

Key Concept

• A good rule of thumb: Avoid *if* statements within *for* loops.

Returning to the mathematics, by definition, then, we have that

$$f(x) = \sum_{k} a_k \phi_x(x)$$
 or
 $f(x) = \sum_{k} f(x_k) h_k(x),$

which in C++ is implemented as follows:

```
double LagrangeInterpolant(double x, int npts, double *xpts,
   double * funcvals){
   int i;
   double sum = 0.0;

   for(i=0;i<npts;i++){
      sum = sum + funcvals[i]*LagrangePoly(x,i,npts,xpts);
   }
   return sum;
}</pre>
```

We can reuse the program presented for the Newton divided differences (previous section) to now do Lagrange interpolation. First, we can remove the declaration of the Newton Differencing coefficients, and we can also remove the call to the function NewtonDiffTable. Now, instead of calling NewtonInterpolant(...), we will call LagrangeInterpolant(...) with its appropriate arguments. We have extracted the relevant code from the program presented earlier, and now present a modified version of the code. The key thing to observe is that by writing modular code, we have increased code reuseability.

```
double * poly_xpts = new double[polypnts];
double * func_vals = new double[polypnts];
```

²In recent years, there has been considerable work in processor design to incorporate what is referred to as "branch prediction." During runtime, the processor monitors the behavior of branches, and attempts to predict which branch of the *if* will be taken. By doing so (i.e., if it predicts correctly) it can pipeline instructions with no additional cost due to the branching statement. If it is wrong, it will incur the normal branching penalty (in terms of pipelining).

Key Concept

• Modular code is reusable code!

The coefficients in this polynomial representation are the function values at the grid points. Therefore, this approach is very efficient, especially for a "static" grid, as the Lagrangian polynomials can be constructed once, stored, and be used repeatedly. However, in a dynamic grid where the grid points $\{x_i\}$ change, we need to recompute *all* Lagrangian polynomials, and this may be costly.

3.1.4 The Runge Phenomenon

Up to this point, we have assumed that all polynomial interpolation formulas converge irrespective of the distribution of points x_i , i.e., uniform with equidistant spacing or non-uniform with arbitrary spacing. We have to wonder, however, if the type of function we try to approximate matters? Usually, polynomial approximation in the neighborhood of a point is safe, but as we try to extend this interpolation in the large, i.e., away from the given data, proper conditions have to be satisfied, as we will illustrate in this section. First, we state a fundamental theorem that justifies polynomial approximation.

Weierstrass Theorem (1885): Any continuous function defined in a finite range can be approximated to any degree of accuracy by polynomial powers.

While this theorem is re-assuring it does not suggest what is the appropriate type of polynomial approximation. Specifically, the question still remains if equidistant data always produce stable and convergent polynomial approximations. Experience shows that the answer to this question is negative, and in fact it was O. Runge in 1901, who first discovered this troubling fact by considering the function

$$f(x) = \frac{1}{1 + 25x^2}, \quad x \in [-1, 1]. \tag{3.6}$$

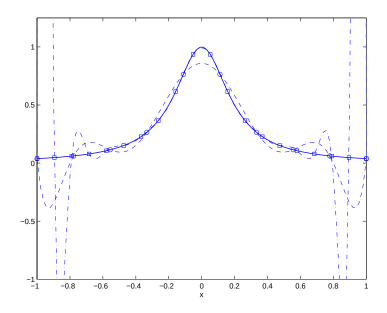


Figure 3.3: Plot of the Runge function (equation (3.4); solid line) and approximations using 10 equidistant points (dashed line) and 20 equidistant points (dashed-dot line).

In figure 3.3 we plot this function along with a polynomial approximation on ten and twenty equidistant points. We observe that the approximation is accurate around the origin but at $x \ge \pm 0.72$ the interpolating polynomial does not converge; taking more than ten points makes the oscillations worse. On the other hand, by interpolating the Runge function at special non-equidistant points, obtained from

$$x_k = cos(k\pi/N), i = 0, \dots, N$$

we obtain a stable polynomial approximation, which converges fast to the exact solution as the number of grid points increases; this is shown in figure 3.4. These special points are the roots of the derivatives of Chebyshev polynomials, which we will study in the next section. Other type of approximations, such as trigonometric interpolation, may also be stable, see section 3.2.4.

The question then remains for equidistant polynomial interpolation how can we know in advance which functions have stable polynomial approximations? We know that for Taylor type expansions this question relates to the analyticity of a function in the complex plane, and we are perhaps familiar with the Taylor circle inside which there are no singularities. The same picture emerges here also except that the circle is replaced by an oval shape

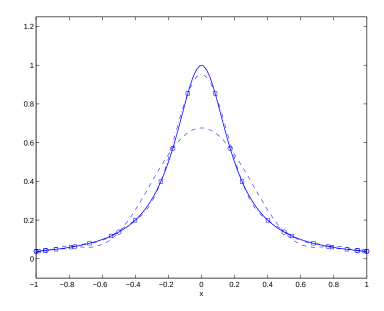


Figure 3.4: Plot of the Runge function (equation (3.4); solid line) and approximations using 10 points (dashed line) and 20 points (dashed-dot line) following the *cosine-law* distribution of grid points.

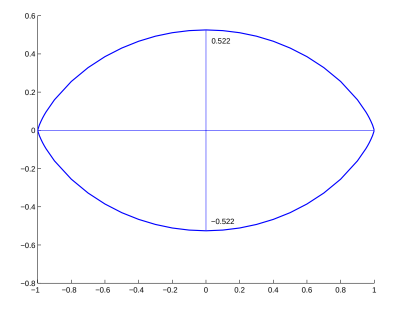


Figure 3.5: Region of required analyticity for a function for stable equidistant polynomial interpolation.

region, which is shown in figure 3.5. The proof is rather elaborate and the interested reader is referred to the book by Lanczos [66].

Theorem: The necessary and sufficient condition for convergence of equidistant polynomial interpolation is that a function not have any singularities in the oval region Ω corresponding to the definition interval of f(x) with $x \in [-1, 1]$.

The Runge function of equation (3.4) has singularities at $z = \pm i/5$, and this is the reason for the unstable behavior close to the end-points. The more general Runge function

$$f(x) = \frac{\epsilon^2}{\epsilon^2 + x^2}, \quad x \in [-1, 1]$$

has a singularity at $x = \pm i\epsilon$. This is a very tough function to approximate, as at x = 0 we have f(0) = 1 but at distance only ϵ away we have $f(\epsilon) = \frac{1}{2}$, independent of the value of ϵ . You can appreciate what happens when $\epsilon = 10^{-6}$!

3.1.5 Chebyshev Polynomials

Spectral approximations and more specifically polynomial approximations using Chebyshev polynomials are a very effective means of representing relatively smooth data and also numerical solutions of partial differential equations. Just as before, we can write our polynomial approximation $p_N(x)$ as truncated series of the form

$$f(x) \approx p_N(x) = \sum_{k=0}^{N} a_k T_k(x),$$

where $T_k(x)$ is the k^{th} Chebyshev polynomial. The Chebyshev polynomial series converges very fast; the polynomials are determined from recursion relations such as:

$$T_0(x) = 1;$$
 $T_1(x) = x;$ $T_{n+1}(x) = 2xT_n(x) - T_{n-1}(x),$ $n \ge 1.$ (3.7)



The following code implements this recursive formula; plots of $T_k(x)$, k = 0, 1, 2, 3, 4 are shown in figure 3.6.

double ChebyshevPoly(int degree, double x){
 double value;

```
switch(degree){
case 0:
  value = 1.0;
  break;
case 1:
  value = x;
```

In this example, there are two things that we want to point out. First, notice that for this particular example, we have *two* explicit stopping conditions: when k = 0 and when k = 1. This is because our recurrence relation contains references to (k-1) and (k-2), and hence we need both to be valid in order to get the k^{th} term. The second thing to observe in this example is the use of a not previously mentioned C++ statement, the **switch** statement.

The SWITCH Statement

The switch statement is a convenient way of replacing a collection of *if-else* statements. In the example above, we have the following logic: If the value of degree is 0, then return the number 1.0, else if the value of degree is 1, return the value of x, else the value is what is given by the recurrence relation. We could implement this using *if-else* statements as follows:

However, C++ has a statement named **switch** which accomplishes this type of logic for you. The syntax of a switch statement is as follows:

```
switch( variable ){

case a:
    statement 1;
    break;

case b:
    statement 2;
    statement 3;
    break;
```

```
default:
    statement 4;
}
```

Here, 'variable' is the variable that you want to test; 'a', 'b', etc. are the *constant* values that you want to test against (these cannot be variables, they must be constants). When the *switch* is executed, the first *case* in which the variable is equivalent to the case is where execution begins. All statements below that *case* statement are executed. Hence, when all the statements that you want done for a particular case have been executed, you must use a **break** statement to exit the switch. The *default* case is the case that is executed if no other cases have matched the variable.

WARNING

Programmer Beware!

• Do not forget to put break statements between independent cases!

Because of the "flow-through" nature of the switch statement, one can group cases together. Suppose, for example, that we wanted to implement a *statement* which executes statement 1 if cases 0 and 1 are true, and statement 2 otherwise. The following pseudo-code example demonstrates the implementation of this logic:

```
switch(degree){
   case 0:
   case 1:
       statement 1;
      break;
   default:
      statement 2;
}
```

In this example, if either case 0 or case 1 is true, then statement 1 (and only statement 1, due to the *break* statement) will be executed. For all other values of the variable degree, statement 2 will be executed.

Key Concept

• **Switch** is a nice organizational tool for implementing *if-else* relationships.

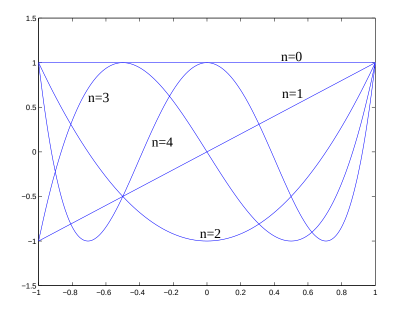


Figure 3.6: Chebyshev polynomials of order 0 through 4 in the interval [-1, 1].

Properties of Chebyshev Polynomials

Next, we summarize some important properties of the Chebyshev polynomials:

- Symmetry: $T_n(-x) = (-1)^n T_n(x)$.
- The leading coefficient is $2^{n-1}, n \ge 1$.
- **Zeros**: The roots of $T_n(x)$ are $x_k = \cos\left(\frac{2k+1}{n} \cdot \frac{\pi}{2}\right)$, $k = 0, 1, \dots, n-1$. These are called Gauss points and we will use them later in numerical integration. The roots of its derivative $T'_k(x)$, which are the locations of extrema for $T_k(x)$, are the Gauss-Lobatto points and are given by $x'_k = \cos\frac{k\pi}{n}$. We also have that

$$T_n(x'_k) = (-1)^k, \quad k = 0, 1, 2, \dots$$

• Orthogonality in the continuous inner product is:

$$\int_{-1}^{1} T_i T_j \frac{dx}{\sqrt{(1-x^2)}} = (T_i, T_j) = \begin{cases} 0, & i \neq j \\ \pi/2, & i = j \neq 0 \\ \pi, & i = j = 0 \end{cases}$$

We often use orthogonality in the discrete inner product:

$$\sum_{k=0}^{m} T_i(x_k) T_j(x_k) = \begin{cases} 0, & i \neq j \\ \frac{m+1}{2}, & i = j \neq 0 \\ m+1, & i = j \end{cases}$$

ullet Lagrangian Interpolant: The Chebyshev Lagrangian interpolant through N Gauss points has a simple form:

$$h_k(x) = \frac{T_N(x)}{T'_N(x_k)(x - x_k)}, \quad x \neq x_k.$$

• Grid Transformation: The following grid transformation maps the Gauss-Lobatto points $x_k = \cos(k\pi/N)$, k = 0, ..., N to a new set of grid points ξ_k obtained from:

$$\xi_k = \frac{\sin^{-1}(\alpha x_k)}{\sin^{-1}(\alpha)},\tag{3.8}$$

where $\alpha \in (0,1]$ defines the exact distribution. For $\alpha \to 0$ the new points are equidistant and the Chebyshev approximation resembles the Fourier method. However, for stability of the approximation the new points cannot be exactly equidistant, and thus $\alpha > 0$.

• MiniMax Property: Of all the n^{th} -degree polynomials with leading coefficient 1, the polynomial $2^{1-n}T_n(x)$ has the smallest maximum norm in the interval [-1,1]. The value of its maximum norm is 2^{1-n} .

Approximation Error and Convergence Rate

Let us assume that we are given the values of a function f(x) on a grid of (m+1) points, and we use a polynomial p(x) to represent the data on this grid. The error (or remainder r(x)) in the approximation of a function f(x) is then

$$|f(x) - p(x)| = \frac{(x - x_0)(x - x_1) \dots (x - x_m)|f^{(m+1)}(\xi)|}{(m+1)!},$$

where $\xi \in [x_0, x_m]$. This error behaves like the polynomial curve

$$r(x) \sim (x - x_0)(x - x_1)(x - x_2) \dots (x - x_m),$$

which oscillates, similar in fact to the least-squares approximation (section 3.1.7), and unlike the Taylor expansion approximation where the error increases exponentially as $\sim (x-x_0)^{m+1}$.

Now, we can attempt to find the optimum distribution of the grid points, which means that we seek to minimize the maximum magnitude of

$$q(x) \equiv (m+1)! p(x) = (x-x_0)(x-x_1) \dots (x-x_m).$$

To this end, we can use the minimax property to obtain

$$q(x) = 2^{-m} T_{m+1}(x),$$

thus $\{x_k\}$ are the zeros of the Chebyshev polynomial $T_{m+1}(x)$, and thus the grid points x_k are the roots of $T_{m+1}(x)$, i.e.,

$$x_k = \cos\left(\frac{2k+1}{m+1}\frac{\pi}{L}\right), k = 0, 1, \dots m$$

We now state Rivlin's minimax error theorem:

MiniMax Error Theorem: The maximum pointwise error of a Chebyshev series expansion that represents an arbitrary function f(x) is only a small constant away from the minimax error, i.e., the smallest possible pointwise error of any N^{th} degree polynomial. The following inequality applies:

$$\| f(x) - \sum_{k=0}^{N} a_k T_k(x) \|_{\infty} \le 4 \left(1 + \frac{\ln N}{\pi^2} \right) \cdot \| f(x) - mm(x) \|_{\infty},$$

where mm(x) is the best possible polynomial.

Note that for N=128 the prefactor is less than 5 and for N=2,688,000 the pre-factor is $4\left(1+\frac{\ln N}{\pi^2}\right)\approx 10$. Therefore, the Chebyshev expansion series is within a decimal point of the minimax approximation.

The convergence of Chebyshev polynomial expansions is similar to Fourier cosine series, as the following transformation applies

$$x = \cos \theta$$
 and $T_n(\cos \theta) = \cos(n\theta)$.

Assuming an infinite expansion of the form

$$f(x) = \sum_{k=0}^{\infty} a_k T_x(x)$$

then

$$a_k = \frac{1}{\pi c_k} \int_0^{\pi} f(\cos \theta) \cos k\theta d\theta = \frac{2}{\pi c_k} \int_{-1}^{1} f(x) T_k(x) \frac{dx}{\sqrt{1 - x^2}},$$

where we have defined

$$c_k = \begin{cases} 2, k = 0 \\ 1, k > 0 \end{cases}$$

The convergence rate of the expansion series is defined by the decaying rate of the coefficients a_k . To this end, if:

- $f^{(p)}(x)$ is continuous $\forall |x| \leq 1$, $p = 0, 1, 2, \dots, n 1$, and
- $f^{(n)}(x)$ is integrable

then

$$a_k \ll \frac{1}{k^n}$$
.

This implies that for infinitely differentiable functions the convergence rate is extremely fast. This convergence is called *exponential*, and it simply means that if we double the number of grid points the approximation error will decrease by a two orders of magnitude (i.e., a factor of 100), instead of a factor of four which will correspond to interpolation with quadratic polynomials and second-order convergence rate. The above estimate also shows that in the Chebyshev approximation we can exploit the regularity, i.e., *smoothness*, of the function to accelerate the convergence rate of the expansion. Also, notice that unlike the

Fourier series (see section 3.2), the convergence of Chebyshev series does not depend on the values of f(x) at the end points, because the boundary terms vanish automatically.

Finally, an important consequence of the rapid convergence of Chebyshev polynomial expansions of smooth functions is that they can be differentiated normally term-wise, i.e.,

$$\frac{d^p f(x)}{dx^p} = \sum_{k=0}^{\infty} a_k \frac{d^p T_k(x)}{dx^p}.$$

In computing Chebyshev derivatives higher than the first, inaccurate results may be obtained due to round-off. In particular, it has been observed that round-off may be significant for the second derivative for N > 128, for the third derivative for N > 64, and for the fourth derivative for N > 32. This round-off can be reduced if the grid transformation given by equation (3.8) is employed.

Example: The following example, first presented in Gottlieb & Orszag [49], shows the fast convergence of Chebyshev discretization. The exact representation for the sine function corresponding to wave number M is

$$\sin M\pi(x+\alpha) = 2\sum_{n=0}^{\infty} \frac{1}{c_n} J_n(M\pi) \sin(M\pi\alpha + \frac{1}{2}n\pi) T_n(x),$$

where $J_n(x)$ is the Bessel function of order n. We can argue that $J_n(M\pi) \to 0$ exponentially fast for $n > M\pi$, given that the Bessel function can be approximated by

$$J_n(M\pi) \approx \frac{1}{\sqrt{2\pi n}} \left(e \underbrace{\frac{M\pi}{2n}}_{\leq 1} \right)^n \quad \text{if} \quad \frac{n}{M} > \pi.$$

This result leads to the following heuristic rule for Chebyshev series approximation, proposed by Gottlieb & Orszag:

Quasi-sinusoidal rule-of-thumb: In order to resolve M complete waves it is required that $M\pi$ modes be retained, or in other words π polynomials should be retained per wavelength.

Although very good, such a resolution capability is less than that of a Fourier method that requires approximately two points per wave! In fact, the above is an asymptotic result, and a more practical rule for the total number of points N is:

$$N = 6 + 4(M - 1),$$

which has been verified in many numerical experiments.

3.1.6 Hermite Interpolation and Splines

We now turn to piecewise polynomial interpolation using relatively low-order polynomials, unlike the single domain global interpolation. As we know from our previous discussion, the more (distinct) grid points that we introduce into the polynomial approximation, the higher the order of the interpolating polynomial. As the degree of the polynomial increases,

the interpolating polynomial becomes more oscillatory. One solution to this problem is to use multiple piecewise polynomials of low-order instead of one high-order polynomial. This has the advantage that the interpolation error is proportional to a lower order derivative resulting in better accuracy and more flexibility. The method of splines, first used in naval architecture, is a very effective way of such an approach and facilitates smooth transition between subdomains.

To proceed, let us first consider cubic *Hermite interpolation* where both function and derivatives are interpolated. The interpolation problem can be stated as follows:

• Given data for the function values y and slopes s, $(x_L; y_L; s_L)$ and $(x_R; y_R; s_L)$ at the left (x_L) and right (x_R) boundaries of one-dimensional domain, find a cubic polynomial

$$p(x) = a_0 + a_1(x - x_L) + a_2(x - x_L)^2 + a_3(x - x_L)^2(x - x_R)$$

with the four unknown coefficients obtained from

$$p(x_L) = y_L, \qquad p(x_R) = y_R$$

$$p'(x_L) = s_L, \qquad p'(x_R) = s_R.$$

The first derivative is

$$p'(x) = a_1 + 2a_2(x - x_L) + a_3[2(x - x_L)(x - x_R) + (x - x_L)^2]$$

and by substituting the known data at the boundaries, we have

$$p(x_L) = y_L = a_0;$$
 $a_0 + a_1 \Delta x + a_2 (\Delta x)^2 = y_R$

$$p'(x_L) = s_L = a_1;$$
 $a_1 + 2a_2\Delta x + a_3(\Delta x)^2 = s_R,$

where $\Delta x \equiv x_R - x_L$ is the domain size. We recast the above in matrix-vector form as follows

$$\begin{bmatrix} 1 & & & O \\ 0 & 1 & & \\ 1 & \Delta x & \Delta x^2 & \\ 0 & 1 & 2\Delta x & \Delta x^2 \end{bmatrix} \begin{bmatrix} a_0 \\ a_1 \\ a_2 \\ a_3 \end{bmatrix} = \begin{bmatrix} y_L \\ s_L \\ y_R \\ s_R \end{bmatrix}$$

and we see that the coefficient matrix is lower-triangular.

The solution is obtained by forward substitution:

$$a_0 = y_L, \qquad a_2 = y_L''$$

$$a_1 = s_L, \qquad a_3 = y_L'''$$

where we have defined the forward differences at the left boundary as:

$$y'_{L} = \frac{y_{R} - y_{L}}{\Delta x}, \quad y''_{L} = \frac{y'_{L} - s_{L}}{\Delta x}, \quad y'''_{L} = \frac{s_{L} - 2y'_{L} + s_{R}}{\Delta x^{2}}.$$

Theorem on Accuracy of Hermite Interpolation: For a function f(x) interpolated by a cubic Hermite polynomial p(x) in a domain of size Δx , the error is bounded from above by

$$|| f(x) - p(x) ||_{\infty} \le \frac{|| f^{(4)}(x) ||_{\infty}}{384} (\Delta x)^4.$$

This theorem guarantees that if the domain is divided into 100 subintervals then the error will be reduced by a factor of 10^8 ! The proof is based on evaluating the *maximum* contribution of the remainder at the mid-point of the domain.

Constructing Cubic Splines: Having obtained simple formulas for Hermite cubic polynomial interpolation, we now proceed to construct formulas for different types of cubic splines. Here we replace the extra information given for first derivatives (s_L, s_R) at the end-points of the domain by imposing continuity at the interior points. We assume, therefore, that the entire domain is subdivided into subdomains or cells and that the function and its derivatives are continuous at the breakup points, which we call the interior points. We can state the problem of constructing splines as follows:

• Given the data points $(x_1, y_1), \ldots (x_n, y_n)$, find a piecewise cubic interpolant S(x) so that S(x), S'(x) and S''(x) are continuous at all interior points x_i , $i = 2, \ldots, (n-1)$.

To achieve this we have to choose the slopes $(s_i, i = 1, ..., n)$ at all points appropriately. To maintain continuity of the slopes we simply assign a single value at each point. However, we have to enforce continuity of the second derivative explicitly. To this end, we consider the subdomain $x \in [x_i, x_{i+1}]$ and apply Hermite interpolation as before where the point x_i is the left boundary and the point x_{i+1} is the right boundary, as shown in figure 3.7.

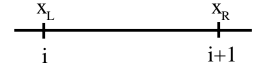


Figure 3.7: Interpolation in the interval $x \in [x_i, x_{i+1}]$.

We then construct the cubic polynomial

$$p_i(x) = y_i + s_i(x - x_i) + y_i''(x - x_i)^2 + y_i'''(x - x_i)^2(x - x_{i+1}),$$

where

$$y'_i = \frac{y_{i+1} - y_i}{\Delta x_i}; \quad y''_i = \frac{y'_i - s_i}{\Delta x_i}; \quad y'''_i = \frac{s_i - 2y'_i + s_{i+1}}{(\Delta x_i)^2},$$

with $\Delta x_i \equiv x_{i+1} - x_i$. We also obtain the second derivative

$$p_i''(x) = 2y_i'' + y_i''' \left[4(x - x_i) + 2(x - x_{i+1}) \right].$$

Next, we move to the adjacent cell $x \in [x_{i+1}, x_{i+2}]$ in order to apply Hermite interpolation on this interval, see figure 3.8.

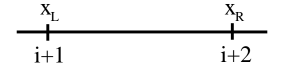


Figure 3.8: Interpolation in the interval $x \in [x_{i+1}, x_{i+2}]$.

We construct the polynomial

$$p_{i+1}(x) = y_{i+1} + s_{i+1}(x - x_{i+1}) + y_{i+1}''(x - x_{i+1})^2 + y_{i+1}'''(x - x_{i+1})^2(x - x_{i+2}),$$

and its second derivative

$$p_{i+1}''(x) = 2y_{i+1}'' + y_{i+1}'''[4(x - x_{i+1}) + 2(x - x_{i+2})].$$

Next, we enforce continuity of second derivative at x_{i+1} and obtain equations for the unknown slopes:

$$\frac{2}{\Delta x_i}(2s_{i+1} + s_i - 3y_i') = \frac{2}{\Delta x_{i+1}}(3y_{i+1}' - 2s_{i+1} - s_{i+2}),$$

which can be re-written as

$$\Delta x_{i+1}s_i + 2(\Delta x_i + \Delta x_{i+1})s_{i+1} + \Delta x_i s_{i+2} = 3(\Delta x_{i+1}y_i' + \Delta x_i y_{i+1}')$$

$$i = 1, \dots, n-2.$$
(3.9)

The above equations can be recast in a matrix-vector form with a tridiagonal coefficient matrix. However, we have no information for the slopes s_1, s_n (the end-slopes), and thus we cannot solve for the slopes yet; we need additional information, the type of which specifies different classes of splines:

- I. The Complete Spline: Here the additional information is given at the end-points where the end-slopes are explicitly specified, i.e., $s_1 = S_L$ and $s_n = S_R$.
- II. The Natural Spline: In this case the *curvature* (i.e., p''(x)) at the end-points is set to zero, i.e.

$$p_1''(x_L) = 0 \Rightarrow s_1 = \frac{1}{2}(3y_1' - s_2)$$
 and $s_n = \frac{1}{2}(3y_{n-1}' - s_{n-1}).$

• III. The Not-A-Knot Spline: In this case, no explicit information on the end-points is provided, but instead we use continuity of the third-derivative p'''(x) at the points x_2 and x_{n-1} . Using

$$p_i'''(x) = 6\frac{s_i + s_{i+1} - 2y_i'}{(\Delta x_i)^2},$$

and enforcing the continuity condition, we obtain

$$s_1 = -s_2 + 2y_1' + \left(\frac{\Delta x_1}{\Delta x_2}\right)^2 (s_2 + s_3 - 2y_2').$$

This forms the first row in the triagonal matrix-vector system, which now has a bandwidth of *two* instead of *one*. A similar equation is valid for the other end (last row in the matrix-vector system).

• IV. The B-Spline: An example of a very popular cubic spline derived from the above Hermite interpolation is the *basic* or *B-spline*. It corresponds to zero slopes at the endpoints, i.e., $s_L = s_R = 0$, and it is symmetric. Its support is five points, that is, it is non-zero within four equidistant intervals Δx . It is defined by the five points

$$(x_i, y_i) = [(1,0), (1,1), (2,4), (3,1), (4,0)].$$

Solving the matrix-vector system constructed from equation (3.9):

$$\begin{bmatrix} 2 & 1 & & & \\ 1 & 4 & 1 & & \\ & 1 & 4 & 1 & \\ & & 1 & 4 & 1 \\ & & & 1 & 2 \end{bmatrix} \begin{bmatrix} s_1 \\ s_2 \\ s_3 \\ s_4 \\ s_5 \end{bmatrix} = 3 \begin{bmatrix} 1 \\ 4 \\ 0 \\ -4 \\ -1 \end{bmatrix}$$

we obtain

$$B(x) \equiv S(x) = \begin{cases} x^3, 0 \le x \le 1 \\ 4 - 6(2 - x)^2 + 3(2 - x)^3, 1 \le x \le 2 \\ 4 - 6(2 - x)^2 - 3(2 - x)^3, 2 \le x \le 3 \\ (4 - x)^3, 3 \le x \le 4, \end{cases}$$

and it is zero everywhere else as shown in figure 3.9. Note that the B-spline, which has its origin in applications of beam vibrations, satisfies a *minimum principle*. Specifically, the B-spline has the smallest bending energy

$$E(B(x)) = \int_0^N \left(\frac{d^2B}{dx^2}\right)^2 dx,$$

among all functions or other splines that go through the same data points.

Next, we state a theorem that gives upper bounds for the error of spline interpolation up to the third derivative. This depends on the end-points, so the theorem is for the *complete spline*:

Theorem on Accuracy of Complete Splines: Let S(x) be the cubic spline that interpolates f(x), $x \in [a,b]$ at points $x_i = i\Delta x$, where $\Delta x = \frac{(b-a)}{n}$, $i = 0, \ldots, n$, and also $s_L = f'(a)$ and $s_R = f'(b)$, then:

$$|| S^{(r)}(x) - f^{(r)}(x) ||_2 \le \epsilon_r || f^{(4)} ||_2 (\Delta x)^{4-r}, \quad r = 0, 1, 2, 3$$

where $\epsilon_0 = \frac{5}{384}, \epsilon_1 = \frac{1}{24}, \epsilon_2 = \frac{3}{8}$ and $\epsilon_3 = 1$.

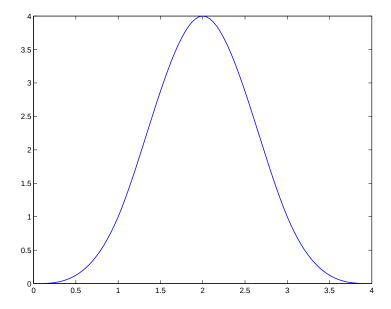


Figure 3.9: Plot of the B-spline. Its support extends over four intervals.

3.1.7 Least-Squares Approximation

What we have dealt with so far are interpolations for which the number of unknowns matches the number of constraints in the function values or slopes. The cubic splines are perhaps a slight exception, as extra information is needed to determine them, but that too could be cast in a form of a linear system

$$Ca = y$$
,

where \mathbf{a} is the vector of unknowns and \mathbf{y} is the vector of prescribed values.

In practical scientific computing, however, the opposite situation may occur where we may have much *more information* than we actually require. For example, consider the case of data analysis in an experiment with many measurements and with the dependent variable following a quadratic trend. It is clear that, unless we are really lucky, not all measurements will lie on a parabola! It is also clear that *not* all measurements have the same confidence associated with them, and this should be reflected in the polynomial interpolant. The question then is what type of parabola fits best the data and has built-in the measurements-accuracy reported.

A similar situation arises in the problem of smoothing the data, a familiar task in engineering, and Gauss was the first to provide a general treatment of this subject. He introduced the $bracket \ notation$ which we will use here, i.e., for a set of m grid points we define

$$[y] \equiv y_1 + \ldots + y_m$$

and the moments

$$[wx^k] = w_1x_1^k + w_2x_2^k + \ldots + w_mx_m^k$$
.

To obtain the least-squares polynomial, we assume that we have m points and the pairs (x_i, y_i) , i = 1, ..., m, and we try to represent them with a polynomial p(x) of degree n

$$p(x) = a_0 + a_1 x + a_2 x^2 + \ldots + a_n x^n$$

where $m \ge n + 1$. The strict inequality corresponds to an over-determined system that we discuss in this section. In matrix-vector form, if we follow the straightforward path, as in the Vandermonde approach (section 3.1.1), we have that

$$\mathbf{C}(m \times (n+1)) \times \mathbf{a}(n+1) = \mathbf{y}(m)$$

so the matrix C is rectangular, and for m > (n + 1) this system does not always have a solution.

However, there is a minimum principle associated with this problem. That is, instead of solving the system $\mathbf{C} \mathbf{a} = \mathbf{y}$ we attempt to find the vector \mathbf{a} such that we minimize the residual

$$r(\mathbf{a}) = \parallel \mathbf{C} \mathbf{a} - \mathbf{y} \parallel_2$$
.

This results in solving a system of the form

$$\mathbf{C}^T \mathbf{C} \ \mathbf{a} = \mathbf{C}^T \mathbf{y},\tag{3.10}$$

where $\mathbf{C}^T\mathbf{C}$ is a symmetric positive definite matrix if \mathbf{C} has linearly independent columns.

This may not be obvious at first but following Gauss, we can derive the *normal equations* that result in the matrix-vector form of equation (3.10). To this end, we compute the residual

$$r_i = a_0 + a_1 x_i + \ldots + a_n x_i^n - y_i$$

at each measurement point x_i , i = 1, ..., m. We now form the sum of the square of all the m residuals

$$R(a_0, a_1, \dots, a_n) = \sum_{i=1}^{m} w_i r_i^2,$$

where w_i is a weight that reflects *confidence* in the accuracy of measurement at point x_i . The next step is to minimize the residual R by taking its derivatives with respect to all a_i and setting them to zero, i.e.,

$$\frac{\partial R}{\partial a_i} = 0 \quad \text{for} \quad i = 1, \dots, m.$$

Using the Gauss notation, this yields

$$a_0[wx^0] + a_1[wx^1] + \dots + a_n[wx^n] = [wy]$$

$$a_0[wx^1] + a_1[wx^2] + \dots + a_n[wx^{n+1}] = [wxy]$$

$$\vdots$$

$$a_0[wx^n] + a_1[wx^{n+1}] + \dots + a_n[wx^{2n}] = [wx^ny].$$

We also note that the unknowns are (a_0, a_1, \ldots, a_n) and that the coefficient matrix **H** is:

$$\mathbf{H} = \begin{bmatrix} [wx^0] & [wx^1] & \dots & [wx^n] \\ [wx^1] & [wx^2] & \dots & [wx^{n+1}] \\ \vdots & \vdots & & & \\ [wx^n] & [wx^{n+1}] & \dots & [wx^{2n}] \end{bmatrix}$$

which is an $(n+1) \times (n+1)$ matrix, the so-called *Hankel matrix*, with its cross-diagonals constant. Also, if we form

$$\mathbf{W} = \begin{bmatrix} w_1 & & & \\ & w_2 & 0 & \\ 0 & & \ddots & \\ & & & w_m \end{bmatrix}$$

the matrix of weights, then we can write the Hankel matrix as

$$\mathbf{H} = \mathbf{V}^T \mathbf{W} \mathbf{V}$$

where

$$\mathbf{V} = \begin{bmatrix} 1 & x_1 & \dots & x_1^n \\ 1 & x_2 & \dots & x_2^n \\ \vdots & \vdots & & & \\ 1 & x_m & \dots & x_m^n \end{bmatrix}$$

is the rectangular Vandermonde matrix; it is non-singular if at least (n + 1) points (out of the total m points) are distinct. The normal equations can be recast in matrix-vector form as

$$(\mathbf{V}^T \mathbf{W} \mathbf{V}) \mathbf{a} = \mathbf{V}^T \mathbf{W} \mathbf{y}$$

where

$$\mathbf{y}^T = (y_1, y_2, \dots, y_m)$$

are the measurements.

The normal equations are, in some sense, a generalization of the Vandermonde approach where the set

$$(1, x, x^2, \dots, x^n)$$

is used as basis $\phi_k(x)$ in the expansion

$$f(x) \approx \sum_{k=0}^{n} a_k \phi_k(x).$$

We can *change the basis*, as we did in the case of deterministic interpolation, in order to arrive at a better algorithm to compute the coefficients $(a_0, a_1, \ldots a_n)$ via *recursion*. To this end, minimization of

$$R(a_0, a_1, \dots a_n) = \sum_{i=1}^{m} w_i [f(x_i) - y_i]^2$$

results in a coefficient matrix $\mathbf{G}((n+1)\times(n+1))$ with elements

$$g_{ij} = \sum_{k=1}^{m} w_k \phi_i(x_k) \phi_j(x_k)$$

for the system Ga = b or

$$g_{ij}a_j = b_i$$
 with $b_i = \sum_{k=1}^m w_k \phi_i(x_k) y_k$.

If this basis is *orthonormal*, i.e.,

$$\sum_{k=1}^{m} w_k \phi_i(x_k) \phi_j(x_k) = \begin{cases} 0, & i \neq j \\ 1, & i = j \end{cases}$$

then the matrix G is a diagonal matrix and the computation of the unknown vector \mathbf{a} becomes a trivial matter. We have seen that the Chebyshev polynomials are orthonormal but their construction requires special Gauss-Lobbato points (see section 3.1.5). Here the key is to discover a similar three-term recurrence formula for a given arbitrary distribution of points $\{x_i\}$. We expect that such a three-term recurrence formula is possible given that the Gram-Schmidt orthogonality procedure of this form for vectors leads to that form, see section 2.2.9; similar results should be expected for polynomials.

Let us call the basis $\phi_k(x) = q_k(x)$ where $q_k(x)$ is the orthonormal polynomial which, by assumption, satisfies the recursion

$$q_{j+1}(x) = xq_j(x) - \alpha_{j+1}q_j(x) - \beta_j q_{j-1}(x), \quad j = 1, \dots, (n-1).$$
(3.11)

We need to find recurrence relations for the constants α_{i+1} and β_j as well as initial conditions. From the orthonormality constraint we have that

$$q_0(x) = 1$$

and by assuming that

$$q_1(x) = x - \alpha_1,$$

we require orthonormality, i.e.,

$$\sum_{i=1}^{m} w_i q_0(x_i) q_1(x_i) = 0$$

or

$$\sum_{i=1}^{m} w_i(x_i - \alpha_1) = 0$$

or

$$\sum_{i=1}^{m} w_i \alpha_1 = \sum_{i=1}^{m} w_i x_i \Rightarrow \alpha_1 = \frac{1}{\gamma_0} \sum_{i=1}^{m} w_i x_i$$

where

$$\gamma_0 \equiv \sum_{i=1}^m w_i$$

and in general, we define

$$\gamma_k \equiv \sum_{i=1}^m w_i q_k^2(x_i).$$

Similarly, we can obtain the coefficients α_2 and β_2 by insisting that $q_2(x)$ be orthogonal to $q_1(x)$ and $q_0(x)$, which are the two previous polynomials. By induction we obtain the general result from the following two orthogonality constraints:

$$\sum_{i=1}^{m} w_i q_{j+1}(x_i) q_j(x_i) = 0; \quad q_{j+1} \perp q_j$$
(3.12)

$$\sum_{i=1}^{m} w_i q_{j+1}(x_i) q_{j-1}(x_1) = 0, \quad q_{j+1} \perp q_{j-1}$$
(3.13)

These conditions are sufficient to prove that $q_{j+1} \perp q_k$, $k = 0, \ldots, j-2$.

By substituting the recurrence formula (equation (3.11)) for q_{j+1} in equations (3.12) and (3.13), we obtain

$$\alpha_{j+1} = \frac{1}{\gamma_j} \sum_{i=1}^m w_i x_i q_j^2(x_i)$$

and

$$\beta_{j} = \frac{1}{\gamma_{j-1}} \sum_{i=1}^{m} w_{i} \underbrace{\left[x_{i} q_{j-1}(x_{i})\right]}_{eqn.3.11} q_{j}(x_{i})$$

$$= \frac{1}{\gamma_{j-1}} \sum_{i=1}^{m} w_{i} \left[q_{j}(x_{i}) + \alpha_{j} q_{j-1}(x_{i}) + \beta_{j-1} q_{j-2}(x_{i})\right] \cdot q_{j}(x_{i})$$

$$= \frac{1}{\gamma_{j-1}} \sum_{i=1}^{m} w_{i} q_{j}^{2}(x_{i}) = \frac{\gamma_{j}}{\gamma_{j-1}}$$

$$\Rightarrow \beta_{j} = \frac{\gamma_{j}}{\gamma_{j-1}}.$$

We can now write the following recursive algorithm for computing the orthogonal polynomials $q_k(x)$:

Initialize:

$$\gamma_{0} = \sum_{i=1}^{m} w_{i}; \ q_{0}(x) = 1; \ q_{1}(x) = x - \frac{1}{\gamma_{0}} \sum_{i=1}^{m} w_{i} x_{i}$$
Begin Loop $j = 1, n - 1$

$$\gamma_{j} = \sum_{i=1}^{m} w_{i} q_{j}^{2}(x_{i})$$

$$\beta_{j} = \frac{\gamma_{j}}{\gamma_{j-1}}$$

$$\alpha_{i+1} = \frac{1}{\gamma_{j}} \sum_{i=1}^{m} w_{i} x_{i} q_{j}^{2}(x_{i})$$

$$q_{j+1}(x) = x q_{j}(x) - \alpha_{j+1} q_{j}(x) - \beta_{j} q_{j-1}(x)$$

End Loop

Having constructed all the orthogonal polynomials $q_k(x)$, the unknown coefficients are computed from

$$a_k = \frac{1}{\gamma_k} \sum_{i=1}^m w_i q_k(x_i) y_i, \quad k = 0, 1, \dots, n$$

and finally

$$f(x) \approx \sum_{k=0}^{n} a_k q_k(x)$$

is the least-squares polynomial.

Remark: There is a similarity between the procedure we just described and the **QR** decomposition presented in section 2. Recall that the problem of finding a least-square polynomial is equivalent to solving the system

$$\mathbf{C}^T \mathbf{C} \mathbf{a} = \mathbf{C}^T \mathbf{y}.$$

We can apply QR decomposition to matrix C, to obtain

$$C = QR$$

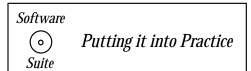
and the above equation becomes

$$\mathbf{R}^T \mathbf{R} \ \mathbf{a} = \mathbf{R}^T \mathbf{Q}^T \mathbf{y}$$

or

$$\mathbf{R} \ \mathbf{a} = \mathbf{Q}^T \mathbf{y}$$

with **R** being upper triangular. The vector of unknown coefficients **a** is then obtained by back substitution. In practice, another version of QR factorization is preferable, the so-called *Householder triangulaziation*, which we will study in chapter 9.



From the discussion above, we see that given a set of data, we need to calculate and store three quantities: α_i , β_i , and the set of least-squares coefficients. We will accomplish this with the function presented below. This function takes as input the number of points over which the least-squares approximation is to be calculated (npts), an array of positions x, data values (or function values) at the previously mentioned spatial points stored in the array funcvals, the degree of the least-squares approximation ndeg, and then as output arrays this routine will fill in the arrays alpha, beta and lscoeffs. Note that this function assumes that all the arrays have been allocated:

}

```
xi = 0.0;
for(i=0;i<npts;i++){</pre>
  xi += xpts[i];
}
xi /= (double) npts;
gamma[0] = npts;
alpha[0] = beta[0] = 0.0;
alpha[1] = xi;
for(j=1;j<=ndeg-1;j++){
  gamma[j]
           = 0.0;
  alpha[j+1] = 0.0;
  for(i=0;i<npts;i++){</pre>
    tmpd = LS_OrthoPoly(j,xpts[i],alpha,beta);
    gamma[j] += tmpd*tmpd;
    alpha[j+1] += xpts[i]*tmpd*tmpd;
  alpha[j+1] /= gamma[j];
  beta[j] = gamma[j]/gamma[j-1];
}
gamma[ndeg] = 0.0;
for(i=0;i<npts;i++){</pre>
  tmpd = LS_OrthoPoly(ndeg,xpts[i],alpha,beta);
  gamma[ndeg] += tmpd*tmpd;
}
beta[ndeg] = gamma[ndeg]/gamma[ndeg-1];
for(j=0;j<=ndeg;j++){
  lscoeffs[j] = 0.0;
  for(i=0;i<npts;i++)</pre>
    lscoeffs[j] = lscoeffs[j] + funcvals[i]*
                  LS_OrthoPoly(j,xpts[i],alpha,beta);
  lscoeffs[j] /= gamma[j];
}
delete[] gamma;
return;
```

There are two issues that we would like to point your attention to as you examine this function:

- 1. **Remark 1:** If you examine the mathematical formulation carefully, you notice that the above function relies on the orthogonal polynomial function being defined. The definition for the orthogonal polynomial requires the α and β to be defined. At first glance, there appears to be a circular dependency! However, it is not. Observe that whenever the orthogonal polynomial needs to know α and β , they have already been properly calculated. The point: this is a highly inductive process. The ordering of calculation is very important in this routine. You should take the time to chart out the dependencies, and observe how timing is everything!
- 2. Remark 2: Dynamic memory allocation within functions is quite natural, and most programmers have no problem allocating arrays within functions. However, many programmers become negligent, and do not deallocate the temporary memory that they needed. In the function above, the array gamma is allocated for temporary usage within this function. Note that this temporary array is deallocated (using the delete[] command) just prior to the function returning. Recall that the local variables within a function are local to the function, and go away when the function concludes. Hence, if memory is allocated to gamma and is not returned to the system prior to the function returning, that memory is "lost" for the remainder of the runtime of your program! This is what is referred to as a memory leak. If you view memory as a conserved quantity (that is, that for every allocation there is a deallocation), then if you forget to deallocate a piece of memory prior to it being inaccessible by the user (in this case due to the pointer variable going away when the function returns), then memory has "leaked," and hence the total amount of memory available for dynamic memory allocation is reduced.

WARNING

Programmer Beware!

Beware of memory leaks!
 For every allocate
 (new) there should be a deallocate (delete[])

Using the Switch



Once again, we use the concept of recursion for quickly implementing the mathematical definition of the orthogonal polynomial. Again, we have two base cases, when the index of the polynomial is 0 or 1; for all other positive values of the index, the value of the polynomial is calculated using the recursion relation.

```
double LS_OrthoPoly(int j, double x, double *alpha, double *beta){
  int i;
  double value;
  switch(j){
  case 0:
   value = 1.0;
   break;
  case 1:
   value = x - alpha[j];
   break;
  default:
    value = (x-alpha[j])*LS_OrthoPoly(j-1,x,alpha,beta) -
      beta[j-1]*LS_OrthoPoly(j-2,x,alpha,beta);
   break;
 }
 return value;
}
```

Once the coefficients have been calculated, then the least-squares approximating polynomial can be evaluated at any point. Below we present the implementation of this function.

```
double LSApproximatingPoly(int ndeg, double x, double *alpha,
   double *beta, double *lscoeffs){
   double value = 0.0;

   for(int i=0;i<=ndeg;i++)
     value += lscoeffs[i]*LS_OrthoPoly(i, x, alpha, beta);

   return value;
}</pre>
```

C++ Compound Assignment

Look carefully at the function above. You will notice something new: the "+=" operator. This is a convenient C++ shorthand used for accumulation. The C++ statement

```
a += b;
is equivalent to the statement
a = a + b;
```

Shorthand	Description
i++	Pre-increment, $i = i + 1$
++i	Post-increment, $i = i + 1$
i–	Pre-decrement, $i = i - 1$
-i	Post-decrement, $i = i - 1$
i += j	i = i + j
i -= j	i = i - j
i *= j	i = i * j
i /= j	i = i / j

Table 3.1: C++ compound assignment operations.

which is to be interpreted as taking the value of b and accumulating it to a. Table 3.1 gives a collection of these "shorthand" programming notations used in C++.

Pre- and post-incrementing/decrementing may be somewhat confusing at first, but consider the following code

```
j = i++;
k = ++p;
```

in which we use the post-incrementor in the first line, and the pre-incrementor in the second line. If we expand this shorthand notation into its traditional C++ code, we obtain the following:

```
j = i;
i = i + 1;
p = p + 1;
k = p;
```

Notice that in the first example, the post-incrementor is used, so the assignment is accomplished first, and then the increment. The exact opposite happens when the pre-incrementor is used. When used as an individual statement (such as we have used it in **for** statements), the two give identical results.

value! Not what was intended!!

Programmer Beware! • C++ shorthands can be convenient but deadly! A slip of the finger, and += can be =+, which is a valid C++ statement, setting one value as the positive value of another

Below, we present a program which uses the functions described above. Notice the general structure of this program:

- 1. Query the user to obtain information.
- 2. Allocate necessary memory (dynamically).



- 3. Produce a grid, and evaluate the function to be approximated.
- 4. Compute least squares coefficients by calling LS_ComputeCoeffs.
- 5. Evaluate approximating polynomial on a fine grid for plotting.
- 6. Deallocate dynamic memory used within the program.

```
#include <iostream.h>
#include "SCchapter3.h"
double func(double x);
int main(int argc, char * argv[]){
  int i;
  int degree, polypnts;
  int npts = 1000;
                         //number of points used for plotting
  double xpt, soln, approx;
  cout << "Enter the degree of the least squares polynomial: ";</pre>
  cin >> degree;
  cout << "Enter the number of points to use for evaluation: ";</pre>
  cin >> polypnts;
  double * poly_xpts = new double[polypnts];
  double * func_vals = new double[polypnts];
                     = new double[degree+1];
  double * alpha
  double * beta
                     = new double[degree+1];
  double * lscoeffs = new double[degree+1];
  CreateGrid_EvenlySpaced(polypnts, poly_xpts, -1.0, 1.0);
  for(i=0;i<polypnts;i++){</pre>
```

```
func_vals[i] = func(poly_xpts[i]);
  }
  LS_ComputeCoeffs(polypnts, poly_xpts, func_vals, degree,
                    alpha, beta, lscoeffs);
  for(i=0;i<npts;i++){</pre>
    xpt = -1.0 + 2.0*i/(npts-1);
    soln = func(xpt);
    approx = LSApproximatingPoly(degree, xpt, alpha,
                                    beta, lscoeffs);
    cout << xpt << " " << soln << " " << approx << endl;</pre>
  }
  delete[] alpha;
  delete[] beta;
  delete[] lscoeffs;
  delete[] poly_xpts;
  delete[] func_vals;
}
double func(double x){
  double y;
  y = 1.0 + 25.0*x*x;
  y = 1.0/y;
  return y;
}
```

Key Concept

• As a programmer, you should have a gameplan! Always take a few moments to formulate the general structure of your program; this will save you much time in the end!

3.1.8 Introduction to Classes

In the previous chapter, we discussed the two fundamental concepts within C++, i.e., the idea of *functions* and the idea of *classes*. In this section, we will present a brief overview of

how to declare, define, and use classes. We will use as an example the class SCVector found in the *software suite*. We will then illustrate the rationale of classes by defining a new class to be used in a least-squares example.

This section is meant only to be a brief overview of classes for the cases where classes are used in this book. To discover the full power of classes (through inheritance, etc.), we refer the reader to [86].

Class Declaration



Classes are *user-defined* data types. We first present the declaration of our user-defined class SCVector, and then will comment on the specifics of the declaration:

```
class SCVector{
private:
  int
        dimension;
  double *data;
public:
 SCVector(int dim);
 SCVector(const SCVector& v);
  SCVector(int col, const SCMatrix &A);
  ~SCVector();
         Dimension() const;
  int
  double Length();
                       /* Euclidean Norm of the Vector */
  void
        Normalize();
  double Norm_11();
  double Norm_12();
  double Norm_linf();
  double MaxMod();
  double ElementofMaxMod();
  int MaxModindex();
  //*************
  // User Defined Operators
  //*************
  int operator==(const SCVector& v) const;
  int operator!=(const SCVector& v) const;
  SCVector & operator=(const SCVector& v);
         operator()(const int i) const;
```

```
double& operator()(const int i);

void Print() const;

void Initialize(double a);

void Initialize(double *v);
};
```

We now present some remarks concerning the code above:

- Observe the structure of a class declaration. First, there is the use of the key word class, followed by the user-defined variable name which will be used later when creating instantiations of this class. Within the {}, there are three key words used to denote accessibility:
 - 1. private those variables and functions that cannot be accessed from outside of the object. Access is non-inheritable.
 - 2. protected those variables and functions that cannot be accessed from outside of the object. Access is inheritable.
 - 3. public those variables and functions that can be accessed from outside of the object.

Those variables/methods within the private section are only accessible from within the object, whereas variables/methods in the public section are accessible outside the object.

- The declaration of a class must be concluded by a ";".
- You may have noticed that we used the term *object* in the above definitions, and you may have assumed that we really meant class. We did not. In C++ nomenclature, a *class* refers to the declaration of the user-defined variable, while an *object* is an instance of a variable of that type. Take for example the predefined variable type *int*. In C++ parlance, we would refer to our declaration of *int* as the class, and every variable that we create as "an object of type *int*."
- Within the private section of this class, we have declared two variables *dimension* and *data* which can be used within the object.
- In the public section, we have declared a collection of methods which access or modify the data contained within the object.

Method Definitions



We now discuss some of the method definitions for this class found within the *software suite*.

Each method has the following structure:

```
<return type> ClassName::MethodName(<argument list>)
```

The return type, method name, and argument list are similar to what we have seen with functions. In the case of a class method, we also designate the class to which the method is assigned (using the "ClassName:" syntax).

Constructor: A constructor is the first function which is called when the object is instantiated. In this case, the constructor requires the input of the dimension of the vector. The constructor then uses this information to initialize the local variable *dimension*, and to allocate memory which is assigned to the local variable *data*.

```
SCVector::SCVector(int dim){
  dimension = dim;
  data = new double[dimension];
  for(int i=0;i<dimension;i++)
    data[i] = 0.0;
}</pre>
```

Copy Constructor: A copy constructor is used whenever a copy is required (either by the programmer or the program). The object to be copied is passed as an argument to this method. Notice that the argument is passed as a const (meaning that within the method we cannot change the value of the object v), and that it is passed by reference (denoted by the "&") so that no new memory allocation is required to store the contents of v (as opposed to if we had passed by value). The current object is initialized so that it is a copy of v.

```
SCVector::SCVector(const SCVector &v){
  dimension = v.Dimension();
  data = new double[dimension];

for(int i=0;i<dimension;i++)
   data[i] = v.data[i];
}</pre>
```

Destructor: This method is called automatically when the object is released back to the operating system. Its purpose is to clean up the storage contained within the object.

```
SCVector::~SCVector(){
  dimension = 0;
  delete[] data;
  data = NULL;
}
```

General Methods: From the class declaration presented above, we present two method definitions, one which merely accesses the data with the object to provide a result, and the second with acts upon the data contained within the object. Consider the following two method definitions:

```
double SCVector::Norm_12(){
   double sum = 0.0;
   for(int i=0;i<dimension;i++)
      sum += data[i]*data[i];
   return(sqrt(sum));
}

void SCVector::Normalize(){
   double tmp = 1.0/Norm_12();
   for(int i=0;i<dimension;i++)
      data[i] = data[i]*tmp;
}</pre>
```

In the first method, we use the information stored within the class (contained within data) to compute the L_2 norm of the vector, and we return this information at the conclusion of the function. In the second method, we act upon the data contained within the object by normalizing the value of the vector to one. Notice within the *Normalize* method that we call local class method Norm 2 to obtain the discrete L_2 norm of the vector.

Overloaded Operators

In addition to class methods, we can also *overload* operators so that they are appropriately defined for our new user-defined data type. Consider the following operator declaration:

```
SCVector operator+(const SCVector& v1, const SCVector& v2);
and corresponding operator definition:

SCVector operator+(const SCVector& v1, const SCVector& v2){
  int min_dim = min_dimension(v1,v2);
  SCVector x(min_dim);
  for(int i=0;i<min_dim;i++)
    x(i) = v1(i) + v2(i);
  return x;
}

The basic syntax is as follows:

<return type> operator<symbol>(<argument 1>,<argument 2>)
```

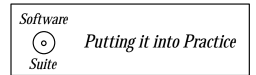
where the return type, symbol, and arguments are to be supplied by the programmer. Using this syntax we appropriately define what it means to add (using the binary operator '+') two SCVector objects. We will illustrate how this is used below.

Object Allocation and Usage

To understand how all the information presented above is used, consider the following code:

```
SCVector a(3),b(3),c(3); //allocate SCVectors a,b,c
                       //Constructor is called automatically to
                       //allocate memory and provide default
                       //initialization
a(0) = 1.0;
            //Initialize the values of 'a' using () operator
a(1) = 2.0;
            //defined for the class SCVector
a(2) = 3.0;
b(0) = -2.0; //Initialize the values of 'b'using () operator
b(1) = 1.0;
             //defined for the class SCVector
b(2) = 3.0;
             //Use overloaded operator '+' to compute
c = a+b;
             //the sum of a and b, and use of the
             //overloaded operator '=' to assign
             //the value to 'c'
c.Print();
             //Use print method to print the value of c
```

Execution of this code within a program would yield the result [-1.0; 3.0; 6.0] printed to standard output.



The least-squares example presented earlier provides a good motivation for using classes. Observe that associated with each least-squares approximating polynomial we form, we need to keep track of three arrays, alpha, beta and lscoeffs, for each approximating polynomial! Is this do-able? Certainly. But from an organizational standpoint, we would like to be able to automatically associate the appropriate arrays with the right polynomials. Imagine that we were asked to handle 20-100 least squares approximations simultaneously! Doable, yes; messy, yes; but recall that classes can provide us a means of organizing our data.

First, we begin by giving the class *declaration*:

```
class LSPoly{
```

In addition to the four variables ndeg, alpha, beta, and lscoeffs declared within the class, five methods associated with this class have declared. We present the definitions of these five methods and provide a brief explanation of each.

Default Constructor: This method is called automatically when the object is instantiated if no other constructor is called. Observe that in this case we initialize variables to either zero or NULL (whichever is appropriate for the variable type).

```
LSPoly::LSPoly(){
  ndeg = 0;
  alpha = NULL;
  beta = NULL;
  lscoeffs = NULL;
}
```

Destructor: This method is called automatically when the object is destroyed.

```
LSPoly::~LSPoly(){
  delete[] alpha;
  delete[] beta;
  delete[] lscoeffs;

ndeg = 0;
  alpha = NULL;
  beta = NULL;
  lscoeffs = NULL;
}
```

The next method is a "private" method; that is, it can only be called from within the object. This means that the only valid places that this function can be called are within other methods defined within the object.

```
double LSPoly::LSPolyOrtho(int j,double x){
  int i;
  double value;
  switch(j){
  case 0:
    value = 1.0;
    break;
  case 1:
    value = x - alpha[j];
    break;
  default:
    value = (x-alpha[j])*LSPolyOrtho(j-1,x) -
      beta[j-1]*LSPolyOrtho(j-2,x);
    break;
  }
 return value;
}
```

The next three methods are "public" methods; that is, they can be accessed from outside the object. The first method accomplishes the initialization of the class which consists of computing the values stored in *alpha*, *beta*, and *gamma*. The second method allows us to print the contents of the object, and the third method allows us to evaluate the least-squares approximation using the information stored within the object.

```
double * gamma = new double[ndeg+1];
// Compute average first
xi = 0.0;
for(i=0;i<npts;i++){</pre>
 xi += xpts[i];
}
xi /= (double) npts;
gamma[0] = npts;
alpha[0] = beta[0] = 0.0;
alpha[1] = xi;
for(j=1; j \le ndeg-1; j++){
 gamma[j] = 0.0;
  alpha[j+1] = 0.0;
  for(i=0;i<npts;i++){</pre>
    tmpd = LS_OrthoPoly(j,xpts[i],alpha,beta);
    gamma[j] += tmpd*tmpd;
    alpha[j+1] += xpts[i]*tmpd*tmpd;
 alpha[j+1] /= gamma[j];
 beta[j] = gamma[j]/gamma[j-1];
}
gamma[ndeg] = 0.0;
for(i=0;i<npts;i++){</pre>
  tmpd = LSPolyOrtho(ndeg,xpts[i]);
 gamma[ndeg] += tmpd*tmpd;
}
beta[ndeg] = gamma[ndeg]/gamma[ndeg-1];
for(j=0;j<=ndeg;j++){
 lscoeffs[j] = 0.0;
 for(i=0;i<npts;i++)</pre>
    lscoeffs[j] = lscoeffs[j] + funcvals[i]*LSPolyOrtho(j,xpts[i]);
 lscoeffs[j] /= gamma[j];
}
delete[] gamma;
```

```
return 1;
}
void LSPoly::PrintCoeffs(){
  cout << endl;</pre>
  cout << "i\talpha\tbeta\tlscoeffs" << endl;</pre>
  for(int j=0; j \le ndeg; j++){
   cout << j << "\t" << alpha[j] << "\t";</pre>
   cout << beta[j] << "\t" << lscoeffs[j] << endl;</pre>
 }
  cout << "********************* << endl << endl;</pre>
 return;
}
double LSPoly::Evaluate(double x){
 double value = 0.0;
 for(int i=0;i<=ndeg;i++)</pre>
   value += lscoeffs[i]*LSPolyOrtho(i,x);
 return value;
}
```

Now, we want to put it all together into one piece of code. Using the information above, we have now declared a class named LSPoly, and we have provided definitions to all its methods. We now use this new user-defined variable in the program below.

```
LSPoly poly; // Our user-defined class!
  cout << "Enter the degree of the least-squares polynomial: ";</pre>
  cin >> degree;
  cout << "Enter the number of points to use for evaluation: ";</pre>
  cin >> polypnts;
  double * poly_xpts = new double[polypnts];
  double * func_vals = new double[polypnts];
  CreateGrid_EvenlySpaced(polypnts, poly_xpts, -1.0, 1.0);
  for(i=0;i<polypnts;i++){</pre>
    func_vals[i] = func(poly_xpts[i]);
  }
 poly.Initialize(polypnts,degree,poly_xpts,func_vals);
  for(i=0;i<npts;i++){</pre>
    xpt = -1.0 + 2.0*i/(npts-1);
    soln = func(xpt);
    approx = poly.Evaluate(xpt);
    cout << xpt << " " << soln << " " << approx << endl;</pre>
  }
  delete[] poly_xpts;
  delete[] func_vals;
}
double func(double x){
  double y;
  y = 1.0 + 25.0*x*x;
  y = 1.0/y;
  return y;
```

We want to draw your attention to certain key items within the program above:

- We begin by instantiating a variable of type LSPoly, just like creating a "regular" (pre-defined) variable. As stated above, when the variable is initiated, the constructor is called.
- To access both variables and methods which are public, we use the "." notation.

<variable name>.<object variable>

or

```
<variable name>.<method>( ... <method argument list> ... )
```

In the example above, we access the *Initialize* method as follows:

```
poly.Initialize(polypnts,degree,poly_xpts,func_vals);
```

where *poly* is the name of the object, and *Initialize* is the name of the public method that we want to access.

If, instead of the object, we were using a pointer to the object, we use the "-¿" notation as follows:

<pointer variable name>-><object variable>

or

```
<pointer variable name>-><method>( ... <method argument list> ... )
```

• All the information necessary for the least squares approximation is stored within the object, and hence the call for evaluating the least squares polynomial is merely the call:

```
approx = poly.Evaluate(xpt);
```

3.1.9 Multi-Dimensional Interpolations

We can extend the interpolation methods we have presented so far in two-dimensions or three-dimensions by constructing appropriate two- or three-dimensional polynomials. In two dimensions, for example, we have

$$f(x,y) = \sum_{k} \alpha_k \phi_k(x,y)$$

where α_k are the unknown coefficients, and the exact form of the polynomial basis $\phi_k(x,y)$ depends on the shape of the computational domain. In order to simplify the presentation, we first consider canonical domains, and subsequently we present mapping techniques to deal with more general domains. The approach presented here is typically followed in finite element methods, see [63], where polynomial approximations in subdomains (the "elements") are required; however this polynomial approximation is general, and also is easy to implement.

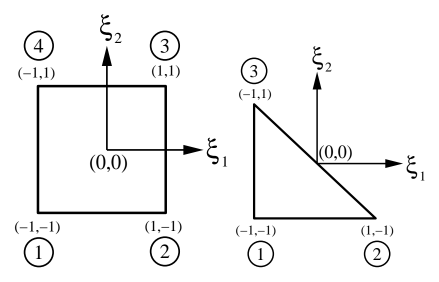


Figure 3.10: Standard domains for the quadrilateral (left), and triangular (right) expansion in terms of the Cartesian coordinates ξ_1, ξ_2 .

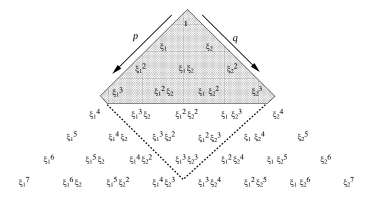


Figure 3.11: Pascal's diagram that shows the polynomial space for the triangular expansion (shaded region) and square region (shaded region plus values within dotted line). The example here is for cubic polynomial approximation.

3.1.10 Simple Domains

The canonical domains we consider are the square and the triangular domains with the coordinates $\xi_1 \in [-1,1]$ and $\xi_2 \in [-1,1]$ being the normalized coordinates as shown in figure 3.10. Similar extensions can be constructed in three-dimensions for a standard hexahedron and a tetrahedron. In figure 3.11 we construct the so-called *Pascal's diagram* to demonstrate graphically the polynomial space for each region.

We first consider the square domain, for which it is computationally efficient to split directions by constructing tensor-products of the form

$$f(\xi_1, \xi_2) \approx \sum_{p} \sum_{q} \alpha_{pq} \phi_p(\xi_1) \phi_q(\xi_2).$$

Let us re-write the above equation using one-dimensional Lagrangian interpolants $h_i(\xi)$, i.e.

$$f(\xi_1, \xi_2) \approx \sum_{p} \sum_{q} f_{pq} h_p(\xi_1) h_q(\xi_2)$$

where f_{pq} are the function values at the node (p, q).

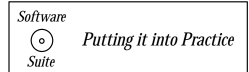
In the standard square, $bilinear\ interpolation$ is achieved by employing the one-dimensional linear interpolants

$$h_i(\xi) = \frac{1}{2}(1 \pm \xi), \quad i = 1, 2$$

and thus

$$f(\xi_1, \xi_2) \approx \frac{1}{4} [f_1(1 - \xi_1)(1 - \xi_2)] + \frac{1}{4} [f_2(1 + \xi_1)(1 - \xi_2)] + \frac{1}{4} [f_3(1 + \xi_1)(1 + \xi_2)] + \frac{1}{4} [f_4(1 - \xi_1)(1 + \xi_2)]$$

where we have used a counter-clockwise convention to number the corners of the domain, and f_i denotes the function values at each vertex starting from the low left corner, see figure 3.10.



We can implement the above formula as follows:

```
double Square_2dInterpolant(SCPoint x, int npts, double *funcvals){
  double value = 0.;
  double h[4];
  if(npts != 4){
    cerr << "Error in Square_2dInterpolant -- ";</pre>
    cerr << "Invalid npts given" << endl;</pre>
    return value;
  }
 h[0] = 0.5*(1.0-x(0));
 h[1] = 0.5*(1.0+x(0));
 h[2] = 0.5*(1.0-x(1));
 h[3] = 0.5*(1.0+x(1));
  value = funcvals[0]*h[0]*h[2] + funcvals[1]*h[1]*h[2] +
          funcvals[2]*h[1]*h[3] + funcvals[3]*h[0]*h[3];
  return value;
}
```

In this routine, we rely on the use of the data type SCPoint, which is a user-defined class included in the software suite. What is important for you to know concerning SCPoint is that if P is a variable of type SCPoint, then the coordinate (x, y) is stored as (P(0), P(1)). Hence, in this 2D example, we are providing an SCPoint which has two accessible values, x(0) and x(1): $x(0) = \xi_0$ and $x(1) = \xi_1$.

Let us observe a few items within this code:

- Notice that because we know how many interpolants we will need, we can use static allocation of an array for holding the temporary values of the interpolants. If instead of four points we knew that we would use nine points, then we could allocate h[9] instead of h[4]. Obviously this is much easier than typing h0,h1, ... h8! It would also be perfectly valid to use dynamic memory allocation here; we have chosen not to for optimization reasons. For such a small number of variables, the cost of dynamic memory allocation outweighs the convenience.
- In this function, we introduce the concept of argument checking. We have written the function above with the intention of providing the two-dimensional bilinear interpolation. From the theory, we see that this requires that four function values be given. What would happen if we are only given three function values (that is, what if the user had only allocated and assigned values for funcvals[0], funcvals[1] and funcvals[2])? Using this function would be invalid, and in all likelihood our program would crash! Quite often programmers introduce checks into their code to help minimize such mistakes. In this case, we check to make sure that we have received npts = 4; if not, we issue a warning that we received an invalid npts value, and we return 0.0.
- Notice that we used *cerr* instead of *cout*. The object *cerr* is an output object like *cout*, and is also declared in *iostream.h*. The difference between *cout* and *cerr* is that *cerr* writes to **standard error** instead of standard output (which may or may not be the same actual output device).

Key Concept

 Carefully designed argument checking can save you hours of debugging time! Plan ahead for your own possible mistakes – and prevent them from occurring.

High-Order Interpolation

Higher order approximations can be constructed if more information about the function f(x) is given. Typically, that information may be available at the midpoints of the edges of the

domain or even at the center of the domain. In this case, we need to first construct higher order one-dimensional Lagrangian interpolants. Let us consider the case of an extra *one* and *two* interior points distributed equidistantly along the edge. From their definition we have that

$$h_3(\xi) = 1 - \xi^2$$
 and $h_4(\xi) = 1 - 27\xi^3 - 9\xi^2 + 27\xi + 9$.

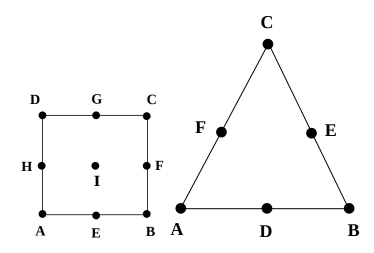


Figure 3.12: Domain and points where the data is specified for square (left) and triangle (right).

With the above four one-dimensional Lagrangian interpolants we can construct up to third-order polynomial approximations in a square domain if information is given at all the points A, B, ..., I as shown in figure 3.12. As an example, let us assume that we have available f(A), f(B), f(C), f(D) and f(E). Then the interpolants (also called *shape functions*) are as follows:

$$h_A = \frac{1}{4}(1 - \xi_1)(1 - \xi_2) - \frac{1}{2}h_E; \quad h_B = \frac{1}{4}(1 + \xi_1)(1 - \xi_2) - \frac{1}{2}h_E;$$

$$h_C = \frac{1}{4}(1 - \xi_1)(1 + \xi_2); \qquad h_D = \frac{1}{4}(1 + \xi_1)(1 + \xi_2);$$

$$h_E = \frac{1}{2}(1 - \xi_1^2) \qquad .$$

Let us now assume that in addition we also have function values at the points I and F. Then the above interpolants can be easily modified to handle this case as well by adding extra terms that reflect this interaction

$$h_A = \frac{1}{4}(1 - \xi_1)(1 - \xi_2) - \frac{1}{2}h_E - \frac{1}{4}h_I; \quad h_B = \frac{1}{4}(1 + \xi_1)(1 - \xi_2) - \frac{1}{2}h_E - \frac{1}{2}h_F - \frac{1}{4}h_I$$

$$h_C = \frac{1}{4}(1 - \xi_1)(1 + \xi_2) - \frac{1}{2}h_F - \frac{1}{4}h_I; \qquad h_D = \frac{1}{4}(1 + \xi_1)(1 + \xi_2) - \frac{1}{4}h_I$$

$$h_E = \frac{1}{2}(1 - \xi_1^2)(1 - \xi_2) - \frac{1}{2}h_I; \qquad h_F = \frac{1}{2}(1 + \xi_1)(1 - \xi_2^2) - \frac{1}{2}h_I$$

$$h_I = (1 - \xi_1^2)(1 - \xi_2^2).$$

These approximations can be extended to *hexahedral domains* using the tensor-product form. For example, in the case of *trilinear interpolation* with the function values specified at the eight vertices the shape functions are:

$$h_i(\xi_1, \xi_2, \xi_3) = \frac{1}{8} (1 \pm \xi_1) (1 \pm \xi_2) (1 \pm \xi_3).$$

For higher order interpolations similar constructions can be obtained with the approximating polynomials defined by the hierarchy shown in figure 3.13(a), which is Pascal's diagram for hexahedral domains. For more details see ([63], chapter 3).

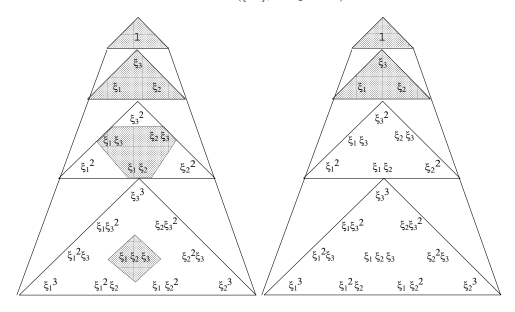


Figure 3.13: Pascal's diagram for hexahedral (left) and tetrahedal domains (right).

Non-Tensor Products

For a triangular region, however, no tensor products are easily constructed unless special coordinate systems and transformations are introduced, see [63]. For linear interpolation, the shape functions are constructed, e.g., by collapsing the corners D and C of the square to obtain

$$h_i^{tr} = h_i^{sq} = \frac{1}{4}(1 + (-1)^i r)(1 - s), \quad i = A, B$$

and

$$h_C^{tr} = h_C^{sq} + h_D^{sq} = \frac{1}{2}(1 + \xi_2).$$

These can be recomputed after we renormalize the coordinates so that $\xi_1 \in [0, 1]$ and $\xi_2 \in [0, 1]$ to obtain

$$h_A = 1 - \xi_1 - \xi_2, \quad h_B = \xi_1, \quad h_C = \xi_2.$$

Furthermore, if we have data at the midpoints of the edges of the triangle (see figure 3.12(b)) we can construct complete quadratic interpolation using the shape functions

$$h_A = 1 - \xi_1 - \xi_2 - \frac{1}{2}h_D - \frac{1}{2}h_F; \quad h_B = \xi_1 - \frac{1}{2}h_D - \frac{1}{2}h_E;$$

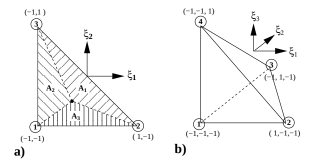


Figure 3.14: (a) The area coordinate system in the standard triangular region. Each coordinate l_1, l_2 , and l_3 can be interpreted as the ratio of areas A_1, A_2 , and A_3 over the total area. (b) The standard tetrahedral region for the definition of volume coordinates.

$$h_C = \xi_2 - \frac{1}{2}h_E - \frac{1}{2}h_F; \quad h_D = 4\xi_1(1 - \xi_1 - \xi_2);$$

 $h_E = 4\xi_1\xi_2; \quad h_F = 4\xi_2(1 - \xi_1 - \xi_2).$

Another popular way of constructing linear interpolations in a triangular region is by using the area (and volume in three-dimensions) coordinates, otherwise known as *barycentric* or *triangular/tetrahedral* coordinates.

The area coordinate system is illustrated in figure 3.14(a) for the standard triangle. Any point in the triangle is described by three coordinates l_1, l_2 , and l_3 , which can be interpreted as the ratio of the areas A_1, A_2 and A_3 over the total area $A = A_1 + A_2 + A_3$, that is,

$$l_1 = \frac{A_1}{A};$$
 $l_2 = \frac{A_2}{A};$ $l_3 = \frac{A_3}{A}.$

Therefore l_1, l_2 , and l_3 have a unit value at the vertices marked 1, 2 and 3 in figure 3.14(a), respectively. By definition these coordinates satisfy the relationship:

$$l_1 + l_2 + l_3 = 1,$$

and they can be expressed in terms of ξ_1, ξ_2 as:

$$l_1 = \frac{1}{2}(1 - \xi_1) - \frac{1}{2}(1 + \xi_2);$$

$$l_2 = \frac{1}{2}(1 + \xi_1);$$

$$l_3 = \frac{1}{2}(1 + \xi_2).$$

A similar construction follows for volume coordinates l_1, l_2, l_3, l_4 , which are defined as having a unit value at the vertices marked 1, 2, 3, 4 in figure 3.14(b). In terms of the local Cartesian coordinates the volume coordinate system is defined as:

$$l_1 = \frac{-(1+\xi_1+\xi_2+\xi_3)}{2};$$
 $l_2 = \frac{(1+\xi_1)}{2};$ $l_3 = \frac{(1+\xi_2)}{2};$ $l_4 = \frac{(1+\xi_3)}{2}.$

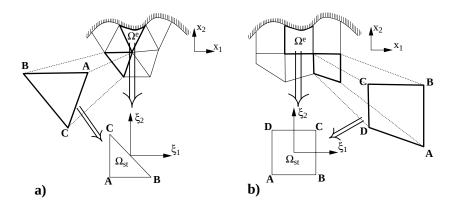


Figure 3.15: To construct a C^0 expansion from multiple elements of specified shapes (for example, triangles or rectangles), each elemental region Ω^e is mapped to a standard region Ω_{st} in which all local operations are evaluated [63].

3.1.11 Curvilinear Domains

In many practical simulation problems we need to perform polynomial interpolation in regions which may be of an arbitrary shape and orientation as illustrated in figure 3.15. Such general domains can be broken into subdomains which are triangular or quadrilateral regions. These can be then transformed to standard regions as shown in figure 3.10. To treat these more general domains we follow the method presented in [63]; we introduce a one-to-one mapping denoted by

$$x_1 = \chi_1^e(\xi_1, \xi_2), \qquad x_2 = \chi_2^e(\xi_1, \xi_2)$$

in two dimensions, and similarly

$$x_1 = \chi_1^e(\xi_1, \xi_2, \xi_3), \qquad x_2 = \chi_2^e(\xi_1, \xi_2, \xi_3), \qquad x_3 = \chi_3^e(\xi_1, \xi_2, \xi_3)$$

in three dimensions.

For elemental shapes with straight sides a simple mapping may be constructed using linear interpolation. For example, to map a triangular region [as in figure 3.15(a)] assuming that the coordinates of the triangle $\{(x_1^A, x_2^A), (x_1^B, x_2^B), (x_1^C, x_2^C)\}$ are known we can use

$$x_1 = \chi(\xi_1, \xi_2) = \frac{1}{2} x_1^A \left(-\xi_2 - \xi_1 \right) + \frac{1}{2} x_1^B \left(1 + \xi_1 \right) + \frac{1}{2} x_1^C \left(1 + \xi_2 \right). \tag{3.14}$$

A similar approach leads to the bilinear mapping for an arbitrary shaped straight-sided quadrilateral where only the vertices need to be prescribed. For the straight-sided quadrilateral with vertices labeled as shown in figure 3.15(b) the mapping is:

$$x_{1} = \chi_{1}(\xi_{1}, \xi_{2}) = x_{1}^{A} \frac{(1 - \xi_{1})}{2} \frac{(1 - \xi_{2})}{2} + x_{1}^{B} \frac{(1 + \xi_{1})}{2} \frac{(1 - \xi_{2})}{2} + x_{1}^{D} \frac{(1 - \xi_{1})}{2} \frac{(1 + \xi_{2})}{2} + x_{1}^{C} \frac{(1 + \xi_{1})}{2} \frac{(1 + \xi_{2})}{2}.$$
 (3.15)

When developing a mapping it is important to ensure that the Jacobian of the mapping to the standard region is non-zero and of the same sign. To ensure this condition is satisfied

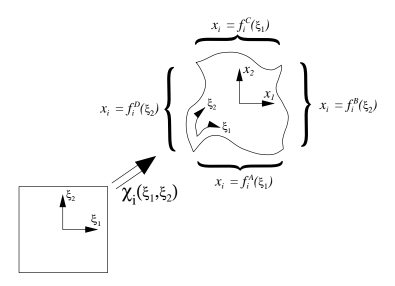


Figure 3.16: A general curved element can be described in terms of a series of parametric functions $f^A(\xi_1), f^B(\xi_2), f^C(\xi_1)$, and $f^D(\xi_2)$. Representing these functions as a discrete expansion we can construct a mapping $\chi_i(\xi_1, \xi_2)$ relating the standard region (ξ_1, ξ_2) to the deformed region (x_1, x_2) . [63].

when using the mappings given above, we require all elemental regions to have internal corners with angles that are less than 180° and so are convex.

To describe a straight-sided region we only need to know the values of the vertex locations. However, to describe a curved region we need more information. Specifically, as illustrated in figure 3.16 we need a description of the shape of each edge in terms of a series of parametric functions which we define as $f_i^A(\xi_1)$, $f_i^B(\xi_2)$, $f_i^C(\xi_1)$ and $f_i^D(\xi_2)$. Since our mapping $\chi_i(\xi_1, \xi_2)$ maps the whole of the standard region to the curvilinear quadrilateral domain, the parameter can be taken as the local coordinate ξ_1 or ξ_2 .

A practical algorithm of doing this is the method of blending function as originally proposed by Gordon and Hall [48]. For the quadrilateral region shown in figure 3.16 the linear blending function is given by

$$\chi_{i}(\xi_{1}, \xi_{2}) = f^{A}(\xi_{1}) \frac{(1-\xi_{2})}{2} + f^{C}(\xi_{1}) \frac{(1+\xi_{2})}{2}
+ f^{D}(\xi_{2}) \frac{(1-\xi_{1})}{2} + f^{B}(\xi_{2}) \frac{(1+\xi_{1})}{2}
- \frac{(1-\xi_{1})}{2} \frac{(1-\xi_{2})}{2} f^{A}(-1) - \frac{(1+\xi_{1})}{2} \frac{(1-\xi_{2})}{2} f^{A}(1)
- \frac{(1-\xi_{1})}{2} \frac{(1+\xi_{2})}{2} f^{C}(-1) - \frac{(1+\xi_{1})}{2} \frac{(1+\xi_{2})}{2} f^{C}(1),$$
(3.16)

where the vertex points are continuous [for example, $f^A(-1) = f^D(-1)$] and so the last four may also be expressed in terms of f^B and f^D . The mapping function of a curved-sided element is constructed by approximating the edge function in terms of the Lagrange

polynomial, that is,

$$f_i^A(\xi_1) \approx \sum_{p=0} f_i^A(\xi_{1,p}) h_p(\xi_1),$$

and then using the linear blending function equation (3.16).

3.2 Fourier Series Representation

In this section, we will consider interpolations based on a basis $\phi_k(x) = e^{ikx}$ or $\cos kx$ or $\sin kx$. This is the Fourier representation in terms of complex exponentials, and cosines or sines, respectively; that is we represent an arbitrary function in terms of pure harmonic functions. The Fourier method works in the frequency domain (or wavenumber domain) to provide information in the physical domain. This ability to relate frequency and space or time domains has made the Fourier transform one of the most useful tools of numerical and mathematical analysis. With the discovery of a fast algorithm in 1965, which reduces its computational complexity from $\mathcal{O}(N^2)$ to $\mathcal{O}(N\log_2 N)$, the FFT (Fast Fourier Transform) is also an extremely effective tool in scientific computing.

3.2.1 Convergence

The main idea of Fourier series is to represent a function y = f(x) with a basis consisting of sines and cosines or complex exponentials. However, unlike the polynomial approximation, the convergence of Fourier series is not always guaranteed in the point-wise sense for an arbitrary function. Here, we will follow the exposition of Lanczos to present the basic material, see [66].

In general, if the function f(x) satisfies the so-called *Dirichlet* conditions, it converges, but such conditions are too restrictive. Specifically, a function satisfies the Dirichlet conditions if:

- f(x) is defined at every point in the interval $x \in [-\pi, \pi]$.
- f(x) is single valued, piecewise continuous, and finite; e.g., the function $f(x) = \log x$ is excluded.
- f(x) is of bounded variation, that is f(x) cannot have an infinite number of maxima and minima.

A function that satisfies the Dirichlet conditions can be expanded into the following convergent infinite series:

$$f(x) = \frac{1}{2}a_0 + a_1\cos x + a_2\cos 2x + \dots + b_1\sin x + b_2\sin 2x + \dots$$
 (3.17)

where the *Fourier coefficients* are computed from:

$$a_k = \frac{1}{\pi} \int_{-\pi}^{\pi} f(x) \cos kx \, dx,$$

and

$$b_k = \frac{1}{\pi} \int_{-\pi}^{\pi} f(x) \sin kx \ dx.$$

Equivalently, we can use complex exponentials as the basis, to write

$$f(x) = \sum_{k=-\infty}^{\infty} c_k e^{ikx}$$
 with $c_k = \frac{1}{2\pi} \int_{-\pi}^{\pi} f(x)e^{-ikx} dx$. (3.18)

Also, the two sets of coefficients are related, i.e.,

$$a_k = c_k + c_{-k} (3.19)$$

$$b_k = \frac{1}{i}(c_k - c_{-k}). {(3.20)}$$

The truncated version of the above series is called *discrete Fourier series*, and has the form:

$$f_N(x) = \sum_{k=-(N-1)}^{N-1} c_k e^{ikx}.$$

As we will see below, it can be constructed by sampling the function at N equidistant points

$$x_k = \frac{2\pi k}{N}, \quad k = 0, \dots N - 1.$$

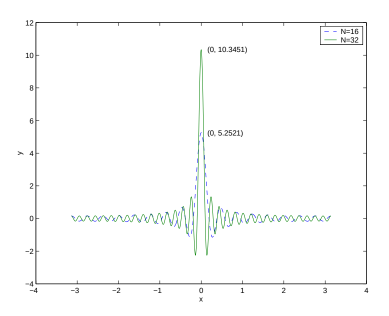


Figure 3.17: Fourier series of $\delta(x)$ for N=16 and N=32.

Example: The Fourier coefficients of the Dirac function $\delta(x)$ are:

$$a_k = \frac{1}{\pi} \int \delta(x) \cos kx dx = \frac{1}{\pi}$$

$$b_k = \frac{1}{\pi} \int \delta(x) \sin kx dx = 0$$

$$c_k = \frac{1}{2\pi} \int \delta(x) e^{ikx} dx = \frac{1}{2\pi}.$$

The Fourier series of $\delta(x)$ for N=16 and N=32 are shown in figure 3.17. The peak is sharper at higher values of N. The Dirac function does not satisfy the Dirichlet conditions.

Dirichlet Kernel

We now return to the question of convergence, and specifically the validity of equations (3.17) or (3.18). What Dirichlet did was to substitute back into equations (3.17) or (3.18) the expressions for the Fourier coefficients a_k, b_k, c_k and truncate after N terms; then he let $N \to \infty$ to obtain

$$f_N(x) = \int_{-\pi}^{\pi} f(z) D_n(x-z) dz$$

where

$$D_N(t) = \frac{\sin(N+1/2)t}{2\pi\sin\left(\frac{t}{2}\right)},$$
 Dirichlet kernel

The Dirichlet kernel contains the partial sum

$$\sum_{-N}^{N} e^{ikt} = e^{-iNt} \left[1 + e^{it} + e^{2it} + \dots + e^{2Nit} \right] = \frac{e^{i(N+1/2)t} - e^{-i(N+1/2)t}}{e^{i(1/2)t} - e^{-i(1/2)t}}$$
$$= \frac{\sin\left(N + \frac{1}{2}\right)t}{\sin\frac{1}{2}t}.$$

The above Dirichlet kernel can be thought as a "lens" that focuses the action at the point z = x. In order to ensure that $\lim_{N\to\infty} f_N(x) = f(x)$, it requires a very strong focusing power around the point t = 0 of the kernel $D_N(t)$. Mathematically, this can be expressed by the following two conditions, see [66]:

• (A)
$$\lim_{N \to \infty} \int_{\epsilon}^{\pi} |D_N(t)| dt = 0$$

• (B)
$$\lim_{N \to \infty} \int_{-\epsilon}^{\epsilon} D_N(t) dt = 1$$

Condition (A) is not satisfied by the Dirichlet kernel because its secondary maxima are comparable to the primary maximum at t = 0.

Fejer's Construction

A different path to convergence was suggested by Fejer who proposed an alternative summation procedure. He only imposed the constraint

$$\int_{-\pi}^{+\pi} |f(x)| dx < \infty \,,$$

which states that f(x) is an absolutely integrable function. For example, the function $y = \log x$ is absolutely integrable but the function y = 1/x is not. He then considered the partial sums

$$S_0 = \frac{1}{2}a_0,$$

$$S_1 = \frac{1}{2}a_0 + a_1 \cos x + b_1 \sin x \dots,$$

$$\vdots$$

$$S_N = \frac{1}{2}a_0 + a_1 \cos x + \dots + a_N \cos Nx + b_1 \sin x + \dots + b_N \sin Nx,$$

and constructed the sequence:

$$f_1 = S_0, f_2 = \frac{S_0 + S_1}{2}, \dots, S_N = \frac{s_0 + s_1 + \dots + s_{N-1}}{N}.$$

This sequence leads to the kernel

$$F_N(t) = \frac{\sin^2(Nt/2)}{2\pi N \sin^2(t/2)},$$

which is very focused and satisfies conditions (A) and (B). Unlike the previous case, the Fejer sequence converges at all points including points of discontinuity, where it converges at the *arithmetic mean* of the values at either side of the discontinuity.

3.2.2 Periodic Extension of Functions

Consider now that f(x) is defined in the interval $x \in [0, \pi]$. We can still represent the function as a Fourier series by appropriately extend it, i.e., define it in the interval $x \in [-\pi, x]$ as well by representing it either by an even or an odd function. The Fourier series will then involve *cosines* or *sines*, respectively. If f(x) is not zero at the boundary points the reflection as an odd function will cause a discontinuity at $x = 0, \pi$, which will lead to the so-called "Gibbs phenomenon." This is manifested by wiggles around the discontinuity that affect the solution everywhere. This is avoided for an even function.

An example of a periodic extension for the function $y = x/\pi$, where $x \in [0, \pi]$ is shown in figure 3.18. Both odd and even constructions are shown, with the latter resulting in a C^0 continuous extended function.

Let us now assume that f(0) = 0 and also $f(\pi) = 0$. Then we should expand f(x) using a *sine series* as both the (extended) function and first derivative are continuous. We can compare the convergence rates of the two representations by performing integration by parts, i.e., for the *cosine series*

$$\int f(x)\cos kx \, dx = \frac{f(x)\sin kx}{k} - \frac{1}{k} \int f'(x)\sin kx \, dx$$

and the second term gives

$$\int f'(x)\sin kx \ dx = -\frac{f'(x)\cos kx}{k} + \frac{1}{k} \int f''(x)\cos kx \ dx.$$

Thus, the convergence of

$$\int_0^\pi f(x)\cos kx \ dx$$

is dictated by the boundary term

$$\frac{(-1)^k f'(\pi) - f'(0)}{k^2} \sim \frac{1}{k^2}.$$

In contrast, the *sine series* representation leads to

$$\int_0^{\pi} f(x) \sin kx \, dx \to \frac{f(0) - (-1)^k f(\pi)}{k} \sim \frac{1}{k}.$$

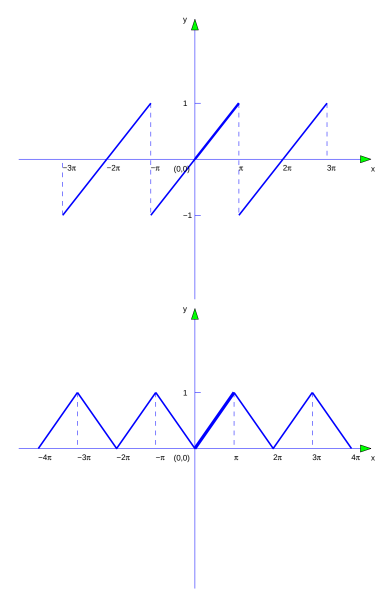


Figure 3.18: Periodic extensions of the function $y = x/\pi$, where $x \in [0, \pi]$ using an odd function extension (upper) and an even function extension (lower).

Now if

$$f(0) = f(\pi) = 0 \Rightarrow \int_0^{\pi} f(x) \sin kx \, dx \to \frac{f''(x) \cos kx}{k^3} \to \frac{(-1)^k f''(\pi) - f''(0)}{k^3}$$
$$\sim \frac{1}{k^3}.$$

Therefore, we conclude that in this case the sine series converges like $1/k^3$ versus $1/k^2$ for the cosine series, and therefore the former is a better approach.

Some Properties of the Fourier Series

Fourier series have many interesting properties, see for example [62]. Two of the most important ones that are useful in theoretical work are:

• Bessel's inequality, which states that projections are smaller than the projected function, and thus:

$$\int_{-\pi}^{\pi} \left[\frac{1}{2} a_0 + a_1 \cos x + \dots + b_N \sin Nx \right]^2 dx \le \int_{-\pi}^{\pi} \left[f(x) \right]^2 dx.$$

• Parseval's formula, which is a direct consequence of the above, and states that

$$\pi[a_0^2 + a_1^2 + b_1^2 + a_2^2 + b_2^2 + \ldots] = \int_{-\pi}^{\pi} [f(x)]^2 dx.$$

This is obtained from Bessels' inequality, which in the limit of $N \to \infty$ becomes an equality; it is simply the definition of the Fourier series. Integrating both sides and using orthogonality we obtain Passeval's formula. This equation is very useful as it connects a vector of coefficients to a function in a unique way.

3.2.3 Differentiation and the Lanczos Filter

Taking derivatives of Fourier series of functions which are discontinuous is not a straightforward matter. Lanczos has devised a clever method in dealing with this, which we explain next.

Let us consider the truncated Fourier series:

$$f_N(x) = \sum_{k=-(N-1)}^{N-1} c_k e^{ikx}$$

and from that the truncation error, which we also call the residual

$$\eta_N(x) = \sum_{k=N}^{\infty} (c_k e^{ikx} + c_{-k} e^{-ikx}) = e^{iNx} \underbrace{\sum_{k=0}^{\infty} c_{N+k} e^{ikx}}_{\rho_N(x)} + e^{-iNx} \underbrace{\sum_{k=0}^{\infty} c_{-N-k} e^{ikx}}_{\rho_{-N}(x)}.$$

Examining the residual more carefully we observe that it consists of two contributions, $\rho_N(x)$ which is slowly varying, and of e^{iNx} which is rapidly varying. Thus, the error in the Fourier series has the structure of a *modulated wave*. This will have consequences when we attempt to take the derivative, the error of which is

$$\eta'_{N}(x) = iNe^{iNx}\rho_{N}(x) + e^{iNx}\rho'_{N}(x) - iNe^{-iNx}\rho_{-N}(x)e^{-iNx}\rho'_{-N}(x).$$

We see from the above that the differentiation of the high frequency wave produces divergent terms proportional to $N, N \to \infty$.

In order to overcome this difficulty, Lanczos introduced a new differentiation operator $\mathcal{D}_{\mathcal{N}}$ so that

$$\mathcal{D}_{\mathcal{N}} \to \frac{d}{dx}, \quad N \to \infty.$$

It is defined by:

$$\mathcal{D}_{\mathcal{N}}f = -\frac{f(x + \pi/N) - f(x - \pi/N)}{2\pi/N}.$$

If we apply, for example, this operator to the function $f(x) = e^{ikx}$ we obtain:

$$\mathcal{D}_{\mathcal{N}}e^{ikx} = i\frac{\sin k/N \pi}{\pi/N}e^{ikx} = \left(\frac{\sin \frac{\pi k}{N}}{k\frac{\pi}{N}}\right) (ik)e^{ikx},$$

where we recognize the term in parenthesis (ik) being the d/dx derivative of e^{ikx} . We can then write

$$\mathcal{D}_{\mathcal{N}} = \sigma_k \frac{d}{dx}$$

where

$$\sigma_k = \frac{\sin \pi k / N}{k \pi / N}$$
 is the Lanczos filter. (3.21)

A plot of the Lanczos filter is shown in figure 3.19. It has the value $\sigma_k = 1$ around the origin and it tends to zero for $k \to N$. Therefore, the action of this filter is to attenuate the contribution of the high frequencies.

We now apply the new differentiation operator to the residual obtained above, i.e.,

$$\mathcal{D}_{\mathcal{N}}\eta_N(x) = -e^{iNx}\mathcal{D}_{\mathcal{N}}\rho_N(x) - e^{iNx}\mathcal{D}_{\mathcal{N}}\rho_{-N}(x).$$

Both $\rho_N(x)$ and $\rho_{-N}(x)$ are smooth functions and their differentiation produces bounded terms. In addition, we observe that there are now no divergent terms (e.g. terms proportional to N) and thus the residual error for the new differentiation tends to zero, and in fact in the limit as $N \to \infty$ the derivative of f(x) is obtained correctly at all points.

Example 1: We first examine the action of the Lanczos filter in the approximation of f(x) by a Fourier series. We choose the function shown in figure 3.20 which has a discontinuity at x = 0 and it is constant at values $\pm 1/2$ in the rest of the interval $xx \in [-\pi, \pi]$.

Its Fourier representation is a sine series of the form

$$y(x) = \frac{2}{\pi} \left(\sin x + \frac{\sin 3x}{3} + \frac{\sin 5x}{5} + \dots \right),$$

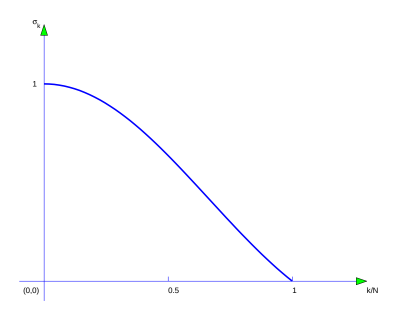


Figure 3.19: Plot of the Lanczos filter of equation (3.21).

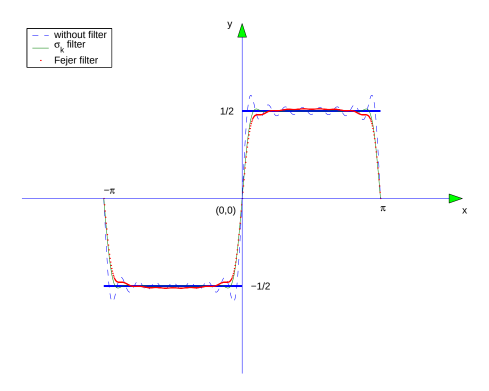


Figure 3.20: Step function and approximations using the Fourier series, the Lanczos-filtered series, and the Fejer construction; N=8.

which we also plot in figure 3.20 for N=8. We observe the Gibbs phenomenon around the origin and also around $x=\pm \pi$, i.e., the loss of convergence, which is not improved for higher values of N. We also plot the filtered Fourier series, i.e., the modified series where each of the coefficients has been multiplied by σ_k ; in this case a smooth representation is obtained. Finally, for comparison we also include the Fejer construction, e.g. the arithmetic mean of the partial sums. We see that this representation is also smooth but not as accurate as the Lancozs-filtered function.

We now turn to the question of differentiation of the above function. Clearly

$$y'(x) = 0,$$

except at x = 0 where the derivative does not exist. The term-wise standard differentiation of y = f(x) gives

$$y'_N(x) = \frac{2}{\pi}(\cos x + \cos 3x + \cos 5x + \ldots)$$

which does not converge at any point x, except at $x = \pm \pi/2$. Thus, the singularity at x = 0 propagates its effect globally!

We then apply the Lanczos recipe, i.e.,

$$y_N'(x) = \frac{2}{\pi} \left[\frac{\sin \frac{\pi}{2N}}{\pi/2N} \cos x + \frac{\sin \frac{3\pi}{2N}}{\frac{3\pi}{2N}} \cos 3x + \dots \right].$$

Now y'(x) converges at all points, except at x = 0, where it grows to ∞ , so that $\int_{-\epsilon/2}^{\epsilon/2} y'_N(x) dx \Rightarrow 1$, which is the $\delta(x)$ function!

Example 2: We can also use the Lanczos recipe to construct Fourier series expansions for functions that are not absolutely square integrable, such as the function y = 1/x. To this end, we consider the function $y = \log x$, where $x \in [0, \pi]$, which is absolutely square integrable, and thus its Fourier series converges (recall Fejer). We then use the Lanczos recipe to take its derivative, and observe that

$$x^{-1} = \frac{d}{dx} \log x$$

thus the Fourier series of

$$x^{-1} = \sum_{k} \sigma_k \frac{d}{dx}$$
 [Fourier series of log x].

3.2.4 Trigonometric Interpolation

Trigonometric interpolation is a powerful tool in representing data and does not suffer from some of the problems we encountered with polynomial interpolation for certain functions, e.g. the Runge function. In fact, we can argue that generally trigonometric interpolation is superior to polynomial interpolation for data on equidistant grids.

We now consider the approximation problem, where we have a number m of basis functions involved less than the number of available data equidistant points n. Let us denote the function values by

$$y_{\alpha} = f(x_{\alpha}),$$

and the representation by

$$\bar{y} = \sum_{k=1}^{m} c_k \varphi_k(x), \ m \le n.$$

As we have already seen in section 3.1.7, in general $y_{\alpha} \neq \bar{y}(\alpha)$, and thus we try to minimize the square of this error. This is the method of least squares, which we apply here first for the case of

$$\varphi_k(x) = e^{ikx}.$$

The coefficients c_k are then computed using the minimization procedure outlined in section 3.1.7 to get:

$$c_k = \frac{1}{\gamma_k} \sum_{\alpha=1}^N y_\alpha \varphi_k^*(x_\alpha)$$

where $\varphi_k^*(x) = e^{-ikx}$ denotes the complex conjugate of $\varphi_k(x)$. Also, γ_k are normalization factors given by

$$\gamma_k = \sum_{\alpha=1}^N \varphi_k(x_\alpha) \varphi_k^*(x_\alpha) .$$

The orthonormality condition implies that

$$\sum_{\alpha=1}^{N} \bar{y}_{\alpha}^{2} = c_{1}^{2} + c_{2}^{2} + \ldots + c_{m}^{2},$$

and thus the error in the approximation is

$$\eta^2 = \sum_{\alpha=1}^N (y_\alpha - \bar{y}_\alpha)^2 = \sum_{\alpha=1}^N y_\alpha^2 - (c_1^2 + c_2^2 + \dots + c_m^2).$$

In summary, assuming that the positions of the data points x_{α} are chosen so that:

$$x_{\alpha} = \alpha \frac{\pi}{N}, \quad a = -N, -(N-1), \dots, (N-1), N$$

then the trigonometric interpolation problem using the complex exponentials as a basis is given by:

$$\begin{cases} \bar{y}(x) = \sum_{k=-m}^{m} c_k e^{ikx}, \ m \le N \\ c_k = \frac{1}{2N} \sum_{\alpha=-N}^{N} y_{\alpha} e^{-ikx\alpha} \end{cases}$$

where in the second sum the end-values should be multiplied by $\frac{1}{2}$.

We can also employ a real basis, either sines or cosines, to perform the approximation depending on the function f(x), i.e., if it is odd or even, respectively.

For example, if the function is odd we employ the *sine series*:

$$f(x) \approx b_1 \sin x + b_2 \sin 2x + \ldots + b_m \sin mx$$

where

$$b_k = \frac{2}{N} \sum_{\alpha=1}^{N-1} y_\alpha \sin k\alpha \frac{\pi}{N}.$$

For m=N we do not have an over-determined problem anymore, and thus we can fit the data exactly to produce a *sine interpolation* instead of approximation. We note here, however, that b_N is undetermined in this case as we effectively have only (N-1) data points. This is because $f(\pi)=0$ in the above approximation since $\sin kx=0$ at $x=\pi$. While the condition f(0)=0 is satisfied automatically because of the odd property, the condition $f(\pi)=0$ may not necessarily be true. This means that a Gibbs phenomenon may develop, which will limit convergence of the approximation to 1/k. To enhance the convergence rate we can construct the following function:

$$g(x) = f(x) - (\alpha + \beta x),$$

which satisfies the conditions $g(0) = g(\pi) = 0$ if

$$\alpha = f(0)$$
 and $\beta = \frac{f(\pi) - f(0)}{\pi}$.

The sine representation of g(x) now leads to convergence rate dictated by $1/k^3$.

We can also employ the *cosine series* to approximate an even function. The expansion now has the form

$$f(x) \approx \frac{1}{2}a_0 + a_1\cos x + \ldots + a_m\cos mx,$$

where

$$a_k = \frac{2}{N} \sum_{\alpha=0}^{N} y_{\alpha} \cos k\alpha \frac{\pi}{N}.$$

Here again in the summation above the end-values are multiplied by 1/2. Notice also that to obtain an *interpolation* instead of an *approximation* the last term is replaced by $1/2a_N \cos Nx$.

Remark: All the above summations can be executed efficiently using the fast Fourier transforms covered in the next section.

3.2.5 Noisy Data

We can also use the properties of trigonometric interpolation to process noisy data, i.e., to remove the major part of the noise which is superimposed on our data; we want to clean up our data! Wavelets is a great way of doing that (see section 3.3) but we can also use Fourier analysis.

We have seen in the previous section that we can construct a sine Fourier series to approximate a function, which converges as $\sim 1/n^3$. On the other hand, any amount of noise, which can be thought of as a pulse of short duration, would lower the convergence of the series significantly, most probably to $\sim 1/n$.

• The key idea is then to differentiate between good data and noise by looking at the decay of coefficients, e.g. $1/n^3$ versus 1/n.

Let us assume that we sample the function $f(x_k)$ over an interval L at the points

$$x_k = 0, h, 2h, \dots, Nh.$$

Then, as before, we construct a new function g(x) based on

$$\begin{cases} g(x) = f(x) - (\alpha + \beta x) \\ g(-x) = -g(x) \end{cases}$$

which leads to

$$g(x) = f(x) - f(0) - \frac{f(L) - f(0)}{L}x.$$

We then expand

$$g(x) = b_1 \sin \frac{\pi}{L} x + b_2 \sin \frac{2\pi}{L} x + \dots$$

where

$$b_k = \frac{2}{N} \sum_{\alpha=1}^{N-1} g(\alpha h) \sin k\alpha \frac{\pi}{N}.$$

From the physical point of view, we may not want any *overtones* contained in the harmonic analysis of f(x) beyond a "cutoff" frequency ν_0 so we set:

$$b_k = 0, \quad k \le m$$

so that

$$\frac{m\pi}{L}x = 2\pi\nu_0 x$$

$$\Rightarrow \boxed{\frac{m}{N} = 2\nu_0 h}.$$

Therefore, we can represent the actual data by

$$g(x) \approx \sum_{k=1}^{m} b_k \sin k \frac{\pi}{L} x,$$

up to wavenumber (or frequency) m determined from the equation in the box. In practice, the lower wavenumbers are also contaminated, i.e., the coefficients $b_k \leq m$, are also influenced by noise but not as much as the upper part of the spectrum which is produced entirely by noise.

3.2.6 Matrix Representation

Let us assume again that the function f(x) is sampled at N equidistant points with spacing $\frac{2\pi}{N}$ between successive neighbors, with the first point at x = 0. The values of the function represented by the truncated Fourier series are given by

$$f_n = \sum_{k=0}^{N-1} c_k e^{ikn\frac{2pi}{N}}, \quad n = 0, 1, \dots, N-1$$

Let

$$w \equiv e^{\frac{2\pi i}{N}}$$

where $w^N = e^{2\pi i} = 1$, so w is the N^{th} root of unity. Then:

$$f_n = \sum_{k=0}^{N-1} c_k w^{kn}.$$

We can then determine the Fourier coefficients c_k by solving the following linear system

$$c_0 + c_1 + c_2 + \dots + c_{N-1} = f_0$$

$$c_0 + c_1 w + c_2 w^2 + \dots + c_{N-1} w^{N-1} = f_1$$

$$c_0 + c_1 w^{N-1} + c_2^{2(N-1)} + \dots + c_{N-1} w^{(N-1)^2} = f_{N-1}$$

which we rewrite in matrix form as

$$\begin{bmatrix} 1 & 1 & \dots & 1 \\ 1 & w & & w^{N-1} \\ \vdots & & & & \\ 1 & w^{N-1} & & w^{(N-1)^2} \end{bmatrix} \begin{bmatrix} c_0 \\ c_1 \\ \vdots \\ c_{N-1} \end{bmatrix} = \begin{bmatrix} f_0 \\ f_1 \\ \vdots \\ f_{N-1} \end{bmatrix}.$$

The coefficients matrix, which we denote by \mathbf{W} , has a special structure similar to the Vandermonde matrix defined in section 3.1.1 of polynomial interpolation. It can be readily inverted, since:

$$\mathbf{W}\mathbf{W}^* = N\mathbf{I} \Rightarrow \mathbf{W}^{-1} = \frac{1}{N}\mathbf{W}^*,$$

where \mathbf{W}^* is the complex conjugate of matrix \mathbf{W} . This can be easily verified by considering any entry of a matrix resulted by multiplying \mathbf{W} with \mathbf{W}^* , and use orthogonality of vectors for the row-column combination.

Note that we can normalize the matrix \mathbf{W} to produce a *unitary matrix* \mathbf{U} by a simple rescaling, i.e.,

$$\mathbf{U} = \frac{\mathbf{W}}{\sqrt{N}}.$$

Circulant Matrices

We first give the definition:

• A circulant matrix is a periodic matrix with constant diagonals of the form

$$\mathbf{C} = \begin{bmatrix} c_0 & c_{N-1} & c_{N-2} & \cdot & c_1 \\ c_1 & c_0 & c_{N-1} & \cdot & c_2 \\ \vdots & \vdots & \vdots & \vdots & \vdots \\ c_{N-1} & c_{N-2} & c_{N-3} & \cdot & c_0 \end{bmatrix}.$$

There is a special relationship between the circular matrix and the Fourier matrix W. In general, the matrix C can be factorized as

$$C = Q\Lambda Q^{-1}$$

where \mathbf{Q} contains the eigenvectors of \mathbf{C} as columns and $\boldsymbol{\Lambda}$ is a diagonal matrix containing the eigenvalues. For a circulant matrix

$$Q = W$$

that is the eigenvectors of the circulant matrix C are the columns of the Fourier matrix W. This can be shown directly by substituting the columns of W into the eigenproblem

$$\mathbf{C}\mathbf{x} = \lambda \mathbf{x}$$
.

3.2.7 The Fast Fourier Transform (FFT)

The fast Fourier transform (FFT) leads to a very smart factorization of the Fourier matrix \mathbf{W} of size N. It produces a product of about $\mathcal{O}(\log_2 N)$ matrices with only about $\mathcal{O}(N\log_2 N)$ non-zero entries (total). Therefore, the cost to compute the discrete Fourier transform or its inverse is $\mathcal{O}(N\log_2 N)$ instead of $\mathcal{O}(N^2)$. The term $\log_2 N$ suggests a fan-in type algorithm (see section 2.3.2) – this is what we will explain in this section.

We start by considering the system

$$C = Wf$$

where **W** is the Fourier matrix of order $N \times N$, **f** is the vector containing the function values at N points, and **C** is the vector of the Fourier coefficients. We recall the Vandermonde structure of the Fourier matrix, for example for N=4 we have

$$\mathbf{W}_{4} = \begin{bmatrix} 1 & 1 & 1 & 1 \\ 1 & w_{4} & w_{4}^{2} & w_{4}^{3} \\ 1 & w_{4}^{2} & w_{4}^{4} & w_{4}^{6} \\ 1 & w_{4}^{3} & w_{4}^{6} & w_{4}^{9} \end{bmatrix} = \begin{bmatrix} 1 & 1 & 1 & 1 \\ 1 & i & -1 & -i \\ 1 & -1 & 1 & -1 \\ 1 & -i & -1 & i \end{bmatrix}$$

where $w_4 = e^{\frac{2\pi i}{4}} = \cos{\frac{2\pi}{4}} + i\sin{\frac{2\pi}{4}} = i$.

The special structure of \mathbf{W}_4 as well as the properties of the principal roots of unity suggest that the matrix \mathbf{W}_8 relates to \mathbf{W}_4 . This is because $w_8^2 = w_4$, and in general

$$w_N^2 = w_M,$$

where $M = \frac{N}{2}$.

For example, the second row of \mathbf{W}_8 is

$$[1 \quad w_8 \quad w_8^2 \quad w_8^3 \quad w_8^4 \quad w_8^5 \quad w_8^6 \quad w_8^7]$$

which we rewrite as

$$\begin{bmatrix} 1 & w_8 & w_4 & w_8w_4 & w_4^2 & w_8w_4^2 & w_4^3 & w_8w_4^3 \end{bmatrix}$$
.

This can be further split into two subvectors of half size. The first one involves only w_4 , i.e.,

$$[1 \quad w_4 \quad w_4^4 \quad w_4^3]$$

formed by taking every other entry, starting with the first one. The second subvector is then:

$$w_8[1 \quad w_4 \quad w_4^2 \quad w_4^3],$$

and it is the same as the first subvector with a pre-factor w_8

We thus see that the rows of \mathbf{W}_8 are closely related to \mathbf{W}_4 , and we can derive similar relations for \mathbf{W}_{10} , \mathbf{W}_{32} and so on. This is the key observation that led Cooley & Tukey in the formulation of FFT in 1965 [17]. It is based on the old idea of **divide-and-conquer**. Assuming that we have a Fourier matrix \mathbf{W}_N where $N=2^m$, then we can split the matrix in successive steps to smaller and smaller matrices. This is only part of the job as we still need to perform matrix-vector multiplies, and finally to compute the Fourier coefficients.

More formally, the n^{th} function value is:

$$f_n = \sum_{k=0}^{N-1} w_N^{nk} c_k = \sum_{k=0}^{M-1} w_N^{2nk} c_{2k} + \sum_{k=0}^{M-1} w_N^{(2k+1)n} c_{2k+1},$$

where $M = \frac{N}{2}$ or

$$f_n = \sum_{k=0}^{M-1} w_M^{nk} c_k^e + w_N^n \sum_{k=0}^{M-1} w_M^{nk} c_k^o$$
 (3.22)

since $w_N^2 = w_M$; also we have denoted by c^e and c^o the even and odd parts of the vector \mathbf{c} , respectively. Notice that the above formula gives us only the *first half* of the Fourier coefficients from $n = 0, \ldots (M-1)$. To obtain the other half we substitute (n+M) instead of n and use the fact that

$$w_M^{k(n+M)} = w_M^{nk} \cdot w_M^{kM} = w_M^{nk}$$

and also

$$w_N^{n+M} = w_N^n \cdot w_N^M = w_N^n \cdot e^{\frac{2\pi i M}{N}} = w_N^n \cdot e^{\pi i} = -w_N^n.$$

Therefore:

$$f_{n+m} = \sum_{k=0}^{M-1} w_M^{nk} c_k^e - w_N^n \sum_{k=0}^{M-1} w_M^{nk} c_k^o.$$
 (3.23)

The matrix-vector product $\mathbf{W} \mathbf{c}$ is then computed from equations (3.22) and (3.23).

The above is the *first level* of splitting into even and odd parts. However, assuming $N = 2^m$, we can repeat this process m more times in a recursive fashion. The cost at each level includes the two matrix-vector multiplies of length M = N/2 plus the multiplication by the prefactor w_N^k in equation (3.22) and (3.23). The basic cost model for this recursion is

$$C(N) = \alpha C\left(\frac{N}{2}\right) + \beta\left(\frac{N}{2}\right),$$

which leads to the estimate

$$C(N) \sim \mathcal{O}(5N \log_2 N)$$

versus $\mathcal{O}(8N^2)$ operations for the direct computation.

Remark 1: Although the FFT is a recursive algorithm, traditional implementations are based on simple nested loops, which have been proved to be more efficient. However, on more modern architectures where cache utilization is important, the recursive procedure is better, see FFTW in section 3.2.8.

Remark 2: In addition to its computational simplicity, the FFT has less round-off error than the direct summation approach.

Remark 3: For N which has the general form

$$N = 2^{\alpha} 3^{\beta} 4^{\gamma} 5^{\delta} 6^{\epsilon}.$$

it is still possible to perform FFTs by splitting into several parts and not only to odd and even components, see [87]. The corresponding cost is

$$\mathcal{O}\left(N\left(5\alpha+9\frac{1}{3}\beta+8\frac{1}{2}\gamma+13\frac{3}{5}\delta+13\frac{1}{3}\epsilon-6\right)\right).$$

Remark 4: The real-to-real transform can be computed in half the number of operations, e.g., in $\mathcal{O}\left(\frac{5}{2}N\log_2 N\right)$. This is done by defining

$$g_k = f_{2k} + i f_{2k+1}, \quad k = 0, 1, \dots M,$$

where M = N/2. Then we take an M-length transform of g_k and set $c_m = c_0$. We extract all the coefficients from

$$c_k = \frac{1}{2}(a\tilde{g}_k + \tilde{g}^*_{M-k}) - \frac{i}{2}e^{\frac{2\pi ik}{N}}[\tilde{g}_k - \tilde{g}^*_{M-k}] \quad k = 0, 1, \dots, M - 1$$

where we have denoted the Fourier coefficients of g with \tilde{g} , and the star denotes complex conjugate.

3.2.8 The Fastest Fourier Transform in the West - FFTW

The FFTW is a C subroutine that contains adaptive software for real and complex onedimensional and multi-dimensional FFTs. Unlike conventional implementations that use loops instead of recursion, in the FFTW explicitly recursive implementation is followed because of theoretical evidence that divide-and-conquer algorithms improve locality.

FFTW is not a new FFT algorithm but it is simply a smart implementation that attempts to exploit a given processor architecture by interacting with its pipeline and its memory hierarchy. It derives its name (the <u>Fastest Fourier Transform in the West</u>) from the fact that in benchmark tests it has proven to be faster than any other publicly available FFT software. FFTW was developed by Frigo & Johnson at MIT in the late 1990s [40]. It is similar in spirit with the ATLAS software described in chapter 2. The key idea is that

the standard Cooley-Tukey algorithm is adapted to the specific hardware employed in the computation.

In order to adapt to any hardware, FFTW runs several diagnostic tests in a pre-processing stage. More specifically, the code is divided into two parts:

- The executor, and
- The codelet generator.

The executor computes the transform by first building a plan. This plan consists of a sequence of instructions that specifies the operation of the executor. The codelets are highly optimized fragments of C code that the executor uses. What exactly combination of codelets will be used depends on the plan, which contains diagnostics and measurements for the particular computer employed. The plan itself is also activated at runtime but it is determined before the actual computation starts. It employs dynamic programming and a cost minimization algorithm, which targets the execution time and not the number of floating point operations.

Therefore, the plan is created first as follows:

```
fftw_plan fftw_create_plan(int n, fft_direction dir, int flags)
```

This function creates a plan for computing an one-dimensional Fourier transform. Here n is the size of the transform, which can be an arbitrary integer, dir = -1 or 1 are flags that denote direction and can be substituted by the aliases $FFTW_FORWARD$ and $FFTW_BACKWARD$, respectively. Also, flags is a boolean, which provides different options. For example, the $FFTW_MEASURE$ finds the optimal plan by computing several FFTs and measuring their execution time. Clearly, this takes time so the first time around when you run FFTW it is expensive. However, the computed plan can be used for subsequent runs which are really fast! An alternative is to use the flag $FFTW_ESTIMATE$ which provides a "best-guess" of an optimal plan without actually running any diagnostic FFTs.

A typical code that uses the complex one-dimensional FFTW looks like:

```
#include<fftw.h>
...
{
fftw_complex in[N], out[N];
fftw_plan p;
...
p=fftw_create_plan(N,FFTW_FORWARD,FFTW_MEASURE);
...
fftw_one(p,in,out);
...
fftw_destroy_plan(p);
}
```

An example of typical performance of the one-dimensional FFTW is provided in figure 3.21 which shows the superior performance of FFTW on the Pentium-4 processor. We also compare the $FFTW_MEASURE$ versus the $FFTW_ESTIMATE$ options and show that even for this relatively small size N of the transform tested, the optimal plan that the former uses makes a substantial difference.

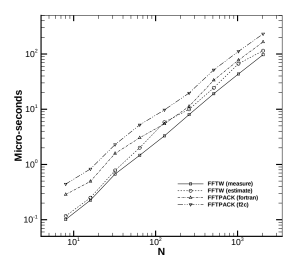


Figure 3.21: Comparisons of FFTW with other FFT implementations on the 1.7GHz Pentium-4 processor. The other FFT implementations are in C and Fortran.

FFTW has also been implemented for parallel environments both for shared memory as well as distributed memory platforms. In particular, the MPI FFTW routines use distributed data, with the array divided according to the rows; that is, each processor gets a subset of the rows of data. FFTW supplies a routine that reports how much data resides on each processor. A typical name of the MPI FFTW routines is

```
#include<fftw_mpi.h>
...
fftwnd_mpi(p, 1, data, NULL, FFTW_NORMAL_ORDER);
...
```

The above uses a complex two-dimensional MPI FFTW.

For details on how to use the FFTW, including the MPI FFTW the reader should consult:

• www.fftw.org,

which is the official internet site maintained by the developers of this adaptive software.

3.3 Wavelet Series Representation

Wavelets are an alternative, relatively recent, family of basis functions to represent a function in the form

$$f(x) = \sum d_{jk}\psi(2^{j}x - k),$$

where d_{jk} are the unknown coefficients. They can be orthonormal or not, smooth or not, compact or not. They allow a decomposition of a function in such a way that its wavenumbers depend on the position x. They can also represent segmented functions, e.g., the multi-step function shown in figure 3.22.

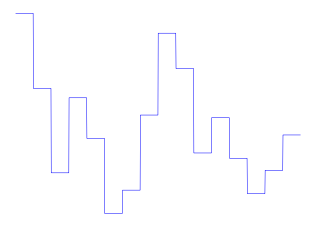


Figure 3.22: Multi-step function.

Such a function is almost impossible to be represented with a Fourier series. However, using wavelets we can construct a special basis, called the *Walsh basis* functions, see figure 3.23, which can approximate the multi-step function very accurately. Specifically, the function shown in figure 3.22 is represented by the first six basis functions of the Walsh family using the following coefficients (from first to sixth):

$$0.3200; \quad 0.9601; \quad 0.7266; \quad 0.4120; \quad 0.7446; \quad 0.2679$$

We note here that the Walsh basis functions are defined in the same interval as the original function, i.e. the basis is global not local, similar to the Fourier representation. In contrast, some of the best wavelets, as we will see in the following, have a compact support, that is they are *local*.

3.3.1 Basic Relations

There are two basic concepts that are used in this field: the scaling function $\phi(x)$ and the wavelet $\psi(x)$. More specifically, we use the integer translations, i.e.,

$$\phi(x-k)$$
 and $\psi(x-k)$, $k \in \mathcal{Z}$

and also, their *dilations*, i.e.,

$$\phi(2^j x)$$
 and $\psi(2^j x)$, $j \in \mathcal{Z}$.

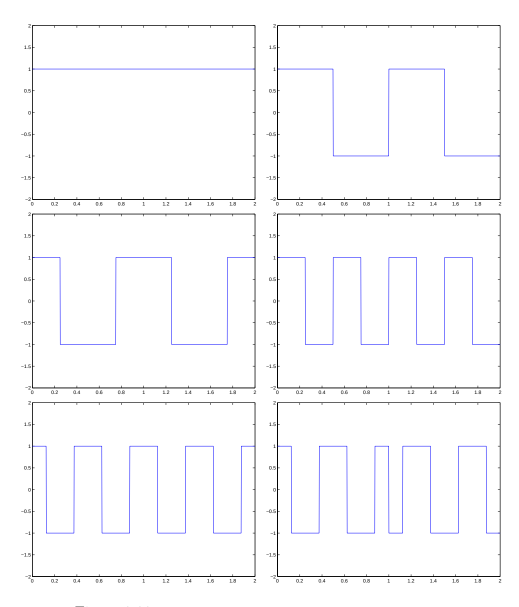


Figure 3.23: First six basis functions of the Walsh family.

Almost always we need both translations and dilations, which we obtain by combining the above concepts, i.e.,

$$\phi(2^jx-k)$$
 and $\psi(2^jx-k)$.

Here, we will refer to 2^{-j} as the scale of level j.

Now using one of the oldest scaling function and wavelets, from the Haar family, we can construct the entire Walsh family. The basic Haar scaling function is a (positive) constant pulse, while the Haar basic wavelet is a combined positive-negative pulse – sometimes we will refer to it as the mother Haar wavelet. In figure 3.24 we show how to construct the first two Walsh basis functions from the Haar basic units.

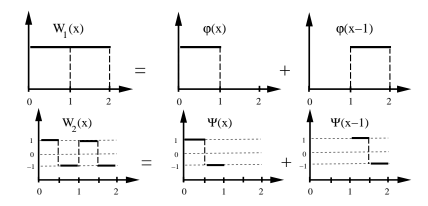


Figure 3.24: The first Walsh basis function (upper) is expressed in terms of the Haar scaling function. The second Walsh basis function (lower) is expressed in terms of the Haar wavelet.

In fact, we can generate the entire Walsh family by using the mother wavelet; we only need the scaling function for the very first Walsh basis. This decomposition of a function into hierarchical components is known as multi-resolution analysis; we will generalize it to arbitrary functions later. Actually, the functions cannot quite be arbitrary – they need to be in L_2 , that is in the space of absolutely square integrable, similar to the condition that Fejer considered in his Fourier construction, see section 3.2.1.

In addition to L_2 , the following two spaces are also very useful in the following:

$$V_j = \left\{ \sum_{k=-\infty}^{\infty} c_{jk} \phi(2^j x - k) : \sum_{k=-\infty}^{\infty} |c_{jk}|^2 < \infty \right\}$$

and

$$W_j = \left\{ \sum_{k=-\infty}^{\infty} d_{jk} \psi(2^j x - k) : \sum_{k=-\infty}^{\infty} |d_{jk}|^2 < \infty \right\}.$$

Based on the above definitions, we have the following important *hierarchy* of spaces:

$$\ldots \subset V_{-1} \subset V_0 \subset V_1 \ldots \subset L_2.$$

We conclude that the subspaces for the scaling functions are nested. Schematically, we have, for example, for the Haar scaling function the picture shown in figure 3.25.

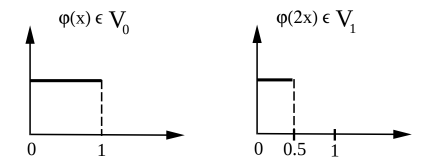


Figure 3.25: Schematic illustration of the spaces V_0 and V_1 .

Using the Haar wavelet we can also infer a relationship between the spaces V_j and W_j . Let us, for example, consider the scaling function $\phi(2x-1)$, then we can write

$$\phi(2x-1) = \frac{1}{2}\phi(x) - \frac{1}{2}\psi(x),\tag{3.24}$$

as shown in the sketch of figure 3.26.

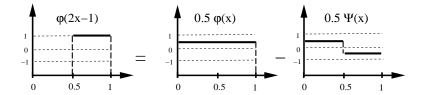


Figure 3.26: Schematic illustration of equation (3.24).

In terms of spaces the above equation can be written as $V_1 = V_0 + W_0$, but it turns out that this can be generalized to:

$$V_{j+1} = V_j + W_j, \quad \forall j \in \mathcal{Z}.$$

In addition, the Haar scaling function and wavelet are orthogonal, which means that the corresponding spaces are orthogonal, i.e.,

$$V_j \perp W_j$$
.

In general, however, the above orthogonality condition may not be valid for all types of wavelets, and instead the following condition is satisfied

$$V_j \cap W_j = \emptyset \quad \forall_j \in \mathcal{Z}$$

that is, the intersection of the scaling function and wavelet spaces is the empty set.

The fact that there is hierarchy in spaces $V_j \subset V_{j+1}$, and also that $W_j \subset V_{j+1}$, leads to the following two fundamental equations, also known as two-scale relations:

$$\phi(x) = \sum_{k} p_k \phi(2x - k) \tag{3.25}$$

and

$$\psi(x) = \sum_{k} q_k \phi(2x - k). \tag{3.26}$$

Thus, the wavelet can be obtained from the scaling function. For special cases, e.g. orthonormal wavelets, we have

$$\psi(x) = \sum_{k} (-1)^{k} p_{1-k} \phi(2x - k), \tag{3.27}$$

which will be discussed in detail later.

Examples:

- For the Haar wavelet $\psi(x) = \phi(2x) \phi(2x 1)$ where $\phi(x)$ is the Haar scaling (box) function, we have: $p_0 = 1$ and $p_1 = 1$.
- For the hat wavelet, corresponding to hat function $N_2(x)$: $\psi(x) = \phi(2x) \frac{1}{2}\phi(2x-1) \frac{1}{2}\phi(2x+1)$, the coefficients are: $p_0 = 1/2$, $p_1 = 1$ and $p_2 = 1/2$ from equation (3.27).

3.3.2 Dilation Equation

The nested property of the spaces V_i leads to the two-scale dilation equation of the form

$$\phi(x) = \sum_{k} p_k \phi(2x - k).$$

For example, for the Haar scaling function, we have that

$$\phi(x) = \phi(2x) + \phi(2x - 1)$$

since

$$p_0 = p_1 = 1.$$

 $p_k = 0, |k| > 1.$

Let us also consider the new scaling function we introduced above

$$\phi(x) = N_2(x) = \int_0^1 N_1(x-t)dt,$$

where $N_1(x)$ is the box Haar function. Then, the dilation equation is

$$\phi(x) = \frac{1}{2}\phi(2x) + \phi(2x - 1) + \frac{1}{2}\phi(2x - 2)$$

since

$$p_0 = \frac{1}{2}$$
; $p_1 = 1$; $p_2 = \frac{1}{2}$

as we have already seen above for the wavelet $\psi(x)$, and

$$p_k = 0, \quad |k| > 2.$$

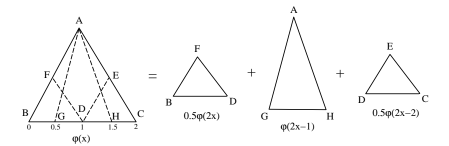


Figure 3.27: Solution of the dilation equation for the hat wavelet $N_2(x)$.

Schematically, this can be justified as shown in figure 3.27.

So far we have shown that the coefficients p_k form a finite set, typically very small consisting of the first few non-zero values. This shows *compact support*, but there are cases that p_k are all non-zeros. How to obtain them is not always easy. Dilation equations are interesting on their own right, and here we review three different approaches of solving them, see [85]:

Iteration for Dilation Equation

Here we iterate on the two-scale dilation equation

$$\phi_n(x) = \sum_k p_k \phi_{n-1}(2x - k)$$

where $\sum_k p_k = 2$. This condition is based on the normalization $\int \phi(x)dx = 1$ and can be derived by multiplying the dilation equation by 2 and integrating, i.e.,

$$2\int \phi dx = \sum_{k} p_k \int \phi(2x - k) d(2x - k) \Rightarrow \sum_{k} p_k = 2.$$

To initialize we need to set the value of ϕ_0 for n = 1. For Daubechies wavelets (see below) we set ϕ_0 equal to the Haar scaling function $N_1(x)$. An appropriate initial choice is made for other families of wavelets.

- If $p_0 = 2$ then $p_k = 0$, |k| > 1 and the iteration above converges to a delta function.
- If $p_0 = p_1 = 1$ the iteration converges to the Haar scaling function, i.e., we have invariance, since $\phi_n = \phi_0$.
- If $p_0 = \frac{1}{2}$, $p_1 = 1$; $p_2 = \frac{1}{2}$ we recover the hat function $N_2(x)$ as demonstrated in figure 3.27.
- If $p_0 = \frac{1}{8}$; $p_1 = \frac{4}{2}$; $p_2 = \frac{2}{3}$; $p_3 = \frac{1}{2}$; $p_4 = \frac{1}{8}$ the iteration converges to the cubic B-spline $N_4(x)$. This can be written as

$$N_4(x) = \int_0^1 N_3(x-t)dt,$$

where $N_3(x)$ is the quadratic B-spline, which in turn is

$$N_3(x) = \int_0^1 N_2(x-t)dt$$

and $N_2(x)$ is the hat function of figure 3.27.

Fourier Transform for Dilation Equation

We can also obtain the scaling function by Fourier transforming the two-scale difference equation, i.e.,

$$\hat{\phi}(z) = \sum_{k} p_{k} \int \phi(2x - k)e^{izx} dx$$

$$= \sum_{k} p_{k} e^{ikz/2} \int \phi(2x - k)e^{i\frac{2x - k}{2}z} \frac{1}{2} d(2x - k)$$

$$= \left(\frac{1}{2} \sum_{k} p_{k} e^{i\frac{kz}{2}}\right) \cdot \int \phi(t)e^{itz/2} dt$$

$$\equiv P\left(\frac{z}{2}\right) \cdot \hat{\phi}\left(\frac{z}{2}\right)$$

where

$$P(z) = \frac{1}{2} \sum_{k} p_k e^{ikz} \tag{3.28}$$

is the transfer function and P(0) = 1. We have thus obtained

$$\hat{\phi}(z) = P\left(\frac{z}{2}\right) \cdot \hat{\phi}\left(\frac{z}{2}\right)$$

where also $\hat{\phi}(0) = 1 = \int \phi(x) dx$ by the normalization condition.

Using the above equation for $\hat{\phi}$ we have that:

$$\hat{\phi}\left(\frac{z}{2}\right) = P\left(\frac{z}{4}\right) \cdot \hat{\phi}\left(\frac{z}{4}\right)$$

and so on for $\hat{\phi}\left(\frac{z}{4}\right)$, $\hat{\phi}\left(\frac{z}{8}\right)$, etc. Thus,

$$\hat{\phi}(z) = \Pi_{n=1}^{N} P\left(\frac{z}{2^{n}}\right) \cdot \hat{\phi}\left(\frac{z}{2^{N}}\right),\,$$

and for

$$N \to \infty, \quad \frac{z}{2^N} \to 0 \Rightarrow \hat{\phi}\left(\frac{2}{2^N}\right) \to 1.$$

Thus

$$\hat{\phi}(z) \to \prod_{n=1}^{\infty} P\left(\frac{z}{2^n}\right),$$

and the inverse transform of $\hat{\phi}(z)$ gives the scaling function $\phi(x)$.

For example:

- Let $p_0 = 2$, then $P(z) = 1 \Rightarrow \hat{\phi}(z) = 1 \Rightarrow \phi(x) = \delta(x)$, the Dirac function.
- Let $p_0 = p_1 = 1$, so $P(z) = \frac{1}{2} [1 + e^{iz}]$. Then

$$P\left(\frac{z}{2}\right)P\left(\frac{z}{4}\right)P\left(\frac{z}{8}\right)\dots P\left(\frac{z}{2^{N}}\right) = \frac{1}{2}(1+e^{iz/2})$$
$$\cdot \frac{1}{2}(1+e^{iz/4})\cdot \frac{1}{2}(1+e^{iz/8})\dots \frac{1}{2}(1+e^{\frac{iz}{2^{N}}})$$
$$\Rightarrow \frac{1-e^{iz}}{(-iz)} = \int_{0}^{1} e^{izx} dx.$$

The inverse transform then gives $\phi(x) = N_1(x)$, which is the Haar scaling function.

Recursion for Dilation Equation

The idea here is that we can construct $\phi(x)$ numerically, point-by-point. That is, assuming that we know $\phi(k)$ (at the integer points), then we can use the dilation equation to obtain ϕ at $\frac{k}{2}$, then at $\frac{k}{4}$ and ultimately at $\frac{k}{2^N}$. The question then becomes how do we obtain the values $\phi(k)$? These are obtained as solutions of the eigenvalue problem constructed from the algebraic equations at points

$$x = 0, \pm 1, \pm 2, \dots$$

within the range of support. Specifically, we set up an eigensystem

$$oldsymbol{\phi} = -\mathbf{P}oldsymbol{\phi}$$

where

$$\boldsymbol{\phi} = [\dots \phi(1), \phi(2), \phi(3) \dots]^T$$

and **P** is a relatively sparse matrix containing the coefficients p_k .

We will demonstrate this approach later for the Daubechies wavelets.

3.3.3 Discrete Wavelet Transform: Mallat's Algorithm

Let $f(x) \in L_2$, and we want to approximate it by

$$f_n(x) \approx f(x), \quad f_n(x) \in V_n.$$

Then, given the hierarchy of spaces and the fact that

$$V_{j+1} = V_j + W_j,$$

a unique decomposition of $f_n(x)$ in terms of its components $g_j(x) \in W_j$ exists. In particular, we can write

$$f_n(x) = \underbrace{f_{n-m}(x)}_{\in V_{n-m}} + \left\{ \underbrace{g_{n-m}(x) + \ldots + g_{n-1}(x)}_{\in W_{n-m} \dots W_{n-1}} \right\},$$
(3.29)

where m is a positive integer that depends on the filter size.

That is, the function is decomposed into one component (the "blur" or "DC") that can be represented in the space V_{n-m} of scaling functions plus contributions at various scales that can be represented by wavelets. This decomposition makes sense as the "blur" represents the "mean" while the wavelets represent the "fluctuations," since they have zero mean by construction. The wavelet contributions correspond to the following wavelet series

$$g_j(x) = \sum_k d_{jk} \psi_{jk}(x).$$

Most but not all wavelets are orthogonal, so here we extend the concept of orthogonality to bi-orthogonality. To this end, we introduce the dual wavelet $\tilde{\psi}(x) \in L_2$, where the following inner product expresses the bi-orthogonality condition:

$$<\psi_{jk}(x), \tilde{\psi}_{jk}(x)> = \int_{-\infty}^{\infty} \psi_{jk}(x) \tilde{\psi}_{\ell m}^{*}(x) dx = \delta_{j\ell} \delta_{km},$$

where the star * denotes complex-conjugate – some wavelets are complex functions. Also, the following normalization condition is usually employed:

$$\psi_{jk} = 2^{j/2} \psi(2^j x - k)$$

to relate the wavelet basis functions to mother wavelet.

With the above definitions, we can now obtain the discrete wavelet transform in the general case. To this end, we re-write equation (3.29)

$$g_i(x) = f_n(x) - [f_i(x) + g_{n-1}(x) + \dots + g_{i+1}(x)],$$

and taking the inner product, we get

$$\langle g_i(x), \tilde{\psi}(2^j x - k) \rangle = \langle f_n(x), \tilde{\psi}(2^j x - k) \rangle,$$

as the rest of the terms drop out due to bi-orthogonality. Thus,

$$d_{jk} = \langle g_j(x), \tilde{\psi}_{jk}(x) \rangle = \langle f_n(x), \tilde{\psi}_{jk}(x) \rangle$$

$$= 2^{j/2} \int_{-\infty}^{\infty} f_n(x) [\tilde{\psi}(2^j x - k)]^* dx,$$
(3.30)

which is the discrete wavelet transform of $f_n(x)$.

We now turn to searching for algorithms that enable us to decompose or reconstruct the approximation $f_n(x)$ fast. Using the scaling function as basis, we can write

$$f_n(x) = \sum_k c_{nk} \phi(2^n x - k).$$

Also, from the relation $V_1 = V_0 + W_0$, we have that

$$\phi(2x - m) = \sum_{k} [a_{m-2k}\phi(x - k) + b_{m-2k}\psi(x - k)] \quad \forall m \in \mathcal{Z}.$$
 (3.31)

Similarly, from $V_n = V_{n-1} + W_{n-1}$, we conclude that

$$\underbrace{f_n}_{\in V_n}(x) = \underbrace{f_{n-1}}_{\in V_{n-1}}(x) + \underbrace{g_{n-1}(x)}_{\in W_{n-1}},$$

where

$$f_{n-1}(x) = \sum_{k} c_{n-1,k} \phi(2^{n-1}x - k)$$
 (3.32)

$$g_{n-1}(x) = \sum_{k} d_{n-1,k} \psi(2^{n-1}x - k).$$
 (3.33)

The above relations are useful in helping us to obtain the set of coefficients

$$\{c_{nk}\}\$$
and $\{d_{nk}\}\$ from $\{c_{n-1,k}\}\$ and $\{d_{n-1,k}\}.$

Comparing for example, equations (3.31) with (3.32) and (3.33), we obtain:

$$\begin{cases} c_{n-1,k} = \sum_{m} a_{m-2k} c_{n,m} \\ d_{n-1,k} = \sum_{m} b_{m-2k} c_{n,m} \end{cases}$$

The above operations represent discrete convolution and can be performed fast. More precisely, there should be a change sign, between e.g., $a_{m-2k} \to a_{2k-m}$; also, only the double-indexed entries should be sampled. The above procedure is known as the Mallat algorithm [69]. It can compute all coefficients in $\mathcal{O}(N)$ arithmetic operations, i.e., even faster than the FFT with $\mathcal{O}(N \log_2 N)$ arithmetic operations.

This analysis gives the decomposition of $f_n(x)$. In terms of reconstruction i.e., to obtain $f_n(x)$ from $\{f_{n-1}(x) \text{ and } g_{n-1}(x)\}$, we start with the two-scale difference equation

$$\phi(x) = \sum_{k} p_k \phi(2x - k)$$

$$\psi(x) = \sum_{k} q_k \phi(2x - k)$$

and thus

$$c_{n,k} = \sum_{m} [p_{k-2m}c_{n-1,m} + q_{k-2,m}d_{n-1,m}],$$

where again the aforementioned corrections are needed to make the above an exact convolution, which can be computed fast. This is the *inverse Mallat's algorithm*.

3.3.4 Some Orthonormal Wavelets

Orthonormal wavelets form a very special class of basis functions. They satisfy the condition among spaces at different scales

$$W_i \perp W_j, \quad i \neq j.$$

Also, the space of finite-energy function L_2 can be represented by an orthogonal sum of the subspaces W_i , which we express as

$$L_2 = \bigoplus_{-\infty}^{\infty} W_j$$
.

It is read as "the direct sum of W_j " and we refer to it as the wavelet decomposition of L_2 . The condition of orthonormality is expressed best by the Fourier transform. That is, if

$$\sum_{k=-\infty}^{\infty} |\hat{\phi}(\omega + 2\pi k)|^2 = 1$$

then the function $\phi(x) \in L_2$ is orthonormal, see [16]. One can show that the orthogonality condition between W_j and V_j , and correspondingly between $\psi(x)$ and $\phi(x)$, gives

$$q_k = (-1)^k p_{1-k},$$

that is, the wavelet coefficients are determined solely by the scaling function coefficients, so

$$\psi(x) = \sum_{k} (-1)^k p_{1-k} \phi(2x - k). \tag{3.34}$$

Based on that, the orthogonal decomposition $V_1 = V_0 \oplus W_0$ leads to:

$$\phi(2x - m) = \sum_{k} \frac{1}{2} [p_{m-2k}\phi(x - k) + (-1)^m p_{2k-m+1}\psi(x - k)],$$

which is a decomposition relation of function $\phi(x)$ and $\psi(x)$.

Next, we provide some examples of orthonormal wavelets.

Haar Wavelet

The oldest of all is the Haar wavelet, which we have already studied. It corresponds to the box scaling function with $p_0 = p_1 = 1$ and $p_k = 0$ for all the other coefficients (k > 1). Using equation (3.34), we can write:

$$\psi(x) = \phi(2x) - \phi(2x - 1)$$
$$= N_1(2x) - N_1(2x - 1).$$

Shannon Wavelet

The Shannon wavelet uses the scaling function

$$\phi(x) = \frac{\sin \pi x}{\pi x}$$

which is identical to the Lanczos filter, see section 3.2.3. The coefficients of the Shannon scaling function are

$$p_k = \begin{cases} 1, & k = 0\\ (-1)^{\frac{k-1}{2}} \frac{2}{\pi k}, & k = 2m - 1\\ 0, & k = 2m \end{cases}$$

Based on equation (3.34), we then obtain

$$\psi(x) = \sum_{k} (-1)^{k} p_{1-k} \phi(2x - k)$$

which leads to equation

$$\psi(x) = \frac{\sin 2\pi x - \cos \pi x}{\pi (x - \frac{1}{2})}.$$
(3.35)

A plot of the Shannon wavelet is shown in figure 3.28 .

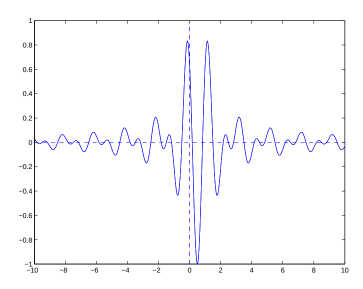


Figure 3.28: Shannon wavelet.

Orthonormal Spline Wavelets

We have discussed splines in section 3.1.6 and here we examine how we can use them as orthonormal wavelets. In general, the translates of B-splines are not orthogonal, and thus a special orthonormalization procedure needs to be employed. The result is that the orthonormalized splines will no longer be polynomials.

Following Chui [16], we first define the B-splines recursively from

$$N_m(x) = \int_{-\infty}^{\infty} N_{m-1}(x-t)N_1(t)dt = \int_0^1 N_{m-1}(x-t)dt,$$

where $N_1(t)$ is the box function. The two-scale equation for the B-splines is

$$N_m(x) = \sum_{k=0}^{m} 2^{-m+1} \binom{m}{k} N_m(2x-k),$$

and $N_m(x)$ is an (m-1)-order polynomial with continuity (smoothness)

$$N_m(x) \in C^{m-2}$$
.

Plots are shown in figure 3.29 for various orders; they are smooth but not orthonormal.

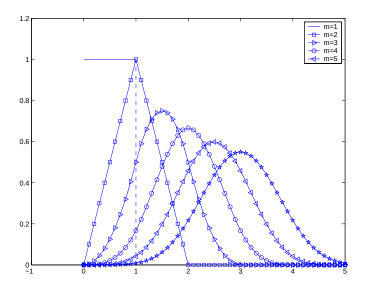


Figure 3.29: B-splines (cardinal) from first- to fifth-order.

In order to orthonormalize $N_m(x)$ to produce $N_m^{\perp}(x)$, Chui has introduced the complex conjugate of the reflection of $N_m(x)$, i.e., $N_m^*(-x)$ as the dual function. To this end, we have

$$N_{2m}(m+x) = \int_{-\infty}^{\infty} N_m(t) N_m^*(t-x) dt = \frac{1}{2\pi} \int_0^{2\pi} \left\{ \sum_{k=-\infty}^{\infty} |\hat{N}_m(\omega + 2\pi k)|^2 \right\} e^{i\omega x} d\omega,$$

where the last term refers to the Fourier transform of $N_m(x)$.

Now defining

$$E_m(z) \equiv \sum_{k=-m+1}^{m-1} N_{2m}(m+k)z^k$$

the orthonormalization procedure leads to

$$N_m^{\perp}(\omega) = \frac{\hat{N}_m(\omega)}{[E_m(z^2)]^{1/2}}, \quad z = e^{-i\frac{\omega}{2}}.$$

Also, the transfer function can be expressed via $E_m(z)$, see [16]:

$$P(z) \equiv \frac{1}{2} \sum_{k} p_k z^k = \left(\frac{1+z}{2}\right)^m \left[\frac{E_m(z)}{E_m(z^2)}\right]^{1/2}.$$

Finally, the m^{th} -order orthonormal wavelet, also known as **Battle-Lemarié wavelet** is given by

$$\psi_m(x) = \sum_{k} (-1)^k p_{1-k} N_m^{\perp} (2x - k). \tag{3.36}$$

In practice, it is more convenient to define the Battle-Lemarié wavelet in terms of its Fourier transform:

$$\hat{\psi}_m(\omega) = -\left(\frac{4}{i\omega}\right)^m \left[\sin^{2m}\left(\frac{\omega}{4}\right)\right] z \cdot \left[\frac{E_m(-z)}{E_m(z)E_m(z^2)}\right]^{1/2},$$

where $z = e^{-i\omega/2}$.

A plot of the Battle-Lemarié wavelet is shown in figure 3.30 for m=2.

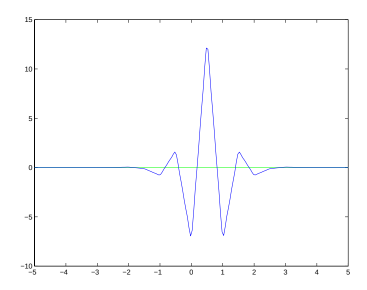


Figure 3.30: Battle-Lemarié wavelet of order m=2 (linear).

The Daubechies Wavelets

The Daubechies wavelets are very special because they are very compact but also orthonormal – no special procedure is needed to orthonormalize them. They can be defined by means of the transfer function

$$P(z) = \frac{1}{2} \sum_{n=0}^{N} p_n z^n = \left(\frac{1+z}{2}\right)^m S_{N-m}(z),$$

where $S_{N-m}(z)$ is a polynomial of degree (N-m) and $S_{N-m}(1)=1$; the latter implies that P(1)=1. How to compute S_{N-m} is not trivial and the interested reader is referred to [22]. The inverse Fourier transform of the product

$$\hat{\phi}_m(\omega) = \prod_{j=1}^{\infty} P\left(e^{-i\omega/2^j}\right)$$

gives the Daubechies scaling function.

An alternative way is to use iteration, as discussed earlier, e.g.,

$$\phi_{m;j+1}(x) = \sum_{k=0}^{N} p_k \phi_{m;j}(2x - k)$$

starting with $\phi_{m;0} = N_2(x)$, the linear B-spline, and iterate to a converged solution to obtain $\phi_m(x)$.

Having obtained the scaling function, the Daubechies wavelets can then be obtained from

$$\psi_m(x) = \sum_{k=-N+1}^{1} (-1)^k p_{1-k} \phi_m(2x-k).$$

The Daubechies scaling function and wavelet of order m=4 are shown in figure 3.31, and of order m=7 in figure 3.32.

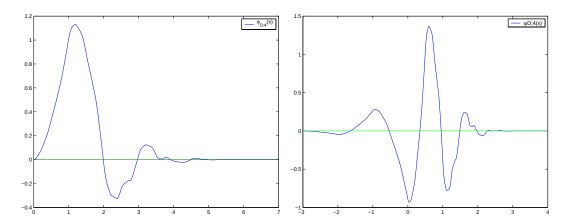


Figure 3.31: Scaling function (left) and wavelet (right) of the Daubechies family; m=4.

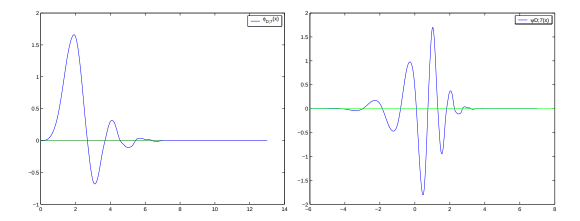


Figure 3.32: Scaling function (left) and wavelet (right) of the Daubechies family; m = 7.

As we see from the plots, the Daubechies wavelets are compact but not very smooth; in fact, here polynomial smoothness has been sacrificed for compactness! The smoothness

of the Daubechies wavelets increases with the order but only by about half derivative each time; e.g., ψ_4 is $C^{0.55}$, i.e., fractionally continuous. In contrast, we recall that the B-splines with m=4 are C^2 continuous.

More specifically, the Daubechies wavelet of fourth-order (m = 4) is defined by the coefficients:

$$p_0 = \frac{1+\sqrt{3}}{4}; \quad p_1 = \frac{3+\sqrt{3}}{4}$$

$$p_2 = \frac{3 - \sqrt{3}}{4}; \quad p_3 = \frac{1 - \sqrt{3}}{4}.$$

We can now use these coefficients to construct $\phi_4(x)$, the scaling function, by recursion, as discussed earlier. First, we set up an eigenvalue problem by applying the dilation equation at the integer points x = 1 and x = 2 in the open interval $x \in (0,3)$:

$$\phi_4(1) = \frac{1+\sqrt{3}}{4}\phi_4(2) + \frac{3+\sqrt{3}}{4}\phi(1)$$

$$\phi_4(2) = \frac{3-\sqrt{3}}{4}\phi_4(2) + \frac{1-\sqrt{3}}{4}\phi_4(1).$$

The eigenvalues of the 2×2 eigensystem are 1 and 1/2; the former leads to

$$\phi_4(1) = \frac{1+\sqrt{3}}{2}$$
 and $\phi_4(2) = \frac{1-\sqrt{3}}{2}$.

Having obtained these values, we now set $x = \frac{1}{2}, \frac{1}{2^2}, \frac{1}{2^3}$ and so on at all dyadic points to compute values of $\phi_4(x)$ at many other points.

The accuracy of the approximation depends on the order p of zeros of the transfer function

$$P(z) = \frac{1}{2} \sum_{k} p_k e^{ikz}.$$

This turns out to be equivalent to the following condition for the coefficients p_k :

$$\sum (-1)^k k^m p_k = 0, \quad m = 0, 1, \dots, p - 1$$

Correspondingly, the first p moments of the wavelet $\psi(x)$ are zero, i.e.,

$$\int x^m \psi(x) dx = 0, \quad \forall m = 0, \dots p - 1.$$

The fourth-order Daubechies scaling function $\phi_4(x)$ has p=2 in contrast with the cubic spline which has p=4. The corresponding error bound is

$$\parallel f(x) - \sum p_k \phi(2^j x - k) \parallel \leq C \cdot 2^{-jp} \parallel f^{(p)}(x) \parallel$$

that is, the error is $\mathcal{O}(h^p)$, where $h = 2^{-j}$.

3.4 Back to Parallel Computing: Send and Receive

The information that we provided to you in chapter 2 was accurate but incomplete. In this section we want to supply more information about sending and receiving in MPI. After this section, throughout the book, we will begin to directly integrate MPI concepts into the lessons.

Recall from our previous discussion that for each call to MPI_Send , there should be a call to MPI_Recv . Below we give the declarations (which can be found in **mpi.h** on your parallel computer) for both MPI_Send and MPI_Recv . Let us investigate each one separately. Throughout the book, we will use the format below for presenting MPI functions. The return value and name will be provided followed by the argument list (both variable type and variable name) and comments denoting whether the variable is intended to act as input (in) to or output (out) of the function.

• MPI_Send

```
int MPI_Send(
        void*
                         message
        int
                         count
                                       in
                                       in
        MPI_Datatype
                        datatype
                                       in
                         dest
        int
                         tag
                                       in
        MPI_Comm
                         comm
```

The first item in the argument list is the starting *address* from which the data item is to be retrieved. You can translate this as "the *pointer* variable." The second piece of information is the count, which is the number of items to send. The third item is the data type, which can be among the following:

- MPI_INT
- MPI_FLOAT
- MPI_DOUBLE
- MPI_CHAR

See Appendix B.2 for a full listing of all the allowable variable types. We only provide here the four basic types that we have introduced thus far. The fourth argument is the processor identification number to which the message is to be sent. This processor id is the integer that is obtained locally on the processor by calling $MPI_Comm_rank()$ (recall, in the previous example, we stored the local process number in the variable mynode). The fifth argument is a tag, which is an integer used to delineate between successive messages. Suppose that you were going to send messages one right after the other to and from one processor to the next. Then you would assign two different tag numbers so that you can guarantee the ordering that the processor will receive the information. The final argument provides information concerning which processes are within the current communication group. For our purposes, this argument will always be set to MPI_COMM_WORLD .

Now, on the receiving side, we have the following:

• MPI_Recv

```
int MPI_Recv(
        void*
                        message
                                       out
        int
                        count
                                       in
        MPI_Datatype
                                       in
                       datatype
        int
                        source
                                       in
                                       in
        int
                        tag
        MPI_Comm
                        comm
        MPI_Status*
                        status
```

Again, the first item in the argument list is the starting address into which the data is to be placed, the second argument is the count, and the third argument identifies the type of information to be sent. Instead of specifying the destination, the fourth argument is the source of the information, which is the process identifier of the source of the message to be received. The fifth and sixth arguments were just discussed above. The final argument is a special type of variable which supplies status information. Examining status.MPI_ERROR can provide the programmer with error information when something goes wrong. For more information concerning the content and use of MPI_Status, we refer the reader to [73].



For the purposes of illustration, let us examine the following MPI code. We want to create an array on each process, but to only initialize it on process 0. Once the array has been initialized on process 0, then it is sent out to each process.

```
#include<iostream.h>
#include<mpi.h>

int main(int argc, char * argv[]){
   int i;
   int nitems = 10;
   int mynode, totalnodes;
   MPI_Status status;

   double * array;

   MPI_Init(&argc,&argv);
   MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
   MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

   array = new double[nitems];

   if(mynode == 0) {
      for(i=0;i<nitems;i++)
         array[i] = (double) i;
}</pre>
```

```
if(mynode==0)
  for(i=1;i<totalnodes;i++)
    MPI_Send(array,nitems,MPI_DOUBLE,i,1,MPI_COMM_WORLD);
else
  MPI_Recv(array,nitems,MPI_DOUBLE,0,1,MPI_COMM_WORLD, &status);

for(i=0;i<nitems;i++){
  cout << "Processor " << mynode;
  cout << ": array[" << i << "] = " << array[i] << endl;
}

delete[] array;

MPI_Finalize();
}</pre>
```

We draw your attention to the following:

- We first do the MPI initialization and information gathering calls described in the previous chapter (MPI_Init, MPI_Comm_size and MPI_Comm_rank).
- We then create, on each process, an array using dynamic memory allocation.
- On process 0 only (i.e., mynode == 0), we initialize the array to contain the ascending index values. Observe that we *explicitly cast* the integer index value i to a double precision number to be stored in array[i].
- On process 0, we proceed with (totalnodes-1) calls to MPI_Send.
- On all other processes other than 0, we call MPI_Recv to receive the sent message.
- On each individual process, we print the results of the sending/receiving pair.
- On each individual process, we deallocate the dynamic memory that we had allocated.

Though this program is simple, it contains most of the necessary components for getting up and running with MPI.

Let us conclude this section by stating a collection of remarks which will be useful both in understanding the program above and in understanding MPI programs to follow.

- Whenever you send and receive data, MPI assumes that you have provided nonoverlapping positions in memory. This remark will be extremely relevant when we study collective routines in the chapters to follow. We will point this fact out again.
- As we pointed out in the previous chapter, MPI_COMM_WORLD is referred to as a **communicator**. In general, a communicator is a collection of processes that can send messages to each other. MPI_COMM_WORLD is pre-defined in all implementations of MPI, and it consists of all MPI processes running after the initial execution of the program.
- In the send/receive discussions above, we were required to use a tag. The tag variable is used to distinguish upon receipt between two messages sent by the same process. The order of sending does not necessarily guarantee the order of receiving. Tags are used to distinguish between messages. MPI allows the tag MPI_ANY_TAG which can be used by MPI_Recv to accept any valid tag from a sender. You cannot use MPI_ANY_TAG in the MPI_Send command, however.
- Similar to the MPI_ANY_TAG wildcard for tags, there is also an MPI_ANY_SOURCE wildcard that can also be used by MPI_Recv. By using it in an MPI_Recv, a process is ready to receive from any sending process. Again, you cannot use MPI_ANY_SOURCE in the MPI_Send command. There is no wildcard for sender destinations.
- When you pass an array to MPI_Send/MPI_Recv , it need not have exactly the number of items to be sent it must have greater than or equal to the number of items to be sent. Suppose, for example, that you had an array of 100 items, but you only wanted to send the first ten items. You can do so by passing the array to MPI_Send and only stating that ten items are to be sent.

Throughout the remainder of this book we will building upon the basic foundation introduced here to accomplish more serious tasks of parallel scientific computing.

3.5 Homework Problems

3.5.1 Homework Problems for Section 3.1

- 1. In the previous chapter's exercises, a recursive relationship was given for generating the Fibonacci sequence. Write a recursively called function which takes as input at least the maximum number of terms to generate (your function may have more inputs) and prints the Fibonacci sequence to the screen.
- 2. The Vandermonde matrix of rank n is defined by

$$V_n \equiv \begin{bmatrix} 1 & x_0 & x_0^2 & \dots & x_0^n \\ 1 & x_1 & x_1^2 & \dots & x_1^n \\ \vdots & \vdots & \vdots & \vdots & \vdots \\ 1 & x_n & x_n^2 & \dots & x_n^n \end{bmatrix}$$

Prove the following result for its determinant

$$\det V_n = d_0 \ d_1 \dots d_{n-1}$$

where

$$d_k \equiv (x_{k+1} - x_k)(x_{k+1} - x_{k-1})\dots(x_{k+1} - x_0).$$

3. Compute (by hand) the second-order polynomial $p_2(x)$ so that

$$p_2(0) = 3, p_2(1) = 2, p_2(3) = 5,$$

using the Vandermonde, Newton, and Lagrange approaches.

4. Construct the third-order Lagrange polynomials for the following functions in the interval $[x_0, x_n]$

(a)
$$f(x) = \ln x$$
, $x_0 = 1$, $x_1 = 1.1$, $x_2 = 1.3$, $x_3 = 1.4$

(b)
$$f(x) = 5\cos x + 3\sin x$$
, $x_0 = 0$, $x_1 = 0.25$, $x_2 = 0.5$, $x_3 = 1.0$

- 5. Consider the Runge function with $\epsilon = 10^{-n}$, where n = 1 (case A) and n = 5 (case B). How many Chebyshev modes are required to approximate it with 10% (engineering) or 1% (scientific) accuracy for both cases A and B?
- 6. Consider the function $f(x) = \sin 2\pi x$, where $x \in [-1, 1]$.
 - (a) Write a program for computing the interpolating polynomial $p_n(x)$ at the points $x_i^n = -1 + \frac{2i}{n}$ with $i = 0, 1, \ldots, n$ for n = 6, 12, 18, 24 using the Lagrange approach. Plot the maximum point-wise error versus n.
 - (b) Do the same as in (a) but use Chebyshev distribution of points to represent f(x).

- (c) Repeat (a) and (b) using the Newton approach.
- (d) Write a program which asks the user which approach to use and which distribution to use. Integers may be used to delineate between approaches and between point distributions (i.e., Lagrange = 0, Newton = 1, etc.)
- 7. Estimate the number of multiplications required to compute $f(x) = \sum_{k=0}^{n} a_k q_k(x)$, where $q_k(x)$ is a general orthogonal polynomial. Show that if $q_k(x)$ is a Chebyshev polynomial then the number of required multiplications can be reduced to half the number required in the general case.
- 8. Modify the class defined for least-squares to be an interpolation class. Encapsulate all the inner-workings of interpolation into one object.
- 9. Represent the data in the table below by a function of the form

$$g(x) = \alpha x^{-1} + \beta x^2,$$

using a procedure similar to least-squares. Determine the values of α, β that minimize the least-squares error.

Note: The resulting system of nonlinear equations may be solved iteratively; for help visit www.netlib.org.

10. Multi-variate polynomial interpolation

In most problems in simulation science the dependent variable is a function of more than one independent variable, e.g., $y = f(x_1, x_2)$ is a bi-variate function. We want to develop a least-squares multi-variate approximation using the quadratic polynomial

$$y = c_0 + \alpha_1 x + \beta_1 y + \alpha_2 x^2 + \beta_2 y^2 + c_1 x y.$$

Obtain the equation for c_i , α_i , β_i that minimize the least-squares error and formulate the problem in matrix form.

11. Determine a B-spline that goes through the end-points for the following control points:

- 12. Construct a fourth-order polynomial that approximates the function $f(x) = \sqrt{x}$ and goes through the points 0, 1, 2, 3, 4. Also, approximate this function with a cubic B-spline and compare the average and maximum approximation errors of the two approaches.
- 13. As stated in section 3.1.9 "Exercise Your Programming Skills," modify the given code to handle more than just bi-linear interpolation. Given nine points, you should be able to go up to bi-cubic interpolation. Follow the guidelines provided in the text.
- 14. Using the MPI_Send and MPI_Recv functions, write an MPI program which accomplishes piecewise Lagrange interpolation. As in the problem above, consider the function $f(x) = \sin 2\pi x$ where $x \in [-1,1]$. Write a program which accomplishes the following:
 - (a) Partition the interval [-1,1] based upon the number of processors used.
 - (b) For each process, use only four interpolation points (third-order polynomial) in which two of the points are the end-points of the subdomain used for that process.
 - (c) For each process, evaluate your interpolant at 30 points on its subdomain.
 - (d) For each process, send all 30 points to process 0 for collection.
 - (e) Have only process 0 print out the resulting interpolant evaluated at 30*P points (where P is the number of processes used).
- 15. Mappings in polynomial approximations: As we have seen in section 3.1.4 regions of rapid variation in a function may lead to global spurius oscillations. An effective remedy is to introduce a new coordinate system $x = q(s, \alpha)$, where x is the physical coordinate and the transformed coordinate satisfies: $-1 \le s \le 1$. The parameter α is associated with the steepness and its location.

Two such mappings have been introduced by [6] that work effectively with Chebyshev approximations.

The first mapping $x \mapsto s$ employs:

$$x = \alpha_2 + \tan[(s - s_0)\lambda]/\alpha_1$$

where

$$s_0 = \frac{k-1}{k+1}$$
, $k = \tan^{-1}[\alpha_1(1+\alpha_2)]/\tan[\alpha_1(1-\alpha_2)]$

and

$$\lambda = \tan^{-1}[\alpha_1(1-\alpha_2)]/(1-s_0).$$

Here α_1 expresses the degree of steepness and α_2 the location of rapid variation.

The second mapping has the form:

$$x = \sin^{-1}(\alpha_1 s) / \sin^{-1}(\alpha_1), \quad 0 \le \alpha_1 \le 1.$$

This mapping expands the boundary regions and compresses the interior regions. As $\alpha_1 \to 1$ the Gauss-Lobatto points become more uniformly spaced.

Consider the Runge function $f(x) = 1/(1 + 25x^2)$ with $x \in [-1, 1]$, and use both mappings to approximate f(x). Systematically vary the values of α_1 and α_2 to find optimum values for N = 4, 8, 12, 24 and 32 grid points. Make a *log-lin* plot of the L_2 and L_{∞} errors versus N and compare your answers to the unmapped case, as in figures 3.3 and 3.4.

- 16. Construct shape functions for a square domain $[-1,1] \times [-1,1]$ and a cubic domain $[-1,1] \times [-1,1] \times [-1,1]$ for which there are only three nodes per edge (i.e., two vertices and a middle node) but not an interior node. Repeat for the case where there are four equidistant nodes per edge but no interior nodes.
- 17. Construct shape functions for a triangular domain and a tetrahedron domain each with edges of unit length and with a middle node per edge as well as one interior node.
- 18. Modify the code for multi-dimensional interpolation presented in section 3.1.9 to handle more than just bi-linear interpolation. Given nine points, you should be able to go up to bi-cubic interpolation. Modify the code presented earlier to handle at least four points, and up to nine points. This implies that your function has to be intelligent concerning the number of points that it is passed. Let us first examine the things that will and will not change:
 - (a) Observe that the argument list itself need not change. Only the information passed into the function will change.
 - b) The user will now specify npts to be some value other than (or possibly including) four. The first thing to change: the static memory allocation of the array. No longer is an array of four elements sufficient. Should we move to dynamic memory allocation based on npts? No. Even though it would be perfectly legitimate to use dynamic memory allocation in this case, it is just as easy to allocate a static array for the maximum number of elements, which in this case is nine. In this case, it is "cheaper" to allocate a few extra doubles (in the case that npts is less than nine) than to both allocate and deallocate memory dynamically.
 - (a) You must now, of course, change the *argument checking*. No longer should you check to see if *npts* is exactly equal to four, but rather if *npts* lies between four and nine (inclusive).
 - (c) Now you must introduce the interpolation formulas for the various interpolants. We have already provided the first four formulas.
 - (d) The final issue is how to check which npts category into which you fall. Did the user want npts = 4, npts = 5, ... or npts = 9? Since there are a small number of possibilities, try using a switch statement. Recall, the layout will look like:

```
switch(npts)
  case 4:
    // formulas for npts = 4
  break;
```

```
case 5:
     // formulas for npts = 5
   break;
   case 6:
     // formulas for npts = 6
   break;
   case 7:
     // formulas for npts = 7
   break;
   case 8:
     // formulas for npts = 8
   break;
   case 9:
     // formulas for npts = 9
   break;
   default:
     //add error statement here for when an
     //invalid npts is given
}
```

One thing that we would like to point out in the example above is that we have made a programming decision to sacrifice optimality for generality. Experienced programmers would be quick to point out that the structure above is not optimal, and that if you were going to be doing thousands or millions of interpolations, that it would be better for you to write a function for each case separately (hence eliminating the switch statement, and even the argument checking). The tradeoff in this case is that we have only one function to maintain. This balance between generality and optimality is one which you will constantly be confronted with as a programmer. The balance you reach is almost always application and situation dependent.

3.5.2 Homework Problems for Section 3.2

- 1. Let $f = \sum a_k e^{ikx}$ and $g = \sum b_k e^{ikx}$, and we construct the product $fg = \sum c_k e^{ikx}$. What is the relationship between c_k and b_k, c_k ?
- 2. Construct the cosine and sine series of the function $f(x) = x^3$. Do the two series converge at the same rate? Plot the partial sums for each representation retaining N = 2, 4, 8, 16, 32 terms. What do you observe?
- 3. Compute the square of the Fourier matrix \mathbf{W} of order N=1024 using matrix-matrix multiplication, e.g. dgemm from BLAS3, and also using the discrete fast Fourier trans-

forms using the fact that this is a product of two circulant matrices. What do you observe?

- 4. Consider the function f(x) = 1, $x \in [0,1]$ and f(x) = -1, $x \in [1,2]$. Obtain its Fourier coefficients and use the Lanczos filter to improve the approximation. What is the asymptotic rate of convergence before and after the filtering in the L_1 and L_2 norms? You can obtain that either analytically or numerically.
- 5. Use a double Fourier series representation to expand in real double Fourier series the function

$$f(x,y) = x + y$$
 $x, y \in [-\pi, \pi].$

6. (Chebyshev Transforms)

Derive an algorithm for a fast Chebyshev transform based on the fast Fourier transform. Write a C++ code for it and obtain timings on your computer for various sizes N. How does it compare to the matrix-vector multiplication, and specifically at what value of N we have the break-even point?

3.5.3 Homework Problems for Section 3.3

- 1. Use the Fourier transform approach to plot the Battle-Lemarié wavelet of order m=4.
- 2. Use the iteration approach to plot the Daubechies scaling functions and wavelets of order m = 5 and 9.
- 3. Obtain the discrete wavelet transform of the function f using the Daubechies wavelets of fourth-order.
- 4. Derive equation 3.35 for the Shannon wavelet.

Chapter 4

Roots and Integrals

In this chapter we apply the approximation theory we presented in chapter 3 to find solutions of linear and nonlinear equations and to perform integration of general functions. Both subjects are classical, but they serve as basic tools in scientific computing operations and in solving systems of ordinary and partial differential equations. With regards to root finding, we consider both scalar as well as systems of nonlinear equations. We present different versions of the Newton-Raphson method, the steepest descent method, and the conjugate gradient method; we will revisit the latter in chapter 9. With regards to numerical integration we present some basic quadrature approaches, but we also consider advanced quadrature rules with singular integrands or in unbounded domains.

On the programming side, we first introduce the concept of passing a function to a function; in the previous chapter we were passing variables. This allows an easy implementation of recursion, which is so often encountered in scientific computing. We offer several C++ examples from root finding and numerical integration applications that make use of recursion, and show an effective use of classes and overloaded operators. We also address parallel programming with emphasis on domain decomposition, specifically the concept of reduction operations. We introduce the MPI commands MPI_Reduce and $MPI_Allreduce$ for accomplishing reduction operations among a collection of processes.

4.1 Root Finding Methods

There are many problems in scientific computing where we need to find the root of a nonlinear equation or systems of algebraic equations. For example, a polynomial equation arises in computing the eigenvalues from the characteristic polynomial, or general transcendental equations need to be solved to obtain the dispersion relation in wave dynamics problems. Another example is the computation of a square-root of a number; e.g., the computation of $\sqrt{3}$ can be turned into finding the root of the equation $f(x) = x^2 - 3 = 0$, and this can be solved *iteratively* very fast, starting from an educated guess! For example, for an initial guess of $x_0 = 1.5$ we substitute in the formula

$$x_{n+1} = \frac{1}{2}(x_n + \frac{3}{x_n}), \quad n = 1, 2, \dots$$

and obtain $x_1 = 1.75$ in one iteration versus the exact value 1.7320508. More iterations will result in predicting accurately more and more digits, in fact, we double the number of correct digits in each iteration! The above formula comes from the Newton-Raphson algorithm, and it is often used in mathematical libraries of computers for the square-root function.



Below we provide our own implementation of this function and construct table 4.1 showing the iteration count and associated error when computing $\sqrt{3}$ using this function.

```
double SquareRoot(double value, double guess, int iterations){
  int i;
  double xn = guess;

  for(i=0; i<iterations;i++)
      xn = 0.5*(xn + value/xn);

  return xn;
}</pre>
```

In the function above, observe that upon calculation of the next value x_{n+1} , we immediately place it in the variable xn. This is because in this iterative scheme, once the new value is computed, the previous value is not needed. An alternative version of the function is the following:

```
double SquareRoot(double value, double guess, double tol){
  int i;
  int maxit = 100;
  double xn2, xn = guess;

for(i=0; i<maxit;i++){
    xn2 = 0.5*(xn + value/xn);</pre>
```

```
if(fabs(xn2-xn)<tol)
    return xn2;
    xn = xn2;
}
cerr << "Maximum number of iterations reached ";
cerr << "without convergence" << endl;
return xn2;
}</pre>
```

In the function above, instead of inputing the number of iterations, we input a tolerance. As we calculate updated values, we check to see if the absolute value of the difference between successive iterates is less than the tolerance. For the code above, we have the following remarks:

- We do need to separately store the new value (xn2) so that it can be compared with the old value.
- We have chosen to set a maximum number of iterations instead of using a while loop.
- We use the function **fabs**, which is a function whose declaration is contained within *math.h*, and which takes as input a double precision number and provides back the double precision absolute value of that number. **NOTE:** A function **abs** exists also, but takes as input integers and returns the integer absolute value. Because of implicit casting, oftentimes people make the mistake of using **abs** when they should be using **fabs**, and they obtain the wrong result.
- We return directly from the *for* loop once the tolerance condition has been met. By using return, we terminate the loop tacitly by exiting the function.

Iteration	Root Approximation	Error
0	1.50000000	2.32050808e-01
1	1.75000000	1.79491924e-02
2	1.73214286	9.20495740e- 05
3	1.73205081	2.44585041e-09
:		

Table 4.1: Numerical approximation of $\sqrt{3}$.

We are familiar with root finding methods for the first- and second-order polynomial equations for which we can derive closed form solutions, and these solutions have been known for centuries to Egyptians and also to Babylonians for more than forty centuries! Clearly, we do not need to use a computer to obtain solutions for these equations, but things become much harder as the order of the polynomial increases, and in fact fifth- or higher order polynomials *cannot* be solved in closed form as Lagrange first discovered in the late

eighteen century. For third- and fourth-order polynomial equations, analytical formulas are available. They were first obtained by the Italian mathematician Ferrari, however they are very complex and thus not very useful.

In this section, we will study methods and corresponding code fragments to obtain solutions of *general nonlinear equations* as well as *systems* of general nonlinear equations, but it instructive to start with polynomial equations. The methods that we will develop are *iterative* and are easy to program but they require good knowledge of the basic theory. We need to know:

- How fast the algorithms converge,
- When to terminate the iterative process, and
- An initial guess to start the iteration, sometimes an educated guess!

In the following, we show how we can use the approximation theory of the previous chapter to achieve this.

4.1.1 Polynomial Equations

Cubic Polynomials: Let us first consider a cubic polynomial equation of the form

$$f(\xi) = \xi^3 + a\xi^2 + b\xi - c = 0$$

where a, b, c are real numbers so that we have at least one real root. Here we will follow the analysis of Lanczos [66]. The first step is to rescale the above equation in a more convenient form by introducing the transformation

$$x = \alpha \xi, \ a_1 = \alpha a, \ b_1 = \alpha^2 b, \ c_1 = \alpha^3 c$$

and by substitution we obtain

$$f(x) = x^3 + a_1 x^2 + b_1 x - c_1 = 0.$$

By taking $\alpha = 1/\sqrt[3]{c}$ we obtain $c_1 = 1$, and therefore

$$f(0) = -1 < 0$$
 and $f(\infty) > 0$.

We also have that the three roots satisfy

$$x_1 x_2 x_3 = 1.$$

Next, we examine the sign of f(x) at x = 1; if f(1) > 0 there exists one *real* root in the interval [0, 1], but if f(1) < 0 there must be one real root in the interval $[1, \infty]$. In this case, we simply introduce another transformation, i.e.,

$$\bar{x} = \frac{1}{x},$$

mapping $[1, \infty] \to [0, 1]$. It is then sufficient to find the real root of a cubic equation in the interval [0, 1].

Telescoping of a power series by successive reductions is an elegant way of reducing a high-order polynomial to a lower order by taking advantage of the properties of the Chebyshev polynomials discussed in section 3.1.5. In particular, we will use the *shifted* Chebyshev polynomials defined by

$$T_k^*(x) = T_k(2x - 1), \quad x \in [0, 1]$$
 (4.1)

which we tabulate in table 4.2. We will make use of the fact that:

- The shifted Chebyshev polynomials have the largest coefficient of the highest power of x among all polynomials defined in the same interval.
- They are bounded by ± 1 , just like the standard Chebyshev polynomials.
- They are defined in the interval of interest; here $x \in [0, 1]$.

Let us consider, for example, the third-order shifted Chebyshev polynomial $T_3^*(x) = 32 x^3 - 48 x^2 + 18 x - 1$, then

$$x^{3} = \frac{48x^{2} - 18x + 1}{32} + \frac{T_{3}^{*}(x)}{32},$$

and because of the boundness of $T_3^*(x)$, we can approximate

$$x^3 \approx 1.5 \ x^2 - 0.5625 \ x + 0.03125 \ \ (\pm 0.03125),$$

where the term in the parenthesis is the error $\pm 1/32$, which in this case is about 3%.

n = 1: 1, 2

n = 2: 1, 8, 8

n = 3: 1,18, 48, 32

n = 4: 1, <u>32</u>, 160, <u>256</u>, 128

n = 5: 1, 50, 400, 1120, 1280, 512

n = 6: 1, <u>72</u>, 840, <u>3584</u>, 6912, <u>6144</u>, 2048

Table 4.2: Coefficients of the first six shifted Chebyshev Polynomials $T_n^*(x)$. The underlined numbers correspond to negative coefficients, and the sequence is from the lowest to highest power; e.g., $T_3^*(x) = -1 + 18x - 48x^2 + 32x^3$; $T_0^*(x) = 1$.

Example: The following example illustrates the main points discussed so far. Let us consider the cubic polynomial equation

$$\xi^3 + \xi^2 - 1.5\xi - 50 = 0.$$

We first perform the transformation

$$\alpha = 1/\sqrt[3]{50} \approx 0.2714 \Rightarrow f(x) = x^3 + 0.2714x^2 - 0.1105x - 1 = 0,$$

and subsequently, we examine the sign of f(x) at x = 1:

$$f(1) = 0.1609 > 0 \Rightarrow x_1 \in [0, 1].$$

The next step is to reduce the cubic equation to a quadratic equation and apply the formula of Babylonians! To this end, we use the shifted Chebyshev polynomial $T_3^*(x)$ and equation (4.1) to obtain

$$1.7714 \ x^2 - 0.673 \ x - 0.9687 = 0.$$

which has a positive root $x_1 = 0.9534$ and a negative root, which we disregard. We now transform back to obtain

$$\xi_1 = \frac{x_1}{\alpha} = \frac{0.9534}{0.2714} = 3.513$$

The residual is f(3.513) = 0.42, and although this value is not 0, it is relatively small compared to the constant c = 50; it corresponds to an 0.4% error! The more accurate value obtained with the Newton-Raphson method is 3.5030. If, however, this is not an acceptable accuracy, then this "educated" guess can serve as initial value in one of the algorithms that we will present in this section.

Fourth-Order Polynomials: We can proceed similarly with fourth-order polynomial equations of the form

$$x^4 + c_1 x^3 + c_2 x^2 + c_3 x + c_4 = 0,$$

which can be turned into the form

$$(x^2 + \alpha x + \beta)^2 = (ax + b)^2,$$

and by taking the square root we reduce this to two quadratic equations. The question is how to obtain α, β, a and b in terms of c_i , i = 1, 2, 3, 4. This is accomplished via a series of simple transformations:

$$\alpha = c_1/2$$
; $A = c_2 - \alpha^2$; $B = c_3 - \alpha A$.

We then form the cubic equation

$$f(\xi) \equiv \xi^3 + (2A - \alpha^2)\xi^2 + (A^2 + 2B\alpha - 4c_4)\xi - B^2 = 0,$$

which has a positive real root since $f(0) = -B^2 < 0$. We now use the previous method for cubic polynomials to determine the real root, which we call ξ_1 . Having obtained ξ_1 , we determine all the coefficients of the two quadratic equations from

$$\alpha = \frac{1}{2}c_1, \ \beta = \frac{1}{2}(A + \xi_1), \ a = \sqrt{\xi_1}, \ b = \frac{a}{2}(\alpha - B/\xi_1).$$

Again, this real root can be used as initial guess to obtain more accurate answers from the algorithms presented later in this section.

High-Order Polynomials: Obtaining good approximations for the roots of high-order polynomials is a much more difficult job and may require many function evaluations in

order to locate approximately real roots. However, high-order polynomials have typically complex roots, so here we review a method first proposed by Bernoulli which provides an approximation to the *absolutely largest* root. Let us consider the polynomial

$$f(x) = x^{n} + a_{1}x^{n-1} + \ldots + a_{n} = (x - x_{1})(x - x_{2}) \ldots (x - x_{n}),$$

were x_i , i = 1, 2, ..., n denote the roots of the polynomial f(x). We can then compute the ratio

$$-\frac{f'(x)}{f(x)} = \frac{1}{x_1 - x} + \frac{1}{x_2 - x} + \ldots + \frac{1}{x_n - x}.$$

If x_0 is an initial guess which happens to be close to one of the roots, say x_1 , then by comparing terms in the above expansion we can see that one term dominates, i.e.,

$$\frac{1}{x_0 - x_1} \sim -\frac{f'(x_0)}{f(x_0)} \,.$$

Thus, we have managed to isolate one root, and, in fact, we can make this estimate sharper by taking the derivative (m-1) times to obtain

$$-\frac{1}{(m-1)!} \left[\frac{f'(x)}{f(x)} \right]^{(m-1)} = \frac{1}{(x_1 - x)^m} + \ldots + \frac{1}{(x_n - x)^m}$$
$$\Rightarrow \frac{1}{(x_1 - x_0)^m} \sim -\frac{1}{(m-1)!} \left[\frac{f'(x_0)}{f(x_0)} \right]^{(m-1)}.$$

By choosing m sufficiently high we can put the "spotlight" on the root x_1 with increasing accuracy. This method, however, requires a lot of work as it involves the computation of derivatives, although approximate ways have been suggested by Lanczos to do this efficiently; for example the so-called *method of moments*. The interested reader should consult Lanczos's book [66] for more details.

4.1.2 Fixed Point Iteration

One approach to solving nonlinear equations is by iteration, where the equation f(x) = 0 is re-arranged as

$$x = g(x), (4.2)$$

with f(x) = x - g(x). We can set up a fixed point iteration of the form

$$x_{n+1} = g(x_n),$$

which upon convergence ("steady state") leads to $x_{n+1} \to s$, and thus equation (4.2) is satisfied. It turns out that the key to convergence, as we will see in the theorem below, is the first derivative of g(x). We demonstrate this by a simple example of a quadratic polynomial, which is used often to model *chaos*. Let us consider

$$g(x) = \alpha x(1-x) = \alpha x - \alpha x^2,$$

with α being the bifurcation parameter, the meaning of which will become clear below.

We want to solve:

$$x_{n+1} = \alpha x_n - \alpha x_n^2. (4.3)$$

We note that the maximum value of g(x) is $g(1/2) = \alpha/4$, and therefore for $\alpha < 4$, $x_n \in [0, 1]$, i.e. the sequence of numbers produced by the iteration process remains bounded within the range [0, 1]. Let us consider a specific example, and take $\alpha = 2$. Then

$$x_{n+1} = 2x_n - 2x_n^2$$

and in steady state, that is upon convergence, we have that $x_n \to s$ and also $x_{n+1} \to s$, so the fixed or *stationary points* are given by the equation

$$s = 2s - 2s^2,$$

which has two roots, the stationary points, at s = 0 and $\frac{1}{2}$.

We then evaluate the first derivative at the stationary points, and if

- |g'(s)| < 1, the stationary point s is attractive,
- otherwise, it is *repulsive*.

This can be seen by constructing the graph of g(x) and follow the sequence generated by the iterative equation. In this case, we have that g'(0) = 2, which is a repulsive point, and g'(1/2) = 0, which is an attractive point, and in fact very attractive!

This process can be easily generalized for the simple iteration equation we have considered to obtain the stationary points $s_1 = 0$ and $s_2 = (\alpha - 1)/\alpha$. At the second point, we have

$$g'(s_2) = 2 - \alpha$$
 and $|2 - \alpha| < 1 \Rightarrow -1 < 2 - \alpha < 1 \Rightarrow 1 < \alpha < 3$.

Therefore, we have found the range within which the bifurcation parameter α results in an attractive point and thus a convergent iteration process. Similarly, it is easy to show that $g'(s_1) = \alpha$, so as long as $\alpha < 1$ we have a convergent iteration process.

We can of course substitute back in equation (4.2) to obtain

$$q_1(x) = q(q(x)) = \alpha(\alpha x - \alpha x^2) - \alpha(\alpha x - \alpha x^2)^2$$

which has four stationary points. By proceeding as above and examining the derivative $g'_1(s)$, we find that we have an instability and thus divergence for $\alpha > 3.45$. In fact, for higher values of α the system bifurcates even further to a *period doubling cascade*. The ratio between the length of one stability window and the next approaches a universal constant $\delta = 4.69920166...$, which is known as the *Feigenbaum's constant* in chaos theory.

Example: Next, we present an example of the iteration process for the quadratic chaos equation, which converges for some small value of the bifurcation parameter; for larger values oscillates between different values even at steady state; and for even larger values it is unstable and diverges. Consider the iteration

$$x_{n+1} = (1+10\alpha)x_n - 10\alpha x_n^2, \quad x_0 = 0.1.$$
 (4.4)

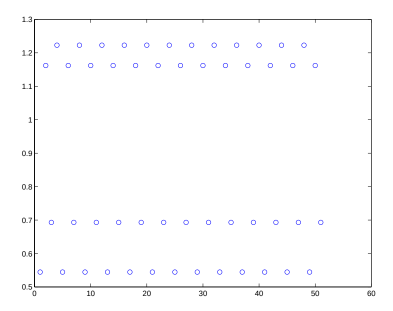


Figure 4.1: Plot of the values (vertical axis) of the time series versus the iteration number (horizontal axis) produced from the quadratic chaos equation (4.3); $\alpha = 0.249$. The iteration converges to four different values at steady state.

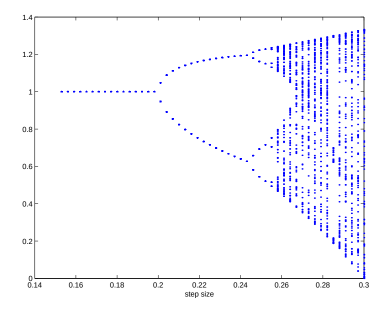


Figure 4.2: Bifurcation diagram. Plotted are the converged values of the iteration in equation (4.4) versus the bifurcation parameter (step size) α .

We find that for $\alpha=0.18$ the above iteration converges to 1, but for $\alpha=0.23$ it jumps between 1.18 and 0.69. For $\alpha=0.25$ it jumps between 1.23, 0.54, 1.16 and 0.70. For the value $\alpha=0.3$ no discernible pattern is displayed; note also a window of *order* just above $\alpha=0.28$ and then a return to chaos again! The plots in figures 4.1 and 4.2 show schematically this strange but interesting behavior.

After this introductory example, we are now ready to state the theorem on convergence of fixed point iteration.

Theorem of Convergence of Fixed Point Iteration: Let s be a solution of x = g(x) and g has a continuous derivative in some interval I containing s. Then if $|g'(s)| \le K < 1$ in I, the iteration process $x_{n+1} = g(x_n)$ converges for any initial value x_0 in I.

Proof: From the mean value theorem we have that there exists a point $t_1 \in [x, s]$: $g(x) - g(s) = g'(t_1)(x - s)$. Then

$$|x_{n} - s| = |g(x_{n-1}) - g(s)| = |g'(t_{1})| |x_{n-1} - s|$$

$$\leq K|x_{n-1} - s|$$

$$= K|g(x_{n-2}) - g(s)|$$

$$\leq K|g'(t_{2})| |x_{n-2} - s|$$

$$\leq K^{2}|x_{n-2} - s|$$

$$\vdots$$

$$\leq K^{n}|x_{0} - s| \to 0.$$

Here $t_i, i = 1, 2, ... \in I$ as well. Note that g(x) is called a *contraction*, and in general a contraction mapping is defined as

$$|g(u) - g(v)| \le K|u - v|, \quad K < 1.$$

Round-off: Associated with such an iteration process is round-off errors, which, however, can be quantified. To this end, consider the iteration

$$x_n = g(x_{n-1}) + \delta_n$$

where $|\delta_n| \leq \delta$ and $\delta \ll 1$ denotes the known machine accuracy. We then have

$$|x_{n} - s| = |g(x_{n-1}) - g(s) + \delta_{n}|$$

$$\leq K|x_{n-1} - s| + \delta$$

$$\leq K(K|x_{n-2} - s| + \delta) + \delta$$

$$\leq K^{n}|x_{0} - s| + \delta(1 + K + K^{2} + \dots + K^{n-1})$$

$$\leq K^{n}|x_{0} - s| + \frac{\delta}{1 - K},$$

where in the last line we used the limit of a geometric series since K < 1.

Error Estimate: Similarly, we can obtain an error estimate by comparing two successive approximations:

$$|x_{n+1} - x_n| = |g(x_n) - g(x_{n-1})|$$

 $\leq K|x_n - x_{n-1}|$
 \vdots
 $\leq K^n|x_1 - x_0|$.

For fixed n and m > n we have

$$|x_{m} - x_{n}| \leq |x_{m} - x_{m-1}| + |x_{m-1} - x_{m-2}| + \dots + |x_{n+1} - x_{n}|$$

$$\leq (K^{m-1} + K^{m-2} + \dots + K^{n})|x_{1} - x_{0}|$$

$$= \frac{K^{n}}{1 - K}|x_{1} - x_{0}|.$$

Since $x_m \to s$ as $m \to \infty$ for convergence, then

$$|x_n - s| \le \frac{K^n}{1 - K} |x_1 - x_0|.$$

We can now define the **convergence rate** as the ratio of two successive error estimates. Let $e_n \equiv x_n - s$ and s = g(s), then

$$e_{n+1} = x_{n+1} - s = g(s + e_n) - g(s) = g'(t)e_n$$

from the mean value theorem, $t \in [x_n, s]$, and thus in the limit:

$$\lim_{n \to \infty} \frac{e_{n+1}}{e_n} = g'(s).$$

If g'(s) = 0 and g'(x) continuous (as in the case of the very attractive point at s = 0 that we encountered in the example earlier), then

$$e_{n+1} = g(s + e_n) - g(s) = g'(s)e_n + g''(s)\frac{e_n^2}{2} + \dots$$

and therefore

$$\lim_{n \to \infty} \frac{e_{n+1}}{e_n^2} = \frac{1}{2}g''(s)$$

which shows quadratic convergence, that is at every iteration the number of correct digits is doubled!

4.1.3 Newton-Raphson Method

We now present the most popular method in rood finding, which was proposed independently by Newton and Raphson. It combines the ideas of iterative process discussed above and of approximation presented in section 3.1. Let us consider again the equation

$$f(x) \equiv x - g(x)$$
 and $f(s) = 0$,

and denote by h the distance between the exact solution s and an initial guess x_0 . Then, we use Taylor expansion around x_0 to obtain

$$f(s) = 0 = f(h + x_0) = f(x_0) + hf'(x_0) + \frac{h^2}{2}f''(x_0) + \dots$$

By neglecting terms higher than second-order in h, we obtain

$$h_0 \equiv h = -\frac{f(x_0)}{f'(x_0)}.$$

We can now propose an improved guess $x_1 = x_0 + h_0$, which leads to a hopefully shorter distance

$$h_1 = -\frac{f(x_1)}{f'(x_1)} \,,$$

and so on.

We summarize this iterative process as follows:

$$x_{n+1} = x_0 + h_0 + \dots + h_n$$

 $h_n = -\frac{f(x_n)}{f'(x_n)}, \quad n = 1, 2, \dots$

where x_0 is the initial guess that starts the iteration. Alternatively, we can write the above two equations as a single equation

$$x_{n+1} = \underbrace{x_n - \frac{f(x_n)}{f'(x_n)}}_{g(x_n)} \tag{4.5}$$

assuming that $f'(s) \neq 0$. The convergence rate can then be obtained by examining the first derivative g'(s):

$$g'(x) = 1 - \frac{f'^2 - ff''}{(f')^2} = f\frac{f''}{(f')^2} \to 0 \text{ as } x \to s.$$

We also have that

$$g''(x) = \frac{(ff'')'(f')^2 - 2f'f''ff''}{(f')^4} = \frac{ff'''f' + f'^2f'' - 2ff''^2}{(f')^3} \neq 0 \text{ as } x \to s.$$

In general, $g''(s) \neq 0$, and therefore the convergence rate of the Newton-Raphson is second-order (quadratic convergence).

Example: Let us revisit the example of finding the square-root of a positive number that we discussed in the introduction of this section. We assume that

$$f(x) = x^2 - C$$
 and therefore $s = \sqrt{C}$

is the exact answer. We now use the Newton-Raphson formula from above

$$f'(x_n)(x_{n+1} - x_n) = -f(x_n),$$

and thus

$$2x_n(x_{n+1} - x_n) = C - (x_n)^2 \Rightarrow x_{n+1} = \frac{1}{2} \left(x_n + \frac{C}{x_n} \right)$$

Therefore, we compute the square-root \sqrt{C} by averaging x and C/x. The convergence is quadratic, i.e.

$$x_{n+1} - \sqrt{C} = \frac{1}{2x_n}(x_n - \sqrt{C})^2.$$

At every iteration step the difference $(x_n - \sqrt{C})$, which is the error, is squared!

Improved Convergence

We can improve the convergence rate of the Newton-Raphson method by retaining the second-order term in the Taylor expansion, i.e.,

$$f(x_0) + h[f'(x_0) + \frac{h}{2}f''(x_0)] = 0$$

$$\Rightarrow h = -\frac{f(x_0)}{f'(x_0) + \frac{h}{2}f''(x_0)}.$$

In the above equation, we can compute a provisional value for h in the denominator, say h_n^* , from the linear Taylor expansion at the previous iteration as

$$h_n^* = -\frac{f(x_n)}{f'(x_n)} \Rightarrow \frac{1}{h_n} = -\frac{f'(x_n)}{f(x_n)} + \frac{1}{2} \frac{f''(x_n)}{f'(x_n)}$$
.

This two-step approach is a general procedure called *predictor-corrector* method and is used very often in scientific computing. The idea is to first predict with a lower order but explicit method and subsequently to correct the answer with a higher order method.

Multiple Roots

So far we have only treated the case of a single root, but we also need to have fast methods to compute multiple roots. In such cases, both the function and some of its derivatives vanish at the root location, depending on the multiplicity.

Let us first examine the case of a **double root** and consider an initial guess x_0 ; if it is close to the exact root s then $f(x_0) \to 0$ and also $f'(x_0) \to 0$. Therefore, the standard Newton-Raphson method will converge slowly due to the fact that the term

$$g'(x_0) = f(x_0) \frac{f''(x_0)}{[f'(x_0)]^2}$$

will be finite or slowly decaying to zero, and

$$\lim_{n\to\infty}\frac{e_{n+1}}{e_n}=g'(s)\neq 0\,,$$

which indicates only first-order convergence for a double root. We can improve convergence by solving the quadratic equation with respect to h

$$\frac{1}{2}f''(x_0)h^2 + f'(x_0)h + f(x_0) = 0,$$

derived from Taylor's expansion to obtain two roots h_{\pm} and then set

$$x_1 = x_0 \pm h_{\pm},$$

and proceed with the iteration.

In the case of a **triple root**

$$f(x_0) \to 0$$
, $f'(x_0) \to 0$, $f''(x_0) \to 0$,

and we need to solve a cubic equation to obtain the proper distance in order to update the guess, i.e.,

$$\frac{1}{6}f'''(x_0)h^3 + \frac{1}{2}f''(x_0)h^2 + f_0'(x_0)h + f(x_0) = 0.$$

By solving this equation we obtain three values h_1 , h_2 , h_3 , and thus we can update $x_1 = x_0 + h_i$, i = 1, 2, 3, and so on for the next iteration.

A better approach in dealing with multiple roots is presented in the following. We assume that the function f(x) has a **root of multiplicity** p, so we can write

$$f(x) = (x - s)^p h(x), \quad p > 1, \quad h(s) \neq 0,$$

and

$$g(x) \equiv x - \frac{f(x)}{f'(x)}$$

whose derivative as $x \to s$ determines the convergence rate. We obtain

$$g'(x) = 1 - \frac{ph^2(x) + (x-s)^2(h'(x))^2 - (x-s)^2h(x)h''(x)}{p^2h^2(x)\left[1 + (x-s)\frac{h'(x)}{ph(x)}\right]^2}.$$

Upon convergence we have that $x \to s$, and thus

$$g'(s) \to 1 - \frac{1}{p} \neq 0$$
 for $p > 1$.

Clearly, |g'(s)| < 1, and the method converges, but it is only first-order. However, we note that if we re-define g(s) so that

$$g(x) \equiv x - p \frac{f(x)}{f'(x)},$$

we obtain quadratic convergence, as $g'(x \to s) \to 0$, and we use the theorem of convergence of the previous section. The iterative process is then

$$x_{n+1} = x_n - p \frac{f(x_n)}{f'(x_n)}, \quad n = 0, 1, \dots$$

Yet, $another\ way$ to obtain second-order convergence for a multiple root is to define the function

$$\phi(x) = \frac{f(x)}{f'(x)},$$

which upon convergence $(x \to s)$ has the simple form

$$\phi(x \to s) \to \frac{x-s}{p}$$
,

with a *single* root at x = s, and thus Newton-Raphson will converge quadratically fast. However, this method requires three function evaluations, i.e. f(x), f'(x) and f''(x), and thus it is more expensive than the previous one.

• In general, it is impossible to achieve second-order convergence with only one function evaluation.

The initial guess is very important in starting the iterative process in the Newton-Raphson algorithm. The following theorem states the proper conditions for convergence:

Theorem of Convergence of Newton-Raphson Method: Suppose that $f'(x) \neq 0$, that f''(x) does not change sign in the interval [a, b], and that f(a)f(b) < 0.

If
$$\left| \frac{f(a)}{f'(a)} \right| < (b-a) \text{ and } \left| \frac{f(b)}{f'(b)} \right| < (b-a)$$

then the Newton-Raphson method converges from an arbitrary initial approximation $x_0 \in [a, b]$.

4.1.4 Passing Functions to Functions in C++

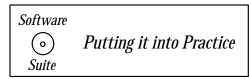
One thing that we will find useful throughout this chapter is the concept of passing functions to functions, just as you pass variables to functions. Why is this useful? When writing modular code, quite often you write essential library routines once, and compile them once, and then merely link them to your current program when you compile. Take for example our Newton-Raphson algorithm above. Notice that the algorithm is described independently of the particular function with which you are working; it merely relies on the fact that you have a function f(x) and its derivative f'(x). We want to write code in such a way that we can change our function and its derivative, and never have to change our core Newton-Raphson implementation. This can be accomplished by passing functions to functions.

Just as in the case of a variable, within the argument of the function to which you are passing the function, you must provide information as to what is being passed. In the case of passing a function, the information that is needed is:

- The name that the function is to be called within the routine.
- The argument types of the function that you are passing.

• The return value of the function that you are passing.

This procedure will be made more clear in the examples below.



In the following examples, there are two main concepts that we would like to draw your attention to:

- Passing functions to functions.
- Using different functions with the same name.

We begin with passing functions to functions. The example below is our implementation of the Newton-Raphson method. Recall that for this algorithm, we require both the function and its derivative. We thus require two functions, here called "func" and "func_der" within our code to be passed to the NewtonRaphson routine.

Concerning the function above, we have the following remarks:

• Observe how the function above is notified that a function is being passed to it. In the second argument of the argument list above, we specify as the argument

```
double (*func)(double)
```

notifying the function of the return type (in this case double), name of the passed function to be used within the function (in this case func), and the argument types of the passed function (in this case double). In the third argument we pass another function with the same return type and argument list types, but a different name $(func_der)$.

- We wrote the *if* statement on one line this is perfectly valid in C++. Since by definition an *if* statement executes the next executable statement given that the check is evaluated to be true, then we can condense an *if* statement having only one executable statement to one line. We suggest that if the executable statement which follows the *if* is complicated, you do not condense both the *if* and the next executable statement to one line.
- We can sacrifice memory for optimization. Notice that within the *if* statement, we evaluate func(x). If the tolerance criterion is not met, we set x0 = x and begin the iteration again, which requires that we evaluate func(x0). If the function func is expensive to compute, it is advantageous us to save the function evaluation to a variable which we use both in the *if* statement and in the case that the tolerance is not met.

Look at the function below: What is the difference between this function and the one directly above?

The answer: The second function does not require you to pass the variable "multiplicity." The second implementation of the *NewtonRaphson* function assumes the multiplicity is one. Why do we point this out? Notice that we used the same function name for both functions. C++ allows you to have multiple functions with the same name *as long as* the argument lists are different. By different, we mean either in number of arguments or in the type of variable.

Key Concept

• C++ allows multiple functions with the same name as long as the argument lists are different.

In the example above, the argument list of one function has less variables than the argument list of the other function, and hence the compiler can distinguish them as two different functions. This concept is useful in writing general software libraries, in that you can handle many different function cases by having functions with the same name but different argument lists.

WARNING

Programmer Beware!

• Different return types does not imply different functions!

The compiler cannot tell the difference between two functions which have identical names and arguments, but only different return types!

Also, observe that in the function above we use the keyword **const**. This identifier should be used in conjunction with a variable declaration and initialization as follows:

```
const <type> <variable name> = <constant value>;
```

where <type> is the variable type, <variable name> is the variable name we have chosen to use, and <constant value> is a value of the appropriate type which we would like to remain constant. Once a variable has been declared constant within a function, it cannot be changed within the function. Using const reinforces that something is meant to be a constant, and guarantees that the programmer does not accidentally reset the value to something else (which would imply that the value was not constant!).

We provide another example of the *NewtonRaphson* routine, now with three functions being passed to this routine. In this case, we pass a functional implementation of the second derivative of the function to this routine so that the convergence rate for multiple roots can be computed.

```
cout << "------" << endl;
cout << "x \t f(x) \t g'(x)" << endl;

for(int i=0;i<max_iter;i++){
   cout << x0 << "\t" << func(x0) << "\t" << (1-p) +
        p*func(x0)*func_secondder(x0)/pow(func_der(x0),2) << endl;;
   x = x0 - p*func(x0)/func_der(x0);
   if(fabs(func(x))<tolerance) break;
   x0 = x;
}

return x;
}</pre>
```

We can now put into practice what we have learned above. The following program uses the last NewtonRaphson function to compute the roots of

$$(x-2)^3(x+2) = 0.$$

```
#include <iostream.h>
#include "SCchapter4.h"
const double alpha = 2.0; // global constant
double func(double x);
double func_der(double x);
double func_secondder(double x);
int main(int argc, char * argv[]){
  int max_iter = 100;
  int multiplicity = 1;
  double x0 = 1;
  double x = NewtonRaphson(x0,func,func_der,func_secondder,
    max_iter,multiplicity);
  cout << x << " " << fabs(func(x)) <<endl;;</pre>
}
double func(double x){
  double value = (x-alpha)*(x-alpha)*(x-alpha)*(x+alpha);
  return(value);
```

```
double func_der(double x){
  double value = 3*(x-alpha)*(x-alpha)*(x+alpha)+
      (x-alpha)*(x-alpha)*(x-alpha);

  return(value);
}

double func_secondder(double x){
  double value = 6*(x-alpha)*(x+alpha)+6*(x-alpha)*(x-alpha);
  return(value);
}
```

Observe that we declare and define our implementations of the function, derivative function and second derivative function just as we have always done previously. Now, when it is time to call the *NewtonRaphson* routine we place into the argument list the function names in the appropriate places within the argument list, just as if we were passing these functions as variables.

4.1.5 Secant Method

The Newton-Raphson method requires that we know the function analytically, so that we can differentiate it analytically. However, there are instances where we only know the function values and therefore we need to construct the derivative using finite differences. General finite difference methods will be covered in detail in sections 5.1 and 6.1, but here we use the simple backward-difference formula to approximate the first derivative, i.e.,

$$f'(x_n) \approx (\Delta f)_1 \equiv \frac{f(x_n) - f(x_{n-1})}{x_n - x_{n-1}}$$
.

The secant method extrapolates along the chord that connects the points x_n and x_{n-1} instead of extrapolating along the tangent at x_{n-1} as the Newton-Raphson method. This is a first-order, $\mathcal{O}(h)$, approximation to the derivative, however the overall convergence rate of the secant method is about 1.6 instead of 2 of the Newton-Raphson method.

In comparing the secant method with the Newton-Raphson method in situations where both approaches can be applied, we have a *break-even point* determined approximately by the equation of cost in computing

$$f'(x) = \theta \times (\text{cost of } f(x)).$$

Then if $\theta > 0.4$, it is more efficient to use the secant method. Note that (0.4 = 2 - 1.6) reflects the difference in convergence rate of the two methods.

Error Analysis of the Secant Method

In the following, we provide a brief proof on the convergence rate of the secant method, following the derivation in [72].

We use Newton's interpolating formula presented in chapter 3 and the mean value theorem:

$$f(x) = f(x_n) + (x - x_n)(\Delta f)_1 + (x - x_{n-1})(x - x_n)\frac{1}{2}f''(\xi),$$

where we defined earlier

$$(\Delta f)_1 = \frac{f_n - f_{n-1}}{x_n - x_{n-1}}.$$

Now x_{n+1} satisfies the secant method equation: $0 = f_n + (x_{n+1} - x_n)(\Delta f)_1$.

We also have for the exact solution:

$$0 = f_n + (s - x_n)(\Delta f)_1 + \frac{1}{2}(s - x_{n-1})(s - x_n)f''(\xi),$$

and by subtracting, we obtain

$$(s - x_{n+1})(\Delta f)_1 + \frac{1}{2}(s - x_{n-1})(s - x_n)f''(\xi) = 0.$$

From the mean value theorem we have that for a special point ξ^* , $(\Delta f)_1 = f'(\xi^*)$, and therefore the error

$$e_{n+1} = \underbrace{\frac{f''(\xi)}{2f'(\xi^*)}}_{C} e_n e_{n-1}.$$

We have assumed in the above that f''(x) is continuous, and thus it is bounded, so we can always find the constant C. Let us now assume that

$$|\epsilon_{n+1}| = K|\epsilon_n|^m,$$

and substitute in the above to arrive at

$$K|e_n|^m \approx C|e_n| (K^{-1/m}|e_n|^{1/m}) \Rightarrow m = 1 + 1/m \to m = \frac{1}{2}(1 \pm \sqrt{5}).$$

We can only accept the (+) sign and thus we obtain that the convergence rate is $m \approx 1.618$ (recall, this value is the *golden ratio*, which we computed from the Fibonacci sequence in the exercises in chapter 2).

4.1.6 Systems of Nonlinear Equations

The Newton-Raphson method can be generalized in a straightforward manner to systems of nonlinear equations by introducing the Jacobian operator. Let us consider the $n \times n$ system:

$$f_1(x_1, \dots, x_n) = 0$$

$$f_2(x_1, \dots, x_n) = 0$$

$$\vdots$$

$$f_n(x_1, \dots, x_n) = 0$$

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which we re-write in compact form as

$$f(x) = 0.$$

Then, the iterative Newton-Raphson method for the above system is:

$$\mathbf{J}_n(\mathbf{x}_{n+1} - \mathbf{x}_n) = -\mathbf{f}(\mathbf{x}_n),$$

where we define the Jacobian operator as

$$\mathbf{J} = \frac{\partial f_i}{\partial x_i} \,.$$

We also note that here the subscripts $n, (n+1), \ldots$ denote iteration number but also components of the vector \mathbf{x} .

Convergence: Next, we examine the conditions that guarantee that the Newton-Raphson method for a system converges at a rate similar to the scalar equations, i.e., at quadratic rate. This is done by examining the partial derivatives of

$$\mathbf{g}(\mathbf{x}) \equiv \mathbf{x} - \mathbf{J}^{-1} \mathbf{f}(\mathbf{x})$$

and by differentiation

$$\frac{\partial \mathbf{g}(\mathbf{x})}{\partial x_j} = \frac{\partial \mathbf{x}}{\partial x_j} - \frac{\partial}{\partial x_j} \left[\mathbf{J}^{-1}(\mathbf{x}) \mathbf{f}(\mathbf{x}) \right]
= \frac{\partial \mathbf{x}}{\partial x_j} - \mathbf{J}^{-1}(\mathbf{x}) \frac{\partial \mathbf{f}}{\partial x_j} - \mathbf{f}(\mathbf{x}) \frac{\partial \mathbf{J}^{-1}(\mathbf{x})}{\partial x_j}.$$

At the solution $\mathbf{x} = \mathbf{s}$, the above expression becomes

$$\frac{\partial \mathbf{g}(\mathbf{s})}{\partial x_j} = \mathbf{I} - \mathbf{J}^{-1}(\mathbf{s})\mathbf{J}(\mathbf{s}) - \mathbf{f}(\mathbf{s})\frac{\partial \mathbf{J}^{-1}(\mathbf{s})}{\partial x_j} \to 0,$$

where **I** is the identity operator. Furthermore, in order to determine the last term $\frac{\partial}{\partial x_j}[\mathbf{J}^{-1}(\mathbf{x})]$, we note that

$$\frac{\partial}{\partial x_j} [\mathbf{J}^{-1} \mathbf{J}] = \frac{\partial \mathbf{I}}{\partial \mathbf{x}} = \mathbf{0} = \mathbf{J}^{-1} \frac{\partial \mathbf{J}}{\partial \mathbf{x}} + \mathbf{J} \frac{\partial \mathbf{J}^{-1}}{\partial \mathbf{x}},$$

and therefore

$$\frac{\partial \mathbf{J}^{-1}}{\partial x_j} = -\mathbf{J}^{-1} \frac{\partial \mathbf{J}}{\partial x_j} \mathbf{J}^{-1}.$$

From the analysis above, we see that we need only require that:

- f(x) have two derivatives, and
- J(x) be non-singular at the solution or root.

Then, the convergence of Newton-Raphson's method is *quadratic*, similar to the scalar case. The theorem below gives the precise conditions for convergence.

Convergence Theorem: Let us assume that

$$\parallel \mathbf{J}^{-1}(\mathbf{x}_0) \parallel \leq a \quad and \quad \parallel \mathbf{x}_0 - \mathbf{x}_1 \parallel \leq b$$

and also that

$$\sum_{k=1}^{n} \left| \frac{\partial^2 f_i(\mathbf{x})}{\partial x_j \partial x_k} \right| \le \frac{c}{n}, \quad \forall \ \mathbf{x} : \| \ \mathbf{x} - \mathbf{x}_0 \| \le 2b, \quad i, j = 1, 2, \dots, n$$

If in addition $abc \le 1/2$, then the Newton-Raphson iterates are uniquely defined and lie in the 2b-sphere, i.e.,

$$\| \mathbf{x}_k - \mathbf{x}_0 \| \le 2b$$

$$\lim_{k \to \infty} \mathbf{x}_k = \mathbf{s}, \quad \| \mathbf{x}_k - \mathbf{s} \| \le \frac{2b}{2^k}.$$

The problem with the Newton-Raphson method for systems is that it is computationally expensive as it requires the solution of a linear system in each iteration to invert the Jacobian matrix \mathbf{J} . It also requires the construction of the n^2 entries in the Jacobian matrix, which are partial derivatives of $\mathbf{f}(\mathbf{x})$. To this end, several modifications have been proposed and many different versions are used in practice, depending on the particular application. Next, we present two simple popular versions.

Modified Newton: Here the Jacobian is computed only initially and used for all subsequent iterations, i.e.,

$$\mathbf{J}_0(\mathbf{x}_{n+1}-\mathbf{x}_n)=-\mathbf{f}(\mathbf{x}_n).$$

This, however, lowers the convergence to first-order, as can be seen for the scalar example presented earlier for the square-root (here $\mathbf{J}_0 = 2x_0$.):

$$(x_{n+1} - \sqrt{C}) = \left[1 - \frac{x_n + \sqrt{C}}{2x_0}\right](x_n - \sqrt{C}),$$

which clearly indicates linear convergence.

Quasi-Newton: Here we use the values from the *first two iterations* $\mathbf{x}_1, \mathbf{x}_0$ and we define $\Delta \mathbf{x} \equiv \mathbf{x}_1 - \mathbf{x}_0$ and $\Delta \mathbf{f} \equiv \mathbf{f}_1 - \mathbf{f}_0$. The idea is to adjust \mathbf{J}_1 to satisfy $\mathbf{J}_1 \Delta \mathbf{x} = \Delta \mathbf{f}$, and thus

$$\mathbf{J}_1 = \mathbf{J}_0 + \frac{(\Delta \mathbf{f} - \mathbf{J}_0 \Delta \mathbf{x})(\Delta \mathbf{x})^T}{(\Delta \mathbf{x})^T (\Delta \mathbf{x})},$$

and then $\mathbf{x}_2 = \mathbf{x}_1 - \mathbf{J}^{-1}\mathbf{f}$. Convergence is faster than linear but not quite quadratic. An extension of this approach with modifications to preserve symmetry of \mathbf{J} is used in finite element methods [84].

Continuation Method

We have seen so far through the theorems and through examples how crucial the initial condition (guess) is for the Newton-Raphson method to converge, and this is especially true for systems of nonlinear equations. In problems where we have to solve nonlinear equations many times, e.g. in time-dependent simulations, we can use the solution of the previous time step. Similarly, assuming that we study the instability of a system which depends on a bifurcation parameter α , we need to solve

$$\mathbf{f}(\mathbf{x}, \alpha) = \mathbf{0}$$

for several values of α . One approach that is often used in practice is to linearize the equations for the critical value of the bifurcation parameter, say at α_c , e.g. right at the onset of the bifurcation (instability), and compute the solution to the linear system of equations. We can then use that solution as a initial guess to obtain a solution for a value of $\alpha > \alpha_c$ but with the difference $(\alpha - \alpha_c)$ not too large. This process can continue for a larger value of α , and so on. This is the so-called *continuation method*, as we assume that the systems of equations define dependent variables which are continuously differentiable in α .

If there is no bifurcation parameter explicitly included in the equations, we can still use the continuation method by introducing a *fictitious* parameter α and rewrite the system of equations as

$$\mathbf{h}(\mathbf{x}, \alpha) = (1 - \alpha)\mathbf{f}(\mathbf{x}_0) - \mathbf{f}(\mathbf{x}), \quad \alpha \in [0, 1]$$

where \mathbf{x}_0 is a first solution guess which may or may not satisfy the original system $\mathbf{f}(\mathbf{x}) = \mathbf{0}$. We see that for $\alpha = 0$ the guess solution is a root for $\mathbf{h}(\mathbf{x})$, and for $\alpha = 1$ we recover the system we want to solve. This suggests that we can start with the guess solution \mathbf{x}_0 and $\alpha = 0$ and continue in small increments of α up to $\alpha = 1$.

4.1.7 Solution via Minimization: Steepest Descent and Conjugate Gradients

In this section we present ways of solving systems of linear and nonlinear algebraic equations by attempting to obtain the minimum of an appropriately defined functional. We will focus on two useful methods: Steepest Descent and Conjugate Gradient.

Method of Steepest Descent

We have already mentioned the computational complexity associated with the computation of the Jacobian J, but even worse, sometimes we may not be able to compute it at all. In this case, we can still use an approach similar to the secant method in which we replace the inverse Jacobian with a scalar constant α , as follows

$$\mathbf{x}_{n+1} - \mathbf{x}_n = -\alpha \mathbf{f}(\mathbf{x}_n). \tag{4.6}$$

The step size α is important, but it is the direction of the path that we follow, namely along the tangent direction $-\mathbf{f}(\mathbf{x}_n)$ that gives this method the name of the *steepest descent*.

In order to appreciate this, let us introduce the anti-derivative of the system of equations we try to solve, which is a scalar function

$$P(x_1, \dots, x_n)$$
 and $f_i = \frac{\partial P}{\partial x_i} = 0$.

We can then interpret $\mathbf{f}(\mathbf{x})$ as the gradient of the parent function $P(\mathbf{x})$. Obtaining the solution is equivalent to minimizing the parent function, as we set all its partial derivatives equal to zero. For example, for a system of *linear equations* $\mathbf{A}\mathbf{x} = \mathbf{b}$, the parent function that we minimize has the simple quadratic form

$$P(\mathbf{x}) = \frac{1}{2}\mathbf{x}^T \mathbf{A} \mathbf{x} - \mathbf{x}^T \mathbf{b}$$

as the location of the minimum coincides with the solution $\mathbf{x} = \mathbf{A}^{-1}\mathbf{b}$. For nonlinear equations, however, the parent function is more difficult to obtain.

Since $\mathbf{f} = \nabla P$ and the gradient operator points towards increasing values of the scalar P, that is along paths of steepest ascent, it is clear that $-\mathbf{f}(\mathbf{x}_n)$ points along paths of steepest descent (see equation (4.6)), and it is perpendicular to iso-contours of $P(\mathbf{x})$. The iso-surfaces of a positive-definite quadratic $P(\mathbf{x})$ is an ellipsoid centered about the global minimum; the semi-axes of the ellipsoid are related to the eigenvalues of the corresponding matrix \mathbf{A} . If the eigenvalues of \mathbf{A} are all equal, then the iso-surfaces of P would be spheres, and thus the steepest descent direction would point towards the sphere center. However, the eigenvalues are typically very different in magnitude; in extreme cases the sphere may be a thin ellipsoid, in which case convergence to the minimum may be very slow or may even fail. This is expressed in the following theorem, which is valid for the linear case:

First Theorem of Convergence for Quadratic P: The eigenspectrum of the matrix A defines the convergence of the steepest descent method as follows:

$$P(x_k) + \frac{1}{2}\mathbf{b}^T \mathbf{A}^{-1}\mathbf{b} \le \left(1 - \frac{1}{\kappa_2(\mathbf{A})}\right) \left(P(x_{k-1}) + \frac{1}{2}\mathbf{b}^T \mathbf{A}^{-1}\mathbf{b}\right).$$

Here

$$\kappa_2(A) = \frac{\lambda_{max}}{\lambda_{min}}$$

is the condition number of the *positive-definite* symmetric matrix \mathbf{A} , and if it is large the convergence is very slow. The condition number expresses the aspect ratio of the ellipsoid, i.e., the ratio of its two semi-axes.

The step size α can be chosen properly so that a monotonic convergence is guaranteed. The following theorem presents the appropriate choice for α .

Second Theorem of Convergence for Quadratic P: Consider the vector $\mathbf{f} \equiv \nabla P = \mathbf{A}\mathbf{x} - \mathbf{b}$. By computing the step size in the steepest descent method from

$$\alpha = \frac{\mathbf{f}^T \mathbf{f}}{\mathbf{f}^T \mathbf{A} \mathbf{f}} \tag{4.7}$$

then

$$P(\mathbf{x}) = P(\mathbf{x} - 2\alpha \mathbf{f}); \quad and \quad P(\mathbf{x} - \alpha \mathbf{f}) - P(\mathbf{x}) \le 0$$

for an arbitrary vector \mathbf{x} .

This theorem simply states that if we move by an amount 2α along the steepest descent, we will end up on the other side of the ellipsoid formed by the iso-contours of $P(\mathbf{x})$. If we move only by an amount α , then we guarantee that we get closer to the global minimum. This is proven by considering:

$$P(\mathbf{x} - 2\alpha\mathbf{f}) = \frac{1}{2}(\mathbf{x} - 2\alpha\mathbf{f})^T\mathbf{A}(\mathbf{x} - 2\alpha\mathbf{f}) - (\mathbf{x} - 2\alpha\mathbf{f})^T\mathbf{b}$$

which is re-arranged to

$$P(\mathbf{x} - 2\alpha \mathbf{f}) = P(\mathbf{x}) - 2\alpha \mathbf{f}^T \mathbf{f} + 2\alpha^2 \mathbf{f}^T \mathbf{A} \mathbf{f},$$

so by choosing α as suggested by the theorem we prove the first statement. The second statement is straightforward to prove with this choice of α , and upon substitution we obtain

$$P(\mathbf{x} - \alpha \mathbf{f}) = P(\mathbf{x}) - \frac{1}{2} \frac{(\mathbf{f}^T \mathbf{f})^2}{\mathbf{f}^T \mathbf{A} \mathbf{f}} \le P(\mathbf{x}).$$

The steepest descent algorithm with adaptive step size α for the linear system $f \equiv \mathbf{A}\mathbf{x} - \mathbf{b} = 0$ can then be summarized as follows:

Steepest Descent Algorithm

• Initialize : Choose $\mathbf{x}_0 \Rightarrow \mathbf{f}_0 = \mathbf{A}\mathbf{x}_0 - \mathbf{b}$.

•Begin Loop: for
$$n = 1, 2, ...$$

$$\alpha_n = \frac{\mathbf{f}_n^T \mathbf{f}_n}{\mathbf{f}_n^T \mathbf{A} \mathbf{f}_n}$$

$$\mathbf{x}_{n+1} = \mathbf{x}_n - \alpha_n \mathbf{f}_n$$

$$\mathbf{f}_{n+1} = \mathbf{A} \mathbf{x}_n - \mathbf{b}$$
endfor

• End Loop

For nonlinear systems the above algorithm can also be used but the step size α_n may not be optimum and convergence could be slow.

Conjugate Gradient Method - CGM

A more effective method than the steepest descent is one that takes not only adaptive steps but also turns adaptively in the pursuit of the minimum of the parent function. To this end, we can improve the search direction so instead of $\mathbf{f}(\mathbf{x}_n)$, we follow the direction of the steepest descent \mathbf{p} , which is conjugate to the previous search directions

$$\mathbf{p}_n = -\mathbf{f}_n + \beta_{n-1} \mathbf{p}_{n-1} \tag{4.8}$$

where β_{n-1} changes in each iteration adaptively and can be computed from the projection

$$\beta_{n-1} = \frac{\mathbf{f}_n^T (\mathbf{f}_n - \mathbf{f}_{n-1})}{\mathbf{f}_{n-1}^T \mathbf{f}_{n-1}}$$
(4.9)

as we will show later. The conjugate gradient formula is then

$$\mathbf{x}_{n+1} - \mathbf{x}_n = \alpha_n \mathbf{p}_n, \tag{4.10}$$

where \mathbf{p}_n is computed from above and the step size α_n could also be computed adaptively. For nonlinear problems there are no known formulas that can do that but for linear problems there is a nice theory that we will present next. We note that for $\beta_{n-1} = 0$ we recover the steepest descent method.

CGM for Linear Systems

We will present several different solution algorithms for linear systems in chapter 9, but CGM is one of the very best. The success of the conjugate gradient method in obtaining solutions of systems of nonlinear equations fast is based upon the theory of conjugate gradients for linear systems of equations and the remarkable properties that are inherent in this method. In the following, we develop this theory for the linear system

$$\mathbf{A}\mathbf{x} = \mathbf{b} \Rightarrow \mathbf{r} \equiv \mathbf{b} - \mathbf{A}\mathbf{x}$$

where \mathbf{r} is defined as the residual. It is zero exactly when we reach the minimum of the the quadratic parent function

$$P(\mathbf{x}) = \frac{1}{2}\mathbf{x}^T \mathbf{A} \mathbf{x} - \mathbf{x}^T \mathbf{b},$$

where we assume that **A** is positive-definite and symmetric.

First, we need to define conjugate directions. In general, the vectors \mathbf{p} and \mathbf{q} are said to be conjugate or A-orthogonal if

$$\mathbf{p}^T \mathbf{A} \mathbf{q} = 0.$$

The idea of the conjugate gradient method is to perform searches in a set of conjugate directions \mathbf{p}_i satisfying the A-orthogonality condition, i.e.,

$$\mathbf{p}_i^T \mathbf{A} \mathbf{p}_i = 0.$$

As an example, the eigenvectors of the matrix A satisfy this property since

$$\mathbf{v}_i^T \mathbf{A} \mathbf{v}_j = \mathbf{v}_i^T \lambda \mathbf{v}_j = \lambda \mathbf{v}_i^T \mathbf{v}_j = 0, \quad i \neq j.$$

Let us now assume that we have a symmetric positive-definite matrix **A** which has size $n \times n$, then at each iteration (k+1) we obtain the conjugate direction, that is the solution, from

$$\mathbf{x}_{k+1} = \mathbf{x}_k + \alpha \mathbf{p}_k,$$

where α will be chosen to minimize the quadratic functional, and \mathbf{p}_k will be computed adaptively, as we will see below. First, we can show that

$$P(\mathbf{x}_{k+1}) = P(\mathbf{x}_k) + \frac{1}{2}\alpha^2 \mathbf{p}_k^T \mathbf{A} \mathbf{p}_k - \alpha \mathbf{p}_k^T \mathbf{r}_k,$$

where we have used $\mathbf{r}_k = \mathbf{b} - \mathbf{A}\mathbf{x}_k$. Next, we minimize the quadratic P with respect to α by setting

 $\frac{\partial P}{\partial \alpha} = 0,$

which leads to

$$\alpha_k \equiv \alpha = \frac{(\mathbf{p}_k, \mathbf{r}_k)}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)}.$$

Here we have defined the inner (dot) product

$$(\mathbf{a}, \mathbf{b}) \equiv \mathbf{a}^T \mathbf{b}$$

to simplify the notation, and we will use that in the following.

Theorem on Conjugate Directions: Let **A** be a symmetric positive-definite matrix of size $n \times n$. Then after n conjugate direction searches in the n-dimensional space, we obtain $\mathbf{r}_n = \mathbf{0}$, and thus we reach a minimum of the parent function $P(\mathbf{x})$.

Proof: We will make use of the A-orthogonality property defining the conjugate directions. First, from the repeated action of the iteration $\mathbf{x}_k = \mathbf{x}_{k-1} + \alpha_{k-1} \mathbf{p}_{k-1}$ we obtain

$$\mathbf{x}_k - \mathbf{x}_0 = \sum_{i=0}^{k-1} \alpha_i \mathbf{p}_i \tag{4.11}$$

where \mathbf{x}_0 is the initial (arbitrary) guess and

$$\alpha_i = \frac{(\mathbf{p}_i, \mathbf{r}_i)}{(\mathbf{p}_i, \mathbf{A}\mathbf{p}_i)}$$

as determined from the minimization of the quadratic form.

Let us now consider the (unknown) solution \mathbf{s} and express it in terms of the n A-orthogonal vectors that form the conjugate directions. This is possible as these vectors are *linearly independent* due to the A-orthogonality. Specifically, we offset the solution by the initial guess and then expand it as follows

$$\mathbf{s} - \mathbf{x}_0 = \sum_{j=0}^{n-1} \gamma_j \mathbf{p}_j,$$

where from the orthogonality again we can compute the coefficients

$$\gamma_j = \frac{(\mathbf{p}_j, \mathbf{A}(\mathbf{s} - \mathbf{x}_0))}{(\mathbf{p}_j, \mathbf{A}\mathbf{p}_j)}.$$

Next, we consider the coefficient γ_k and subtract-off the scalar

$$\frac{(\mathbf{p}_k, \mathbf{A}(\mathbf{x}_k - \mathbf{x}_0))}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)},$$

which is zero considering equation (4.11). We can then combine the two to get

$$\gamma_k = \frac{(\mathbf{p}_k, \mathbf{A}(\mathbf{s} - \mathbf{x}_0))}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)} - \frac{(\mathbf{p}_k, \mathbf{A}(\mathbf{x}_k - \mathbf{x}_0))}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)} \\
= \frac{(\mathbf{p}_k, \mathbf{A}(\mathbf{s} - \mathbf{x}_k))}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)} \\
= \frac{(\mathbf{p}_k, \mathbf{r}_k)}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)},$$

where we have used the definition of the residual, i.e., $\mathbf{r}_k = \mathbf{b} - \mathbf{A}\mathbf{x}_k = \mathbf{A}(\mathbf{s} - \mathbf{x}_k)$.

Now by comparing the coefficients γ_k and α_k we see that they are identical and therefore the n-term expansion of the conjugate direction iteration $(\mathbf{x}_k - \mathbf{x}_0)$ is identical to the exact solution $(\mathbf{s} - \mathbf{x}_0)$. This result is indeed remarkable!

While the above theorem is valid for any set of conjugate directions, efficiency is what distinguishes the conjugate gradient algorithm. For example, the use of the eigenvectors of \mathbf{A} would be prohibitively expensive as it takes much more computational work to compute the eigenvectors of the matrix than solving the linear system! Another approach would be to use the Gram-Schmidt algorithm to obtain conjugate directions but this is also very expensive and would amount to $\mathcal{O}(n^4)$ work.

To this end, the idea of Hestenes & Stiefel [55] to compute iteratively the search directions \mathbf{p} is the keystone of this method. It is very efficient and requires the storage of two or three vectors only. Initially, we set $\mathbf{p}_0 = \mathbf{r}_0$, and then we iterate

$$\mathbf{p}_{k+1} = \mathbf{r}_{k+1} + \beta_k \mathbf{p}_k,$$

where we need to determine the scalar, β_k , by using the A-conjugate property of the search directions \mathbf{p}_k . For this, we take the inner product of the above equation with \mathbf{p}_k^T and impose A-orthogonality on the left-hand-side

$$0 = (\mathbf{p}_k, \mathbf{A}\mathbf{p}_{k+1}) = (\mathbf{p}_k, \mathbf{A}\mathbf{r}_{k+1}) + \beta_k(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k),$$

and this leads to

$$\beta_k = -\frac{(\mathbf{p}_k, \mathbf{A}\mathbf{r}_{k+1})}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)} = -\frac{(\mathbf{r}_{k+1}, \mathbf{A}\mathbf{p}_k)}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)},$$

which can always be computed since the denominator is guaranteed to be non-zero. The last equality is valid since A is symmetric.

Next, we prove some useful relationships between the orthogonal directions and the residual which can be used to reduce even further the computational complexity of the conjugate gradient algorithm. First, it is clear that using its definition we can compute the residual

iteratively as well, as follows

$$\mathbf{r}_{k+1} = \mathbf{b} - \mathbf{A}\mathbf{x}_{k+1}$$

$$= \mathbf{b} - \mathbf{A}(\mathbf{x}_k + \alpha_k \mathbf{p}_k)$$

$$= (\mathbf{b} - \mathbf{A}\mathbf{x}_k) - \alpha_k \mathbf{A}\mathbf{p}_k$$

$$= \mathbf{r}_k - \alpha_k \mathbf{A}\mathbf{p}_k.$$

Using the above equation we now show that

$$(\mathbf{r}_i, \mathbf{p}_i) = 0, \quad i \neq j \tag{4.12}$$

$$(\mathbf{r}_i, \mathbf{p}_i) = (\mathbf{r}_i, \mathbf{r}_i) \tag{4.13}$$

$$(\mathbf{r}_i, \mathbf{r}_j) = 0, \quad i \neq j \tag{4.14}$$

This is shown by induction, that is assuming that the first equation above is valid we will also show that $(\mathbf{r}_{i+1}, \mathbf{p}_j) = 0$. We substitute \mathbf{r}_{i+1} from the iteration equation above and then it becomes obvious; we proceed similarly for the other two equations.

Based on these relations, we can also prove that β_k is the ratio of the square of the magnitudes of two successive gradients, i.e.,

$$\beta_k = -\frac{(\mathbf{p}_k, \mathbf{A}\mathbf{r}_{k+1})}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)} = \frac{(\mathbf{r}_{k+1}, \mathbf{r}_{k+1})}{(\mathbf{r}_k, \mathbf{r}_k)}.$$

We now finalize the CG algorithm for linear systems of equations:

Conjugate Gradient Algorithm

• Initialize : Choose
$$\mathbf{x}_0 \Rightarrow \mathbf{p}_0 = \mathbf{r}_0 = \mathbf{b} - \mathbf{A}\mathbf{x}_0$$
.

• Begin Loop: for
$$k = 1l$$
to n

$$\alpha_k = \frac{(\mathbf{r}_k, \mathbf{r}_k)}{(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k)}$$

$$\mathbf{x}_{k+1} = \mathbf{x}_k + \alpha_k \mathbf{p}_k$$

$$\mathbf{r}_{k+1} = \mathbf{r}_k - \alpha_k \mathbf{A} \mathbf{p}_k$$

$$\beta_k = \frac{(\mathbf{r}_{k+1}, \mathbf{r}_{k+1})}{(\mathbf{r}_k, \mathbf{r}_k)}$$

$$\mathbf{p}_{k+1} = \mathbf{r}_{k+1} + \beta_k \mathbf{p}_k$$

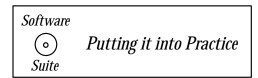
endfor

 $\bullet End\ Loop$

This code represents the ideal algorithm that terminates after exactly n iterations; it assumes that the orthogonality of the conjugate directions is preserved independently of the

matrix \mathbf{A} , i.e., its size and its structure. In practice, however round-off errors may destroy such orthogonality resulting in incomplete convergence. The convergence process then is controlled by the condition number of \mathbf{A} as discussed in chapter 9, where preconditioning techniques for convergence acceleration are also presented.

The simplicity of the CG algorithm both in terms of coding as well as in terms of computational complexity is amazing! We only need to perform one matrix-vector product, i.e., \mathbf{Ap}_k , one dot product, and three daxpy's operations, all of which can be implemented very efficiently using the BLAS routines discussed in section 2.2.7.



Here we provide our implementation of a non-preconditioned ConjugateGradient routine. We also use a stopping criterion for convergence in anticipation of round-off errors. Our program can then be terminated before or after n iterations, depending on the matrix \mathbf{A} and the tolerance level we use. For this implementation, we are using classes from the software suite: namely, SCVector and SCMatrix.

The beautiful thing about the implementation below is that by using classes (and all the mechanics that are available for using classes in C++), the CG method can be described succinctly in a few lines of code. Below, we present our implementation:

```
SCVector ConjugateGradient(SCMatrix A, SCVector b, SCVector x0){
  int dim = x0.Dimension();
  const double tolerance = 1.0e-14;
  SCVector x(dim),r(dim),v(dim),z(dim);
  double c,t,d;
  x = x0;
  r = b - A*x;
  v = r;
  c = dot(r,r);
  for(int i=0;i<dim;i++){</pre>
    if(sqrt(dot(v,v))<tolerance){</pre>
      cerr << "Error in ConjugateGradient: execution "</pre>
      cerr << "of function terminated" << endl;</pre>
      break;
    }
    z = A*v;
```

```
t = c/dot(v,z);
x = x + t*v;
r = r - t*z;
d = dot(r,r);
if(sqrt(d) < tolerance)
    break;
v = r + (d/c)*v;
c = d;
}
return x;
}</pre>
```

In this function, we have used classes for encapsulating the ideas of a vector and a matrix, and we have used the idea of *overloaded operators* to perform the mathematical operations necessary. Notice that we are using operators like '+', '-' and '*' between variables of type SCVector and SCVector, and between SCMatrix and SCVector. As we discussed in 3.1.8, this is accomplished by overloading the operators, that is, extending the definition of these operators to include operations between our newly created data types.

We now present a simple main driving program which uses the function above.

```
#include <iostream.h>
#include "SCmathlib.h"
#include "SCchapter4.h"
int main(int argc, char * argv[]){
  int dim = 4;
  SCVector x(dim),b(dim),x0(dim);
  SCMatrix A(dim);
  // Set our initial guess
  x0(0) = x0(1) = x0(2) = x0(3) = 1.0;
  for(int i=0;i<dim;i++){
    for(int j=0; j<dim; j++){</pre>
      A(i,j) = 1+(i+1)*(j+1);
      /* We do this to make sure that the symmetric matrix that
         we create has a determinant which is non-zero */
      if(i==3 \&\& j == 2)
        A(i,j) = 12;
      if(i==2 \&\& j == 3)
        A(i,j) = 12;
```

}

```
}
}
cout << "The Matrix A that we are using: " << endl;</pre>
A.Print();
cout << endl;</pre>
SCVector y(dim);
y(0) = 2.;
y(1) = -3.;
y(2) = 5.43;
y(3) = -22.56;
cout << "The exact solution is: " << endl;</pre>
y.Print();
cout << endl;</pre>
b = A*y;
cout << "The right hand side, b, of Ax=b: " << endl;</pre>
b.Print();
cout << endl;</pre>
x = ConjugateGradient(A,b,x0);
cout << "The approximate solution using CG is: " << endl;</pre>
x.Print();
cout << endl;</pre>
```

4.2 Numerical Integration Methods

Approximate integration of a function is a very old subject. It was first performed rigorously by Archimedes who used the method of inscribed and circumscribed polygons to obtain lower and upper bounds for the value of the area of a circle. He also computed the center of mass and center of buoyancy for many complicated figures, a task that requires accurate integration. Simpson suggested a very accurate formula in the mid-eighteen century, and Gauss proposed his famous quadrature rules in the early part of the nineteen century. Numerical discretization methods such as finite elements and boundary elements, in particular, depend critically on efficient numerical integration procedures. There exist both simple quadrature rules as well as more advanced approaches, which we present next. More details on integration can be found in the books of Davis & Rabinowitz [23] and Ghizzetti & Ossicini [43].

4.2.1 Simple Integration Algorithms

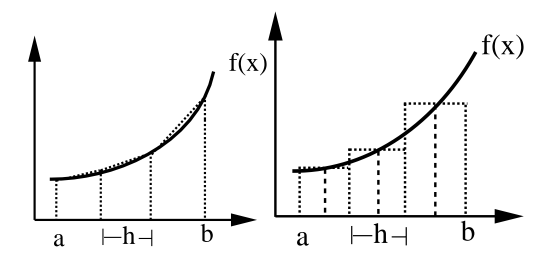


Figure 4.3: Trapezoid (left) versus midpoint-rectangle (right) rule: The accuracy is of the same order, $\mathcal{O}(h^2)$ for both, but surprisingly the midpoint-rectangle wins by a factor of two.

The simplest quadrature formulas are based on either piecewise constant approximations of a function in the interval of interest

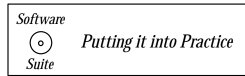
$$I = \int_{a}^{b} f(x)dx,$$

as shown schematically in figure 4.3 (right) or on piecewise linear approximations (left). Assuming that h is the size of each of the n equal cells involved in the discretization, we obtain the approximations:

• Midpoint-Rectangle Rule:

$$I \approx R(h) = h \sum_{i=1}^{n} f_{i-1/2}$$
 (4.15)

This is implemented in the code below.

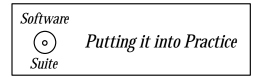


We present a functional implementation of the Midpoint Rule. Notice that we are continuing to use the concept of passing functions to functions.

• Trapezoid Rule:

$$I \approx T(h) = h\left[\frac{1}{2}f_0 + f_1 + \dots + f_{n-1} + \frac{1}{2}f_n\right].$$
 (4.16)

This is implemented in the code below.



We present a functional implementation of the Trapezoidal Rule. Observe the use of short-hand operators in the function below. We are able to short-hand both addition and multiplication.

Note that, by definition, the following identity applies

$$T(h) = \frac{1}{2}[T(2h) + R(2h)],$$

so we can easily relate the two formulas. Intuitively, one would guess that the trapezoid rule may be more accurate than the midpoint-rectangle because of the higher-order approximation involved, i.e. linear versus constants. However, as we will see below the quadrature error in the trapezoid rule is twice the quadrature error in the midpoint-rectangle rule!

Quadrature Error

To obtain the quadrature error for the above methods we assume that f''(x) is continuous in [a, b]. We then consider a grid consisting of the (m + 1) points

$$[x_0, x_1, \ldots, x_m],$$

and construct the unique polynomial

$$p(x) = c_0 + c_1(x - x_0) + c_2(x - x_0)(x - x_1) + \dots + c_m(x - x_0)(x - x_1) \dots (x - x_{m-1})$$
(4.17)

so that $p(x_i) = f(x_i)$, i = 0, 1, ..., m. This is Newton's formula of approximating f(x) in the interval [a, b] where $x_0 = a$ and $x_m = b$. Then, if f(x) possesses continuous derivatives of order at least (m + 1), we obtain

$$f(x) - p(x) = \frac{f^{(m+1)}(\xi)}{(m+1)!} (x - x_0)(x - x_1) \dots (x - x_m)$$
(4.18)

for some point $\xi \in [a, b]$. We can then obtain the quadrature error by simply integrating the right-hand-side in the above equation and by providing an upper bound for the magnitude

of the (m+1) derivative.

For the **Trapezoid** rule we obtain on each cell *i*:

$$\epsilon_i^T = T_i(h) - \int_{x_{i-1}}^{x_i} f(x)dx = -\int_{x_{i-1}}^{x_i} \frac{f''(\xi)}{2} (x - x_{i-1})(x - x_i)dx, \quad \xi \in [x_{i-1}, x_i]$$

and thus

$$\epsilon_i^T \approx -\frac{1}{2}f''(\xi_i) \int_{x_{i-1}}^{x_i} (x - x_{i-1})(x - x_i) dx.$$

Let

$$x = x_{i-1} + h s$$
, $\epsilon_i^T \approx -\frac{1}{2} f''(\xi_i) \int_0^1 h s h(s-1) h ds = \frac{h^3 f''(\xi_i)}{12}$.

The above is the *local quadrature error* on the cell i. We can easily obtain the *global error* by summing over all the cells i, i.e.

$$\epsilon^T = \sum_{i=1}^n \epsilon_i^T \approx f''(\xi) \frac{(b-a)h^2}{12},$$

where

$$b - a = \sum_{i=1}^{n} h.$$

Therefore, the trapezoid rule results in second-order convergence rate $\mathcal{O}(h^2)$ for the global error.

For the **Midpoint-Rectangle** rule we need to expand the function around the midpoint $x_0 = (x_{i-1} + x_i)/2$ in the cell i:

$$f(x) \approx f(x_0) + (x - x_0)f'(x_0) + \frac{(x - x_0)^2}{2}f''(\xi),$$

where $\xi \in [a, b]$. Then the error for the integral $R_i(h)$ is

$$\epsilon_i^R = R_i(h) - \int_{x_{i-1}}^{x_i} f(x)dx = -\int_{x_{i-1}}^{x_i} f'(x_0)(x - x_0)dx - \int_{x_{i-1}}^{x_i} \frac{f''(\xi)}{2}(x - x_0)^2 dx$$
$$= f''(\xi) \frac{h^3}{24}.$$

The above is the *local quadrature error* on the cell i. We can easily obtain the *global error* by summing over all the cells i, i.e.

$$\epsilon^R = \sum_{i=1}^n \epsilon_i^R \approx f''(\xi) \frac{(b-a)h^2}{24}.$$

Comparing now the global quadrature errors e^R and e^T we see that, surprisingly, the midpoint-rectangle rule approximates the integral I better, as its corresponding error is half of that for the trapezoid rule!

Better Trapezoid Rules

We present here two efficient ways of improving the second-order accuracy of the trapezoid rule by

- 1. Correcting the end-values,
- 2. Interval subdivision.

First, the **corrected trapezoid rule** is based on a trigonometric interpolation of the function, unlike the Newton linear approximation used for the standard trapezoid (equation (4.17) with m = 1). It adds to the standard trapezoid formula a correction term at the ends, i.e.

$$T_C(h) = h\left[\frac{1}{2}f_0 + f_1 + \dots + f_{n-1} + \frac{1}{2}f_n\right] + \frac{h^2}{12}[f'(a) - f'(b)], \tag{4.19}$$

with corresponding quadrature error

$$\epsilon^{TC} \approx \frac{h^4}{720} \times [f^{'''}(b) - f^{'''}(a)].$$

Clearly, for functions which are periodic in the interval [a, b], a very high degree of accuracy is achieved without the addition of the extra correction terms.

Another efficient method to increase the formal order of accuracy of the trapezoid rule is **Romberg's method**. This method, which is also called extrapolation to the limit, is a systematic procedure of using subdivision of intervals with the trapezoid rule. It is based on Richardson's extrapolation idea which is used most often in numerical differentiation.

Let us define a series of trapezoid sums

$$T_0, T_1, T_2, \ldots$$

by successively bisecting the interval of integration

$$T_{0} = \frac{h_{0}}{2} [f(x_{0}) + f(x_{0} + h_{0})]$$

$$T_{1} = \frac{h_{1}}{2} [f(x_{0}) + 2f(x_{0} + h_{1}) + f(x_{0} + 2h_{1})]$$

$$\vdots :$$

$$T_{k} = \frac{h_{k}}{2} [f(x_{0}) + 2\sum_{i=1}^{2^{k}-1} f(x_{0} + ih_{k}) + f(x_{0} + 2^{k}h_{k})].$$

Then the error on each grid (k) is: $\epsilon_k = I - T_k$, where I is the exact value of the integral. Because of the quadratic convergence, we have

$$\epsilon_{k+1} pprox rac{1}{4} \epsilon_k.$$

We also have for $k = 0, 1, \dots$

$$I = T_0 + \epsilon_0$$

 $I = T_1 + \epsilon_1 \simeq T_1 + \frac{1}{4}\epsilon_0 = T_1 + \frac{1}{4}(I - T_0) \Rightarrow I \simeq \frac{4T_1 - T_0}{3}$.

In general, we have by induction

$$I_k^1 = \frac{4T_k - T_{k-1}}{3} \,,$$

which defines a *one-step correction*. Expanding T_k and T_{k-1} and substituting in the above formula we obtain a quadrature error for I_k^1 which is proportional to h^4 , competitive with Simpson's rule (see below). Thus, the error in successive terms of I_k^1 is reduced by a factor of 1/16 and we obtain a *two-step correction*:

$$I_k^2 = \frac{16I_k^1 - I_{k-1}^1}{15} \,.$$

Continuing this process (m) times, we get the recurrence formula

$$I_k^m = \frac{4^m I_k^{m-1} - I_{k-1}^{m-1}}{4^m - 1}, \quad m \ge 1, \ k \ge m.$$
 (4.20)

To completely define the recursive process, we also need to specify the initial conditions, i.e.,

$$I_k^0 = T_k; \quad I_{k-1}^0 = T_{k-1}.$$

The corresponding quadrature error in Romberg's method is

$$\epsilon_k^m \sim h^{2m+2} \cdot f^{(2m+2)}(\xi).$$

This method uses a fine grid (k) and a coarse grid (k-1) and extrapolates the standard trapezoid rule. The reason why this algorithm works is because we know that the error structure has the polynomial form

$$c_1h^2 + c_2h^4 + c_3h^6 + \dots$$

for the trapezoid rule. So the idea of the method is to use more than one grid, a *fine* grid and a coarse grid, and then at the first correction level construct the extrapolant, at the second correction level construct the extrapolant of the extrapolant, and so on. By weighting appropriately these successive constructs, we can extrapolate the resulted quadrature values to higher accuracy.

In theory, Romberg's method gives arbitrarily high convergence rate but in practice round-off error slows down convergence and accuracy.

We present a functional implementation of Romberg's integration. Notice that this implementation consists of two components:

Software

Outling it into Practice

Suite

- 1. An implementation of the trapezoid rule.
- 2. A recursive definition for Romberg integration.

In the case below, the stopping condition for the recursion is when the variable m is equal to zero. When this is true, we execute the trapezoid rule. For all other valid values of m we recursively call the Romberg function with appropriately changed arguments.

```
double Romberg(int m, int k, double xleft, double xright,
               double (*func)(double)){
  double RI, I1, I2;
  double coeff = pow(4.0,m);
  if(k < m){
    cerr << "ROMBERG::Value of k must be >= m; setting k=m\n";
  }
  if(m==0){
   RI = TrapezoidRule(k,xleft,xright,func);
  }
  else{
    I1 = Romberg(m-1,k, xleft,xright,func);
    I2 = Romberg(m-1,k-1,xleft,xright,func);
   RI = (coeff*I1 - I2)/(coeff-1.0);
  }
  return RI;
}
```

Simpson's Rule

The idea in Simpson's rule of integration is to connect three consecutive ordinates by a parabola and use Newton's approximation formula to obtain the quadrature error. The standard or 1/3 Simpson's rule employs an even number of cells. For the *i* cell defined by $[x_{i-1}, x_i, x_{i+1}]$ we obtain

$$\int_{i-1}^{i+1} f(x)dx \approx S_i(h) = \frac{h}{3} [f_{i-1} + 4f_i + f_{i+1}], \tag{4.21}$$

and the Simpson quadrature for the entire interval [a, b] is

$$S(h) = \frac{h}{3} [f_0 + 4f_1 + 2f_2 + \dots 4f_{m-1} + f_m]$$
(4.22)

where m is an even number.

To obtain the error in Simpson's formula we employ equation (4.17) with m=2, then

$$\epsilon_i^S = h \int_0^2 \frac{t(t-1)(t-2)}{6} h^3 f'''(\xi) dt = 0,$$

thus the leading term in the error is zero, so we have to integrate the next term

$$\epsilon_i^S = h \int_0^2 \frac{t(t-1)(t-2)(t-3)}{24} h^4 f^{(4)}(\xi) dt = -\frac{h^5}{90} f^{(4)}(\xi),$$

so the local error is $\mathcal{O}(h^5)$. The *global error* can be easily obtained by replacing h=(b-a)/2, i.e.

$$\epsilon^S = -\frac{(b-a)}{2880} h^4 f^{(4)}(\xi).$$

It is interesting to note that in Simpson's rule we obtain $\mathcal{O}(h^4)$ convergence with a quadratic polynomial fit as we take advantage of the *even* number of cells and the corresponding cancellation in the error terms. In fact, a cubic polynomial fit in equation (4.17) leads to the 3/8 rule

$$S_i(h) = \frac{3}{8}h[f_{i-1} + 3f_i + f_{i+1}]$$

with an error

$$\epsilon_i^S = -\frac{3h^5}{80}f^{(4)}(\xi),$$

which is larger than the local error of the 1/3 Simpson's rule corresponding to lower order polynomial approximation. However, this rule is useful when an *odd number of cells* is required in the integration, which can be broken up into two subintervals: The first one consists of 3 cells (3/8 rule), and the second one consists of the rest, which corresponds to an even number of cells where we apply the standard 1/3 Simpson's rule.

Example



As an example, we put into practice several of the functions implemented above (see *software suite*). We are attempting to approximate the integral of x^4 on the interval [0,1], for which we know the exact value to be 0.2. In the program, we compute the midpoint rule, trapezoid rule, and Simpson's rule approximation for ten levels. The output of the program is the following table:

Level	Midpoint	Trapezoidal	Simpsons
0	1.375000e-01	3.000000e-01	8.333333e-03
1	3.984375e-02	8.125000e-02	5.208333e-04
2	1.030273e-02	2.070312e-02	3.255208e-05
3	2.597046e-03	5.200195e-03	2.034505e-06
4	6.505966e-04	1.301575e-03	1.271566e-07
5	1.627326e-04	3.254890e-04	7.947286e-09

6	4.068837e-05	8.137822e-05	4.967053e-10
7	1.017242e-05	2.034493e-05	3.104406e-11
8	2.543125e-06	5.086255e-06	1.940226e-12
9	6.357825e-07	1.271565e-06	1.212364e-13

From the data presented above, we observe the following:

- The trapezoid rule has, in fact, an error which is about twice that of the midpoint rule, just as predicted by the theory!
- Both the midpoint rule and the trapezoid rule exhibit second-order convergence, as predicted by the theory!
- For Simpson's rule, we obtain fourth-order convergence, as predicted by the theory!

Key Concept

• Theoretical results should be used to test the validity of your implementation.

4.2.2 Advanced Quadrature Rules

All integration methods in the previous section are based on Newton's formula and loworder polynomial approximation. We can extend these methods to high-order polynomial interpolation using Lagrangian interpolation, see section 3.1. The objective is to maximize the order of accuracy for a fixed number of points whose locations are allowed to vary. This leads us to **Gauss quadrature**. For example, suppose that we want to evaluate the integral

$$\int_0^1 \frac{1+x}{\sqrt{x}},$$

the exact value of which is 8/3. We can only use n=2 quadrature points and Gauss integration to compute the above integral exactly!

The important difference with the methods of the previous section is the location of the quadrature points. These are special points, as we will explain, and typically they are roots of an appropriate orthogonal polynomial. The Legendre polynomial is the most-often used but Chebyshev, Laguerre and Hermite polynomials are also used. For the aforementioned integral, the two special points are determined from the roots of the second-order Legendre polynomial

$$P_2(s) = \frac{1}{2}(3s^2 - 1), \quad s \in [-1, 1]$$

i.e., $s_{\pm} = \pm 1/\sqrt{3}$. These special points need to be mapped in the interval of integration [a, b], so we obtain

$$x = \frac{a+b}{2} + \frac{b-a}{2}s$$
, where $x \in [a,b]$ and $s \in [-1,1]$.

Let us now derive the general Gauss quadrature. We assume that the values

$$f(x_1), f(x_2), \ldots, f(x_n)$$

cannot provide sufficient information for determining f(x). Let $x = x_k$ and correspondingly $y_k \equiv f(x_k)$, and determine the polynomial $p_{n-1}(x)$ which fits the coordinates $y_1, y_2 \ldots, y_n$. We can employ Lagrange's interpolation by constructing the fundamental polynomial

$$F_n(x) = (x - x_1)(x - x_2) \dots (x - x_n)$$

and define

$$h_i(x) = \frac{1}{F'_n(x_i)} \frac{F_n(x)}{x - x_i}, \quad i = 1, 2, \dots, n$$

$$h_i(x_k) = 0; \quad h_i(x_i) = 1, \text{ by construction.}$$

We then obtain the polynomial

$$p_{n-1}(x) = y_1 h_1(x) + y_2 h_2(x) + \ldots + y_n h_n(x)$$

where

$$p_{n-1}(x_k) = y_k.$$

The Gauss integral can then be approximated as:

$$I_G = \int_{-1}^{+1} p_{n-1}(x) dx = \sum_{k=1}^{n} y_k \int_{-1}^{1} h_k(x) dx = \sum_{k=1}^{n} y_k w_k,$$

where

$$w_k \equiv \int_{-1}^{+1} h_k(x) dx$$

are the weights of integration, which are independent of the integrand.

Gauss's idea is then to add an extra point x_{n+1} without changing $x_k, k \leq n$ or equivalently the term

$$h_{n+1}(x) = \frac{F_n(x)}{F'_{n+1}(x_{n+1})},$$

and thus the (n+1) weight

$$w_{n+1} \sim \int_{-1}^{1} F_n(x) dx$$
.

Similarly, if we add m new points

$$\underbrace{x_{n+1}, x_{n+2}, \dots, x_{n+m}}_{m \text{ points}}$$

then the $(n+1)^{th}$ weight is obtained from

$$w_{n+1} = \int_{-1}^{+1} F_n(x) G_{m-1}^i(x) dx$$

where $G_{m-1}^i(x)$ is a polynomial of order (m-1). Now, if we impose that all moments of $F_n(x)$ up to (m-1) are zero, i.e.,

$$\int_{-1}^{1} F_n(x) x^k dx, \quad k = 0, 1, \dots, m - 1$$

then orthogonality leads to

$$\int_{-1}^{+1} F_n(x) G_{m-1}^i(x) dx = 0,$$

since G is a linear combination of powers x^k . We can add up to n points to the original grid, and thus we can effectively double the number of points considered with only half the weights!

Therefore, the Gauss formula

$$I_G = \sum_{k=1}^n y_k w_k$$

that employs a grid of n points results in quadrature accuracy equivalent to a grid corresponding to 2n coordinates using the simple rules of the previous section.

Jacobi Polynomials

Jacobi polynomials $P_n^{\alpha,\beta}(x)$ are a family of polynomial solutions to the singular Sturm-Liouville problem. A significant feature of these polynomials is that they are orthogonal in the interval [-1,1] with respect to the function $(1-x)^{\alpha}(1+x)^{\beta}$ $(\alpha,\beta>-1)$. We have already presented the Chebyshev polynomials in section 3.1.5 – they are a subset of Jacobi polynomials for the special case that $\alpha=\beta=-\frac{1}{2}$. A detail account of their properties can be found in Abramowitz & Stegun ([1]: chapter 22) and also in Ghizzetti & Ossicini ([43]: chapter 3.4).

Differential Equation

$$(1-x)(1+x)\frac{d^2y(x)}{dx^2} + (\beta - \alpha - (\alpha + \beta + 2)x)\frac{dy(x)}{dx} = -\lambda_n y(x)$$
 (4.23)

or

$$\frac{d}{dx}\left[(1-x)^{1+\alpha}(1+x)^{1+\beta}\frac{dy(x)}{dx}\right] = -\lambda_n(1-x)^{\alpha}(1+x)^{\beta}y(x)$$

$$\lambda_n = n(n+\alpha+\beta+1)$$

$$y(x) = P_n^{\alpha,\beta}(x)$$
(4.24)

Special Cases

Legendre Polynomial
$$(\alpha = \beta = 0) \rightarrow P_n(x) = P_n^{0,0}(x)$$

Chebychev Polynomial $(\alpha = \beta = -\frac{1}{2}) \rightarrow T_n(x) = \frac{2^{2n}(n!)^2}{(2n)!}P_n^{-\frac{1}{2},-\frac{1}{2}}(x)$

Recursion Relations

$$P_0^{\alpha,\beta}(x) = 1$$

$$P_1^{\alpha,\beta}(x) = \frac{1}{2} [\alpha - \beta + (\alpha + \beta + 2)x]$$

$$a_n^1 P_{n+1}^{\alpha,\beta}(x) = (a_n^2 + a_n^3 x) P_n^{\alpha,\beta}(x) - a_n^4 P_{n-1}^{\alpha,\beta}(x)$$

$$a_n^1 = 2(n+1)(n+\alpha+\beta+1)(2n+\alpha+\beta)$$

$$a_n^2 = (2n+\alpha+\beta+1)(\alpha^2-\beta^2)$$

$$a_n^3 = (2n+\alpha+\beta)(2n+\alpha+\beta+1)(2n+\alpha+\beta+2)$$

$$a_n^4 = 2(n+\alpha)(n+\beta)(2n+\alpha+\beta+2)$$

$$(4.25)$$

$$b_{n}^{1}(x)\frac{d}{dx}P_{n}^{\alpha,\beta}(x) = b_{n}^{2}(x)P_{n}^{\alpha,\beta}(x) + b_{n}^{3}(x)P_{n-1}^{\alpha,\beta}(x)$$

$$b_{n}^{1}(x) = (2n + \alpha + \beta)(1 - x^{2})$$

$$b_{n}^{2}(x) = n[\alpha - \beta - (2n + \alpha + \beta)x]$$

$$b_{n}^{3}(x) = 2(n + \alpha)(n + \beta)$$
(4.26)

Special Values

$$P_n^{\alpha,\beta}(1) = \binom{n+\alpha}{n} = \frac{(n+\alpha)!}{\alpha!n!}$$
 (4.27)

$$P_n^{\alpha,\beta}(-x) = (-1)^n P_n^{\beta,\alpha}(x) \tag{4.28}$$

Orthogonality Relations

$$\int_{-1}^{1} (1-x)^{\alpha} (1+x)^{\beta} P_n^{\alpha,\beta}(x) P_m^{\alpha,\beta}(x) dx = 0 \qquad n \neq m$$
 (4.29)

$$\int_{-1}^{1} (1-x)^{\alpha} (1+x)^{\beta} P_n^{\alpha,\beta}(x) P_n^{\alpha,\beta}(x) dx = \frac{2^{\alpha+\beta+1}}{2n+\alpha+\beta+1} \frac{\Gamma(n+\alpha+1)\Gamma(n+\beta+1)}{n!\Gamma(n+\alpha+\beta+1)}$$

Evaluation of the Zeros of Jacobi Polynomials

The formulas for the weights of the general Jacobi polynomials (see table 4.3 for the Legendre case) have a closed form in terms of the grid points x_i . In general, however, there

are no explicit formulas for the grid points. These are defined in terms of the roots of the Jacobi polynomial such that

$$x_i = x_{i,m}^{\alpha,\beta}$$

$$P_m^{\alpha,\beta}(x_{i,m}^{\alpha,\beta}) = 0 i = 0, 1, \dots, m-1.$$

The zeros $x_{i,m}^{\alpha,\beta}$ can be numerically evaluated using an iterative technique, such as the Newton-Raphson we studied in the previous section 4.1.3. However, we note that the zeros of the Chebychev polynomial $(\alpha = \beta = -\frac{1}{2})$ do have an explicit form,

$$x_{i,m}^{-\frac{1}{2},-\frac{1}{2}} = -\cos\left(\frac{2i+1}{2m}\pi\right) \quad i = 0,\dots, m-1,$$

and so we can use $x_{i,m}^{-\frac{1}{2},-\frac{1}{2}}$ as an initial guess to the iteration.

To ensure that we find a new root at each search we can apply *polynomial deflation* or reduction, where the known roots are factored out of the initial polynomial once they have been determined. This means that the root finding algorithm is applied to the polynomial

$$f_{m-n}(x) = \frac{P_m^{\alpha,\beta}(x)}{\prod_{i=0}^{n-1} (x - x_i)},$$

where x_i (i = 0, ..., n - 1) are the known roots of $P_m^{\alpha,\beta}(x)$.

Noting that

$$\frac{f_{m-n}(x)}{f'_{m-n}(x)} = \frac{P_m^{\alpha,\beta}(x)}{[P_m^{\alpha,\beta}(x)]' - P_m^{\alpha,\beta}(x) \sum_{i=0}^{n-1} [1/(x-x_i)]},$$

a root finding algorithm to determine the m roots of $P_m^{\alpha,\beta}(x)$ using the Newton-Raphson iteration with polynomial deflation is

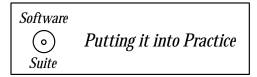
$$\begin{split} \text{for } k = 0, m-1 \\ r = x_{k,m}^{-\frac{1}{2},-\frac{1}{2}} \\ \text{if } (k>0) \ r = (r+x_{k-1})/2 \\ \text{For } j = 1, \text{stop} \\ s = \sum_{i=0}^{k-1} \frac{1}{(r-x_i)} \\ \delta = -\frac{P_m^{\alpha,\beta}(r)}{[P_m^{\alpha,\beta}(r)]' - P_m^{\alpha,\beta}(r)s} \\ r = r + \delta \\ \text{if } (\delta < \epsilon) \text{ exit loop} \\ \text{endfor} \\ x_k = r \end{split}$$
 endfor

Here ϵ is a specified tolerance. Numerically, we find that a better approximation for the initial guess is given by the average of $r = x_{k,m}^{-\frac{1}{2},-\frac{1}{2}}$ and x_{k-1} . The values of $P_m^{\alpha,\beta}(x)$ and $[P_m^{\alpha,\beta}(x)]'$ can be generated using the recursion relationships (4.25) and (4.26).

n	Abscissas x_j	Weights w_j
2	$\pm 0.577350 = \pm \frac{1}{\sqrt{3}}$	1
3	0	$\frac{8}{9}$
	± 0.774597	$\frac{5}{9}$
4	± 0.339981	0.652145
	± 0.861136	0.347855
5	0	0.568889
	± 0.538469	0.478629
	± 0.906180	0.236927

Table 4.3: Zeros of Legendre polynomials $P_n(x)$ and corresponding weights.

In the following, we first use the above formulas to compute the Jacobi polynomials and its derivatives, and subsequently implement the root finding algorithm given above.



Here we present an implementation of the Jacobi polynomials. Observe that our definition relies on the recursive nature of these polynomials, and that we have specifically implemented the three-term recurrence relation immediately into code.

```
switch(degree){
  case 0:
   value = 1.0;
   break;
  case 1:
    value = 0.5*(alpha-beta+(alpha+beta+2.0)*x);
   break;
  default:
   degm1 = degree-1.0;
   tmp = 2.0*degm1+alpha+beta;
    a1= 2.0*(degm1+1)*(degm1+alpha+beta+1)*tmp;
    a2= (tmp+1)*(alpha*alpha-beta*beta);
    a3= tmp*(tmp+1.0)*(tmp+2.0);
    a4= 2.0*(degm1+alpha)*(degm1+beta)*(tmp+2.0);
   value = ((a2+a3*x)*JacobiPoly(degree-1,x,alpha,beta)-
     a4*JacobiPoly(degree-2,x,alpha,beta))/a1;
 }
 return value;
}
```

Similarly, for the derivatives of the Jacobi polynomials, we rely on the three-term recurrence relation to provide us a fast way of implementing the derivative.

```
double JacobiPolyDerivative(int degree, double x,
                            double alpha, double beta){
  double value;
  double tmp;
  double b1,b2,b3;
  switch(degree){
  case 0:
   value = 0.0;
   break;
  default:
    tmp = 2.0*degree+alpha+beta;
   b1 = tmp*(1.0-x*x);
   b2 = degree*(alpha-beta-tmp*x);
   b3 = 2.0*(degree+alpha)*(degree+beta);
   value = (b2*JacobiPoly(degree,x,alpha,beta) +
             b3*JacobiPoly(degree-1,x,alpha,beta))/b1;
 }
```

```
return value;
}
```

To compute the zeros of the Jacobi polynomials, we use a reduction technique. In the code below, you will notice that we use two primary concepts:

- We rely on our previous function definitions for the Jacobi polynomials and their derivatives (given directly above).
- We rely on using Newton-Raphson iteration for obtaining the root.

```
void JacobiZeros(int degree, double *z, double alpha,
                 double beta){
  int i,j,k;
  const int maxit = 30;
  const double EPS = 1.0e-14;
  double dth = M_PI/(2.0*degree);
          poly,pder,rlast=0.0;
  double
  double sum, delr, r;
  double one = 1.0, two = 2.0;
  // If the degree of the polynomial is zero (or less),
  // then there are no roots
  if(degree<=0)
    return;
  for(k = 0; k < degree; k++){
    r = -\cos((two*k + one) * dth);
    if(k) r = 0.5*(r + rlast);
    for(j = 1; j < maxit; ++j){
      poly = JacobiPoly(degree,r,alpha,beta);
      pder = JacobiPolyDerivative(degree,r,alpha,beta);
      sum = 0.0;
      for(i = 0; i < k; ++i)
sum += one/(r - z[i]);
      delr = -poly / (pder - sum * poly);
         += delr;
      if( fabs(delr) < EPS ) break;</pre>
    }
    z[k] = r;
    rlast = r;
```

```
}
return;
}
```

Combining everything that we have done above, we can now implement one function which, when called, returns the zeros and the weights of the Jacobi polynomial of your choice. Notice in this function that we assume that both arrays z and w have already been allocated.

We note that the orthogonality conditions are satisfied automatically by the Legendere polynomials $P_n(x)$. The zeros of these polynomials will then determine the locations of the special points x_j . Some values are shown in table 4.3; these values were computed using the JacobiZW function presented above with both the alpha and beta arguments set to zero.

Also, the fast convergence in the error in Gauss quadrature is due to the fast convergence of the Legendre interpolation. In addition, the computational advantage results from the fact that we employ only half of the (2n) coordinates explicitly. Equidistant interpolation is not a well-convergent process as we have seen in section 3.1.4, and for functions with singularities inside the "oval region" (see section 3.1.4) convergence is not guaranteed. The convergence in Legendre distribution is always guaranteed.

Example: Let us compare the Gauss quadrature with the trapezoid rule for the integral evaluation

$$\int_0^4 x e^x dx = 3e^4 + 1 \cong 164.79445.$$

Using the trapezoid with rule 9-coordinates equidistant we obtain

$$T(h) = (1/2)\left(\frac{1}{2}0 + 0.824361 + \dots + \frac{1}{2}218.3926\right) = 170.42826$$

with error

$$\epsilon^T = 5.63381.$$

Next, we employ Gauss quadrature using 5-coordinates obtained from Legendre's zeros

$$x_k = 2 + 2s_k, \quad s_k \in [-1, 1]$$

which in the domain of interest are

$$2(1 \pm 0.906179846)$$

$$2(1 \pm 0.538469310)$$
.

Table 4.2.2 contains the 5 coordinates in physical space and also the corresponding weights which also need to be corrected, as follows

$$w_k \times \frac{b-a}{2}$$

because

$$\int_{a}^{b} f(x)dx = \frac{b-a}{2} \int_{-1}^{1} f(s)ds.$$

x_k	w_k
0.18764031	0.47385377
0.9230618	0.95725734
2	1.13777778
3.07693862	0.95725734
3.81235969	0.47385377

Table 4.4: Five Gauss-Legendre coordinates and corresponding weights in the interval $x \in [0, 4]$.

The result is:

$$I_G = \sum_{1}^{5} y_k w_k = 1.64794290$$
, with error $\epsilon^G = 1.5981 \times 10^{-4}$,

which is four orders of magnitude less than the result obtained with the trapezoid rule. We, therefore, find that the good interpolation with ninth-order Legendre polynomials leads to good integration with effectively 9 total coordinates.

Gauss Quadrature Error

The error in Gaussian quadrature is relatively difficult to obtain. The standard formula employs the $(2n)^{th}$ derivative (see Lanczos [66]), i.e.,

$$\epsilon^G \sim \left[\frac{(n!)^2}{(2n)!} \right]^2 \frac{2^{2n+1}}{2n+1} \frac{f^{(2n)}(\xi)}{(2n)!},$$
(4.30)

where $\xi \in [-1, 1]$. However, such an error cannot be easily computed, and also this bound is not very sharp as it is based on the evaluation of a very high-order derivative.

An alternative estimate proposed by Lanczos is obtained as follows: Let us start with the identity

$$\int_{-1}^{1} [xf(x)]' dx = \int_{-1}^{1} xf' dx + \int_{-1}^{1} f(x) dx$$
$$= f(1) + f(-1)$$

Then, if we consider the error e^* in the Gauss quadrature of the function (xf') we have that it is equal to the quadrature error of the function (xf)' minus the quadrature error in f because (xf)' = xf' + f. Therefore,

$$\epsilon^* = f(1) + f(-1) - I_G - \sum_{k=1}^n w_k \xi_k f'(\xi_k).$$

In the case of a general interval [a, b] instead of [-1, 1] we have

$$\epsilon^* = \frac{b-a}{2} [f(b) + f(a)] - I_G - \left(\frac{b-a}{2}\right)^2 \sum_{k=1}^n w_k \xi_k f'(x_k).$$

The next key step relies on the assumption that we integrate a function f(x) which is relatively smooth, and thus we can assume that the unknown point is at $\xi \approx 0$. Then the term in the Taylor expansion of the error (equation (4.30)) is approximately equal to the coefficient ξ^{2n} in the Taylor expansion around the origin $\xi = 0$. On the other hand, the expansion of $(\xi f'(\xi))$ is identical to the original expansion except from the shift in coefficients and thus the a_{2n} coefficient is multiplied by (2n+1). By comparison then we can obtain that

$$\epsilon^G pprox rac{1}{2n+1} \epsilon^*,$$

which relates the Gauss quadrature error to the first derivative of the function. If the function f(x) does not meet the smoothness criterion, the above estimate overpredicts the quadrature error. If f'(x) changes sign in the interval [a, b], the above procedure breaks down completely.

Weights and Weighted Moments

Let us consider the grid of figure 4.4.

On this grid, all powers

$$1, x, x^2, \ldots, x^{n-1}$$

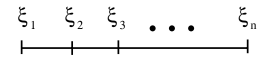


Figure 4.4: Grid to compute the weights.

are interpolated exactly, and thus the corresponding quadrature associated with all these powers will also be exact. Let us compute the k^{th} moment

$$u_k = \int_a^b \rho(x) x^k dx = \sum_{m=1}^n w_m f(\xi_m),$$

where $\rho(x)$ is a weight function. Thus, we can obtain the weights from the known moments, i.e.,

$$w_1 + w_2 + \ldots + w_n = u_0$$

$$w_1\xi_1 + w_2\xi_2 + \ldots + w_n\xi_n = u_1$$

$$w_1 \xi_1^{n-1} + w_2 \xi_2^{n-1} + \dots + w_n \xi_n^{n-1} = u_{n-1}$$

by solving the above system. Alternatively, we can compute the weights directly from

$$w_k = \int_{-1}^1 h_k(x) dx.$$

Theorem: The weights w_k in the Gauss quadrature are positive.

Proof: The formula is exact for $f(x) = h_k^2(x)$, since this is a polynomial of degree 2n. But $h_k(x_j) = \delta_{ij}, \quad k \neq j$. Thus,

$$\int_{a}^{b} (h_k(x)^2 dx = w_k(h_k(x_k))^2 \Rightarrow w_k = \int_{a}^{b} [h_k(x)]^2 dx > 0.$$

Gaussian Quadrature Over Infinite Intervals

The two general strategies in dealing with such important applications of numerical integration are:

- 1. Use knowledge of the integrand to bound the magnitude of the integral from some finite value to infinity by a positive constant, and then use a quadrature formula for the remaining finite interval.
- 2. Use a quadrature formula especially developed for the infinite interval.

Laguerre Integration: Following the second approach here we introduce a weight function w(x)

$$w(x) = e^{-x}, \quad x \in [0, \infty]$$

where we employ the Laguerre polynomial $L_n(x)$ defined by

$$L_0(x) = 1$$

$$L_1(x) = 1 - x$$

$$(n+1)L_{n+1}(x) = -xL_n(x) + (2n+1)L_n(x) - nL_{n-1}(x),$$

and its derivative defined by

$$L'_{n+1}(x) = L'_n(x) - L_n(x).$$

Software

Suite

The corresponding quadrature points are defined by the roots of the Laguerre polynomial (see table 4.2.2 and corresponding software suite). Specifically, the weights are given by

$$w_j = \frac{(n!)^2}{L'_n(x_j)L_{n+1}(x_j)}.$$

Then we approximate the integral

$$\int_0^\infty e^{-x} f(x) = \sum_{j=1}^n w_j f_j + \epsilon^L,$$

and the error is

$$\epsilon^{L} = \frac{(n!)^2}{(2n)!} f^{(2n)}(\xi).$$

Example: Taking n = 3 we can compute the interval

$$\int_0^\infty e^{-x} x^7 dx \cong (0.711093)(0.415774)^7 + (0.278518)(2.294280)^7 + (0.010389)(6.289945)^7 = 4139.9$$

using the values of table 4.2.2. The exact value is 5040, and thus substantial errors occur because $f^{(6)}(x)$ is not bounded. Note, however, that for $n = 4 \Rightarrow f^{(8)}(x) \equiv 0$, and thus we obtain the exact result!



In the *software suite* we present the code necessary to compute the zeros and the weights of the Laguerre polynomials. Four functions are provided as enumerated below:

- 1. The polynomial definition (using recursion) the function Laguerre Poly.
- 2. The derivative definition (using recursion) the function LaguerrePolyDerivative.
- 3. The computation of zeros (using reduction and Newton-Raphson iteration) the function Laguerre Zeros.

n	Abscissas x_j	Weights w_j	n	Abscissas x_j	Weights w_j
2	0.585786	0.853553	4	4.536620	0.038888
	3.414214	0.146447		9.395071	0.000539
3	0.415775	0.711093	5	0.263560	0.521756
	2.294280	0.278518		1.413403	0.398667
	6.289945	0.010389		3.596426	0.074942
4	0.322548	0.603154		7.085810	0.003612
	1.745761	0.357419		12.640801	0.000023

Table 4.5: Zeros of Laguerre polynomials and corresponding weights.

4. Putting it all together to get zeros and weights - the function Laguerre ZW.

Hermite Integration: Here the weight function is

$$w(x) = e^{-x^2},$$

and we can use the Hermite polynomials that are associated with this weight to perform this integration efficiently. We can then compute integrals of the form

$$\int_{-\infty}^{\infty} e^{-x^2} f(x) dx = \sum_{j=1}^{n} w_j f(x_j) + \epsilon^H,$$

with the Hermite polynomial $H_n(x)$ defined by:

$$H_0(x) = 1$$

 $H_1(x) = 2x$
 $H_{n+1}(x) = 2xH_n(x) - 2nH_{n-1}(x),$

and its derivative defined by

$$H'_{n+1}(x) = 2(n+1)H_n(x).$$

The corresponding weights are given by

$$w_j = \frac{2^{n+1} n! \sqrt{\pi}}{\left[H'_n(x_j)\right]^2},$$

and the error is

$$\epsilon^H = \frac{n!\sqrt{\pi}}{2^n(2n)!} f^{2n}(\xi) .$$

n	Abscissas a_j	Weights H_j
2	± 0.707107	0.886227
3	0	1.181636
	± 1.224745	0.295409
4	± 0.524648	0.804914
	± 1.650680	0.081313
5	0	0.945309
	± 0.958572	0.393619
	± 2.020183	0.019953

Table 4.6: Zeros of Hermite polynomials and corresponding weights.

In table 4.2.2 we present the zeros of the Hermite polynomials and corresponding weights for n up to 5.



In the *software suite* we present the code necessary to compute the zeros and the weights of the Hermite polynomials. Four functions are provided as enumerated below:

- 1. The polynomial definition (using recursion) the function *HermitePoly*.
- 2. The derivative definition (using recursion) the function HermitePolyDerivative.
- 3. The computation of zeros (using reduction and Newton-Raphson iteration) the function HermiteZeros.
- 4. Putting it all together to get zeros and weights HermiteZW.

Gauss-Chebyshev Quadrature

The Gauss-Chebyshev quadrature uses the weight function $w(x) = (1-x^2)^{-1/2}$ in the interval $x \in [-1, 1]$, i.e.,

$$\int_{-1}^{1} \frac{f(x)}{\sqrt{1-x^2}} dx = \sum_{k=1}^{n} w_k f_k + \epsilon^C,$$

where the quadrature points are the zeros of the Chebyshev polynomial which are obtained from

$$T_n(x) = \cos(n\cos^{-1}x) = 0$$

and thus

$$x_j = \cos \frac{(2j-1)\pi}{2n}, \ j=1,\dots,n$$

$$w_j = \frac{\pi}{n}.$$

The corresponding quadrature error is

$$\epsilon^C = \frac{2\pi}{2^{2n}(2n)!} f^{(2n)}(\xi).$$

Singular Integrals

We can also follow a weight-function approach to deal with singular integrals, Let us consider integrands with singularities at the end-points of the interval, and write the general problem as

$$\int_{a}^{b} \rho(x)f(x)dx = \sum_{i=1}^{k} w_{k}f(x_{k}) + \epsilon$$

where $\rho(x)$ is a weight function and it may be *singular* at one end-point or both. Accordingly, we distinguish the following cases:

Case 1: $\rho(x) = (1 - x^2)^{1/2}$ on [-1, 1]. Then, we will employ Chebyshev polynomials of the second kind $U_n(x)$ defined as

$$U_n(x) = \frac{\sin[(n+1)\cos^{-1}x]}{\sin(\cos^{-1}x)},$$

which gives the following quadrature points

$$x_j = \cos\frac{j\pi}{n+1}, \quad j = 1, \dots, n$$

with corresponding weights

$$w_j = \frac{\pi}{n+1} \sin^2 \frac{j\pi}{n+1} \,.$$

The error in this quadrature is

$$\epsilon = \frac{\pi}{2^{2n+1}(2n)!} f_{(\xi)}^{(2n)}.$$

Case 2: Here the weight function is $\rho(x) = 1/\sqrt{x}$ on [0,1]. The appropriate polynomial is defined as

$$p_n(x) = P_{2n}(\sqrt{x}),$$

where P_{2n} is the Legendre polynomial. Then the quadrature points are the roots of $p_n(x)$ and the appropriate weights are twice the weights corresponding to $P_{2n}(x)$.

The quadrature error is

$$\epsilon = \frac{2^{4n+1}[(2n)!]^3}{(4n+1)[(4n)!]} f^{(2n)}(\xi).$$

Example: We now return to the integral we mentioned in the introduction of this section, i.e.,

$$\int_0^1 \frac{1+x}{\sqrt{x}} dx$$

and we use n=2. From table 4.3 we obtain the Legendre points

$$x_1 = (0.339981)^2, \quad x_2 = (0.861136)^2$$

and

$$w_1 = 1.304290, \quad w_2 = 0.695710.$$

Therefore,

$$\int_0^1 \frac{1+x}{\sqrt{x}} dx \cong 2.66666,$$

which is equal to the exact value (8/3)!

Case 3: Here the weight function is $\rho(x) = \sqrt{x}$ on [0, 1]. The appropriate polynomial has singularities in the derivative and is defined based on the Legendre polynomial

$$p_n(x) = \frac{1}{\sqrt{x}} P_{2n+1}(\sqrt{x}).$$

The roots x_j of $p_n(x)$ determine the quadrature points and they are then related to the roots of the Legendre polynomial (X_j) by $x_j = X_j^2$. Correspondingly, the appropriate weights are

$$w_j = 2W_j X_j^2 \,,$$

where W_j are the Legendre weights.

The quadrature error is given by

$$\epsilon = \frac{2^{4n+3}[(2n+1)!]^4}{(4n+3)[(4n+2)!](2n)!}f^{(2n)}(\xi).$$

Case 4: Here the weight function is $\rho(x) = [x/(1-x)]^{1/2}$ on [0,1]. The appropriate polynomial is defined in terms of the Chebyshev polynomial, i.e.

$$p_n(x) = \frac{1}{\sqrt{x}} T_{2n+1}(\sqrt{x}).$$

This has a singularity at one point and a derivative-singularity at another point. Here the set of quadrature points, weights, and error, respectively, are given by

$$x_{j} = \cos^{2} \frac{(2j-1)\pi}{4n+2}$$

$$w_{j} = \frac{2\pi}{2n+1} W_{j}$$

$$\epsilon = \frac{\pi}{2^{4n+1}(2n)!} f^{(2n)}(\xi)$$

where W_i refers to the corresponding Chebyshev weights.

Remark: More details on Gauss quadrature based on Jacobi polynomials, which includes both Chebyshev and Legendre polynomials, can be found in Ghizzetti & Ossicini [43], and Karniadakis & Sherwin [63]. In particular, there exist three different approaches in distributing the quadrature points:

- In *Gauss integration* the end-points are not included, and the locations of the quadrature points are determined by the zeros of the Jacobi polynomials.
- In Gauss-Lobatto integration both end-points are included and the interior quadrature points are determined by the zeros of the first derivative of the Jacobi polynomials.
- In Gauss-Radau integration only one point is included and the interior points are determined by the zeros of the Jacobi polynomials with mixed weights.

4.2.3 Multi-Dimensional Integration

Numerical integration in two- or three-dimensions can be accomplished similarly following the algorithms presented above, where *direction splitting* is applied.

We show here how to compute, using Simpson's rule, a two-dimensional integral over a rectangular region as well as a more general region. To this end, we consider the integral

$$Q \equiv \int_{X_1^L}^{X_1^R} \int_{X_2^L}^{X_2^R} f(x_1, x_2) dx_1 dx_2$$

and subdivide the two directions of integration as follows:

$$x_1 = X_1^L + ih_1, i = 0, 1, ..., 2I$$

 $x_2 = X_2^L + jh_2, j = 0, 1, ... 2J$

where

$$h_1 = \frac{X_1^R - X_1^L}{2I}$$
 and $h_2 = \frac{X_2^R - X_2^L}{2J}$.

We apply Simpson's rule direction-by-direction, i.e., we first compute

$$Q_2 \equiv \int_{X_2^L}^{X_2^R} f(x_1, x_2) dx_2$$

SO

$$Q_2 \approx \frac{h_2}{3} \left[f(x_1, x_2^0) + 2 \sum_{j=1}^{J-1} f(x_1, x_2^{2j}) + 4 \sum_{j=1}^{J} f(x_1, x_2^{2j-1}) + f(x_1, x_2^{2J}) \right].$$

Then, the two-dimensional integral is approximated as

$$Q \approx \frac{h_2}{3} \left[\int_{X_1^2}^{X_1^R} f(x_1, x_2^0) dx_1 + 2 \sum_{j=1}^{J-1} \int_{X_1^L}^{X_1^R} f(x_1, x_2^{2j}) dx_1 \right] + 4 \sum_{j=1}^{J} \int_{X_1^L}^{X_1^R} f(x_1, x_2^{2j-1}) dx_1 + \int_{X_1^L}^{X_1^R} f(x_1, x_2^{2J}) dx_1 \right].$$

Next, we need to integrate each one of these terms along the x_1 -direction using again Simpson's rule. This will give

$$\begin{split} Q &\approx \frac{h_1 h_2}{9} \left\{ \left[f(x_1^0, x_2^0) + 2 \sum_{i=1}^{I-1} f(x_1^{2i}, x_2^0) + 4 \sum_{i=1}^{I} f(x_1^{2i-1}, x_2^0) + f(x_1^{2I}, x_2^0) \right] \right. \\ &+ 2 \left[\sum_{j=1}^{J-1} f(x_1^0, x_2^{2j}) + 2 \sum_{j=1}^{J-1} \sum_{i=1}^{I-1} f(x_1^{2i}, x_2^{2j}) + 4 \sum_{j=1}^{J-1} \sum_{i=1}^{I} f(x_1^{2i-j}, x_2^j) \right. \\ &+ \left. \sum_{j=1}^{J-1} f(x_1^{2I}, x_2^{2j}) \right] \\ &+ 4 \left[\sum_{j=1}^{J} f(x_1^0, x_2^{2j-1}) + 2 \sum_{j=1}^{J} \sum_{i=1}^{I-1} f(x_1^{2j}, x_2^{2j-1}) + 4 \sum_{j=1}^{J} \sum_{j=1}^{I} f(x_1^{2i-1}, x_2^{2j-1}) \right. \\ &+ \left. \sum_{j=1}^{J} f(x_1^{2I}, x_2^{2j-1}) \right] \\ &+ \left. \left[f(x_1^0, x_2^{2J}) + 2 \sum_{i=1}^{I-1} f(x_1^{2i}, x_2^{2J}) + 4 \sum_{i=1}^{I} f(x_1^{2i-1}, x_2^{2J}) + f(x_1^{2I}, x_2^{2J}) \right] \right\}. \end{split}$$

The error is additive if the partial fourth-order derivate is continuous along both directions, thus

$$\epsilon^S = -\frac{L_1 L_2}{18} \left[h_1^4 \frac{\partial^4 f(\xi_1, \zeta_1)}{\partial x_1^4} + h_2^4 \frac{\partial^4 f(\xi_2, \zeta_2)}{\partial x_2^4} \right],$$

where (ξ_i, ζ_i) for i = 1, 2 are some unknown points inside the region of integration.

In many applications the region of integration is not rectangular. To this end, we can either use domain decomposing or employ variable step size integration. For example, if we consider the above case with $X_2^L(x_1)$ and $X_2^R(x_1)$ then

$$Q = \int_{\Omega} \int f(x_1, x_2) dx_1 dx_2$$

where Ω is a general, non-cartesian region. In this case, we can proceed as before by allowing the step size

$$h_2(x_1) = \frac{X_2^R(x_1) - X_2^L(x_1)}{2J}.$$

Therefore, for each fixed x_1 location we integrate along x_2 with fixed h_2 step size, as before.

For Gaussian quadrature, we can also form tensor-product interpolations of the integrand as was discussed in section 3.1. This is straightforward for cartesian (orthogonal) domains. For triangular domains the barycentric coordinates (l_1, l_2, l_3) can also be used, and for linear approximations the following exact relations hold:

$$\int_{A} l_{1}^{m} l_{2}^{n} l_{3}^{k} dA = m! n! k! \frac{2A}{(m+n+k+2)!}$$

$$\int_{L_{e}} l_{1}^{m} l_{2}^{n} ds = m! n! \frac{L_{e}}{(m+n+1)!},$$

where A is the area of the triangular region, and L_e is the length of the edge of the triangle.

For integrals in multiple dimensions (greater than 3) or for integrands which are not very smooth it is more efficient to resort to Monte Carlo integration. The convergence of this approach is very slow, e.g. $\mathcal{O}(N^{-1/2})$ compared to $\mathcal{O}(N^{-2})$ for the trapezoid rule or $\mathcal{O}(N^{-4})$ for the Simpson's rule, and thus it is very inefficient for one- two- or three-dimensional integrals. Convergence acceleration is used in practice following standard algorithms such as importance sampling or control variate, see [15] for more details.

4.3 Back to Parallel Computing: Reduction

The concept of numerical integration lends itself to a discussion of *domain decomposition*. Suppose that you want to integrate a function numerically on multiple processors? From our knowledge of calculus, we know that we can write an integral over a interval [a, b] as the sum of the integrals over a disjoint partition of subintervals of [a, b].

For the purposes of this explanation, let us integrate the function $f(x) = \frac{\sin x}{x}$ on the interval [-50, 50]. This is a very important function often used in computer graphics as a filtering function – the Lanczos filter, see section 3.2.

Suppose that we have *five* processors that we can partition this integral operation across. First, we must partition the problem into five processes so that each is doing roughly the same amount of work. This step is graphically accomplished in figure 4.5. Notice that we partition the entire domain [-50, 50] into five equal length subdomains

$$[-50, -30], [-30, -10], [-10, 10], [10, 30], [30, 50].$$

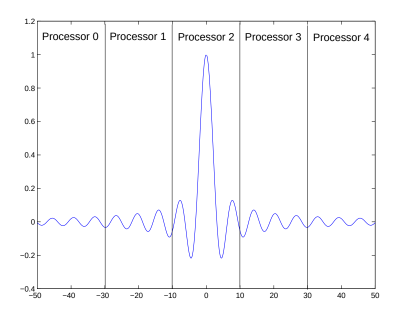


Figure 4.5: Plot of the Lanczos filter $\frac{\sin x}{x}$, and processor assignment.

As we have expressed before, we can now use our numerical integration routines discussed previously in this chapter to accomplish the integration on our local interval. Once this is accomplished, we now want to add up the integral components from each processor. How can this be accomplished? Recall from our previous discussions that the classical way of accomplishing this would be for each process to send its result to one process (process 0), and then for process 0 to accumulate all the results from the individual processes and to print out a result. This type of operation is so common in scientific computing that MPI has a function built in to accomplish all of this in one function call!

Key Concept

• If it seems like a collection of operations is repeatedly done, in all likelihood a function to accomplish those operations already exists to simplify your life.

The function is called *MPI_Reduce* and combines the sending of information with one operation. The admissible operations are given in table 4.3.

Operation Name	Meaning
MPI_MAX	Maximum
MPI_MIN	Minimum
MPI_SUM	Sum
MPI_PROD	Product
MPI_LAND	Logical and
MPI_BAND	Bitwise and
MPI_LOR	Logical or
MPI_BOR	Bitwise or
MPI_LXOR	Logical exclusive or
MPI_BXOR	Bitwise exclusive or
MPI_MAXLOC	Maximum and location of maximum
MPI_MINLOC	Minimum and location of minimum

Table 4.7: Summary of MPI commands for reduction operations.

In our case, we are interested in the MPI_SUM operation; that is, we want to take a piece of information from each processor, and we want the sum of all the pieces to be given to process zero for printing the result. This type of operation is graphically depicted in figure 4.6.

Here, we present the function syntax, argument list description, usage examples, and some remarks for both MPI_Reduce and MPI_Allreduce. The functions are very similar. MPI_Reduce takes information from all processes and sends the result of the MPI operation to only one process. $MPI_Allreduce$ sends the results of the operation back to all processes; it is useful when all processes need the value of the joint operation.

MPI_Reduce:

Function Call Syntax

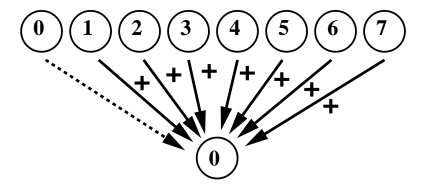


Figure 4.6: Reduction (MPI_Reduce) combines fanning-in within a single operation.

```
int MPI_Reduce(
        void*
                        operand
        void*
                        result
                                       out
        int
                        count
        MPI_Datatype
                        datatype
                                      in
        MPI_Op
                        operator
        int
                        root
        MPI_Comm
                        comm
```

Understanding the Argument List

- operand starting address of the send buffer.
- result starting address of the receive buffer.
- count number of elements in the send buffer.
- datatype data type of the elements in the send/receive buffer.
- operator reduction operation to be executed.
- root rank of the root process obtaining the result.
- *comm* communicator.

Example of Usage

```
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

// Determine datasize and root

double * senddata = new double[datasize];
double * recvdata = NULL;
if(mynode == root)
    recvdata = new double[datasize];

// Fill in senddata on all processes

MPI_Reduce(senddata,recvdata,datasize,MPI_DOUBLE,MPI_SUM, root,MPI_COMM_WORLD);

// At this stage, the process root contains the result
// of the reduction (in this case MPI_SUM) in the
// recvdata array
```

Remarks

- The *recvdata* array only needs to be allocated on the process of rank root (since root is the only processor receiving data). All other processes may pass NULL in the place of the *recvdata* argument.
- Both the *senddata* array and the *recvdata* array must be of the same data type. Both arrays should contain at least *datasize* elements.

MPI_Allreduce:

Function Call Syntax

```
int MPI_Allreduce(
        void*
                        operand
        *biov
                        result
                                      out
                        count
                                      in
        int
        MPI_Datatype
                       datatype
                                      in
        MPI_Op
                        operator
                                      in
        MPI_Comm
                        comm
```

Understanding the Argument List

- operand starting address of the send buffer.
- result starting address of the receive buffer.

- count number of elements in the send/receive buffer.
- datatype data type of the elements in the send/receive buffer.
- operator reduction operation to be executed.
- comm communicator.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units over which
              //
                     reduction should occur
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
// Determine datasize and root
double * senddata = new double[datasize];
double * recvdata = new double[datasize];
// Fill in senddata on all processes
MPI_Allreduce(senddata, recvdata, datasize, MPI_DOUBLE,
              MPI_SUM,MPI_COMM_WORLD);
// At this stage, all processes contains the result
// of the reduction (in this case MPI_SUM) in the
// recvdata array
```

Remarks

- In this case, the *recvdata* array needs to be allocated on all processes since all processes will be receiving the result of the reduction.
- Both the *senddata* array and the *recvdata* array must be of the same data type. Both arrays should contain at least *datasize* elements.



We present a sample MPI code which accomplishes the integration of $f(x) = \frac{\sin x}{x}$ on the interval [-50, 50] across multiple processes, the number of which is specified by the user, i.e., totalnodes in the example code below.

```
#include<iostream.h>
#include<math.h>
#include<mpi.h>
#include "SCchapter4.h"
double func(double x);
int main(int argc, char * argv[]){
  int mynode, totalnodes;
  const double global_a = -50.0;
  const double global_b = 50.0;
  const int levels = 10;
  double local_a,local_b,local_sum,answer;
  MPI_Init(&argc,&argv);
  MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
  MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
  local_a = global_a + mynode
                                 *(global_b-global_a)/totalnodes;
  local_b = global_a + (mynode+1)*(global_b-global_a)/totalnodes;
  local_sum = MidpointRule(levels, local_a, local_b, func);
  MPI_Reduce(&local_sum,&answer,1,MPI_DOUBLE,
             MPI_SUM,O,MPI_COMM_WORLD);
  if(mynode == 0){
    cout << "The value of the integral is: " << answer << endl;</pre>
 MPI_Finalize();
}
double func(double x){
  return(sin(x)/x);
}
```

We would like to draw your attention to the following points:

• Observe that every process has the same global_a and global_b variable, here defined as a constant (by using the keyword const in the declaration of the variable), however every process must maintain its own version of local_a and local_b. This is a very important concept to understand: There exists a global domain, which in this case is the interval

[-50, 50], and each processor must keep track of its "computational responsibility" on this domain.

• Notice the use of the operator "& ." In this case, you can translate "&" to mean "the address of." Hence &local_sum is equivalent to saying "the address of local_sum." Why do we do this? Recall that for this particular argument position, MPI expects us to provide to it a pointer variable (which, if you recall, is an address variable). However, we did not allocate an array, we merely had one value. Instead of creating an array which contained only one element, copying the value of local_sum into that array element, and then passing the pointer variable to MPI_Reduce, we used a neat programming feature of C++. Recall from section 3.1.2 the programming feature "&," which is an operator which allows you to obtain the address of a variable (and hence you can "pass it off" as a pointer). In the example above, you should be mindful that

is invalid, because MPI_Reduce will expect pointer (address) variables in the argument positions for $local_sum$ and answer, and hence you must use:

• Each process puts the result of its local integration in the first argument of MPI_Reduce, and immediately following this call on process zero (because in the sixth argument we have placed a "0", denoting that process zero is to be the recipient of the reduction) the variable answer contains the sum of all the information contained in the local_sum variables across all the processes. If we were to do an MPI_Allreduce, then we would not specify which processor the reduction was going to (because all processes would obtain the result of the reduction), and on each process their local copy of the variable "answer" would contain the sum of all local_sum variables.

4.4 Homework Problems

4.4.1 Homework Problems for Section 4.1

1. Consider the iteration

$$x_{n+1} = ax_n - ax_n^2.$$

Plot the converged solution for different values of a, i.e., a=1,2,3,3.3,3.5,3.7,3.8, etc.

What do you observe?

- 2. Write a C++ program to compute the first dozen roots of $\tan x = x$.
- 3. Modify the function NewtonRaphson given in the text so that it does not accomplish unnecessary re-evaluations of the function (per the remarks made in the text).
- 4. Write a function similar to *NewtonRaphson* which implements the secant method. Your function should allow the user to input an arbitrary function (just as the *NewtonRaphson* function does).
- 5. Find all the intersection points of the function

$$f(x) = e^x$$
 and $g(x) = 2x^2$.

What is the convergence rate that you observed?

6. Write a C++ code to find the intersection of the circle with the ellipse

$$x^2 + y^2 = 1$$

$$\frac{x^2}{4} + 16y^2 = 1$$

to within double precision ($\epsilon = 10^{-14}$).

Prove that we can compute \sqrt{C} with cubic rate of convergence from the formula:

$$x_{n+1} = x_n \frac{x_n^2 + 3C}{3x_n^2 + C}.$$

7. Let us replace the equation $\mathbf{A}\mathbf{x} = \mathbf{0}$ (which has the trivial solution $\mathbf{x} = 0$) with the iteration

$$\mathbf{x}_{n+1} - \mathbf{x}_n = -\mathbf{A}\mathbf{x}_n$$
 where \mathbf{x}_0 is arbitrary.

- (a) What is the appropriate test on the matrix **A** to check convergence?
- (b) We want to apply the steepest descent method to minimize $P = x_1^2 x_1x_2 + x_2^2$ and we set the step-size $\alpha = 1$. First, obtain an expression for the vector \mathbf{x}_{n+1} . Do the iterations converge to the bottom point of P (at x = 0)? Sketch the quadratic $P(x_1, x_2)$ and follow the first iteration graphically starting from an initial vector $\mathbf{x}_0 = (1, 1)$ and $\mathbf{x}_0 = (1, -1)$.

8. Use the method of steepest descent to obtain the solutions of the following systems to within tolerance

(a)
$$\epsilon = 10^{-2}$$
 and (b) $\epsilon = 10^{-5}$.

How does the number of iterations scale with the tolerance level?

(i)

$$x_1^2 + x_2^2 = 2$$
$$-\cosh x_1 + x_2 = 0$$

(ii)

$$x_3 = e^{x_1} + e^{x_2}$$

$$x_2^2 = 4 + 2x_1x_3$$

$$x_1x_3 = x_1^3 + x_1^2x^2 + 6$$

Can you solve the above using Newton's method? What is the relative gain in iteration number versus computational work?

9. Halley's method: Show that the iteration

$$x_{n+1} = x_n - \frac{f_n f_n'}{(f_n')^2 - (f_n f_n^*)/2}$$

finds the roots of f(x) = 0 and estimate the convergence rate. Here $f_n = f(x_n)$.

Hint: Set $\phi(x) \equiv f/\sqrt{f'}$ and use the Newton-Raphson formula for $\phi(x)$.

10. Mueller's method: In this method a quadratic approximation of the given function is assumed given three pairs of data points: (x_1, y_1) ; (x_2, y_2) ; (x_3, y_3) , as follows

$$f(x) \approx y_3 + y'(x - x_3) + y''(x - x_3)(x - x_2)$$

where

$$y' = \frac{y_3 - y_2}{x_3 - x_2}, \quad y'_1 = \frac{y_2 - y_1}{x_2 - x_1}.$$

and

$$y'' = \frac{y' - y'}{x_3 - x_1}$$
 $z = y' = y''(x_3 - x_2)$

Show that we can solve approximately for the root

$$s = x_3 - \frac{2y_3}{z + \operatorname{sign}(z)\sqrt{z^2 - 4y_3y''}}.$$

In the next update we set

$$x_1 = x_2$$
; $x_2 = x_3$; $x_3 = s$,

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and so on, until convergence to specified tolerance.

The rate of convergence of Mueller's method is $m \approx 1.84$ and requires only function evaluations, just like the secant method. It can also be used to find both complex and real roots.

Practice this method by solving the equation

$$f(x) = 2x^{10} - 1 = 0.$$

- 11. The natural frequencies of the vibrations of a beam structure depend on the boundary conditions, i.e., the type of support of the structure.
 - (a) If the beam is pinned at one end and free at the other end, the eigenfrequencies are solutions of the equation

$$\tan \omega - \tanh \omega = 0.$$

(b) If the beam is clamped at one end and free at the other end, the equation is

$$\tan \omega - \tanh \omega + 1 = 0.$$

Here ω is a non-dimensional frequency that depends on the length of the beam, its density, and its flexural rigidity.

Obtain the eigenfrequencies for both cases for *single* and *double* precision and compare the corresponding computational work for each case.

12. Consider the iteration

$$y_i = x_i - \frac{f(x_i)}{f'(x_i)}; \quad x_{i+1} = y_i - \frac{f(y_i)}{f'(x_i)}.$$

This is the Newton-Raphson iteration with the derivative computed only every second step.

(a) Show that if the iteration converges,

$$\lim_{i \to \infty} \frac{x_{i+1} - \alpha}{(y_i - \alpha)(x_i - \alpha)} = \frac{f''(\alpha)}{f'(\alpha)} \text{ as } x_i \to \alpha.$$

(b) Thus conclude that

$$\lim_{i \to \infty} \frac{x_{i+1} - \alpha}{(x_i - \alpha)^3} = \frac{1}{2} \left[\frac{f''(\alpha)}{f'(\alpha)} \right]^2.$$

(c) If the cost of computing f(x) is 1 and f'(x) is C, for what values of C is the method more efficient than (i) the Newton-Raphson method (ii) the secant method?

13. Write a C++ program to calculate the wavenumber corresponding to a given angular frequency for gravity water waves in finite depth, including surface tension effects. Plot κ versus ω for various depths, assuming that for water

$$\tau = 70mN/m, \ \rho = 1000kg/m^3, \ g = 9.81m/s^2.$$

The dispersion relation for finite depth, including surface tension is:

$$(g\kappa + \frac{\tau}{\rho}\kappa^3)tanh(\kappa h) = \omega^2.$$

- 14. Write an MPI code to accomplish a parallel conjugate gradient method. Assume that the matrix is distributed by rows across each process.
 - (a) Given that the matrix is partitioned by rows across each process, outline the parallel decomposition of the conjugate gradient algorithm. What operations can be done concurrently, and when will parallel calls be required?
 - (b) Determining how many calls to $MPI_Reduce/MPI_Allreduce$ are necessary. What operations within the CG algorithm require reduction?
 - (c) Where will MPI_Send/MPI_Recv be required? Why?
 - (d) Define $h = \frac{1}{N}$, $\mu = h^2$, and $q[i][j] = (8\pi^2 + 1)h^2 sin(2\pi hi)sin(2\pi hj)$ where $i, j = 0, \ldots, N-1$.

Let **A** be of the form given in figure 4.7.

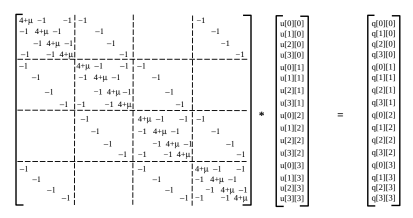


Figure 4.7: Matrix system $\mathbf{A}u = q$ for a N = 4.

Solve the matrix system $\mathbf{A}\mathbf{u} = \mathbf{q}$ for u for both N = 4 and N = 20. Notice that the definition above uses two-dimensional arrays, which we will go over in the next chapter. Implement the concept of a two-dimensional array using only a one-dimensional array construction. What indexing is required?

(e) When you run the program on different numbers of processors, is there a noticeable difference in the performance time?

4.4.2 Homework Problems for Section 4.2

1. Ramanujan proposed that the number of numbers between a and b which are either squares or sums of two squares is given approximately by the integral

$$0.764 \int_a^b \frac{dx}{\sqrt{loq_e x}}$$

Use Gauss quadrature to test numerically this hypothesis for a = 1; b = 30.

2. Evaluate the sine integral Si(x)

$$Si(x) = \int_0^x \frac{\sin z}{z} dz$$

using Simpson, Romberg and the Gauss method for x = 2. What do you observe?

3. Use Simpson's formula to find the length of the ellipse

$$\frac{x^2}{a^2} + \frac{y^2}{b^2} = 1.$$

Plot the error versus the number of quadrature points, and verify the fourth-order convergence of the method.

Hint: Describe the ellipse parametrically using

$$x(\theta) = a \cos \theta$$

 $y(\theta) = b \sin \theta, \theta \in [0, 2\pi]$

and obtain the length using the formula

$$\int \sqrt{dx^2 + dy^2} \,.$$

4. Compute the error function

$$Erf(x) = \frac{2}{\sqrt{\pi}} \int_0^x e^{-t^2} dt$$

for x = 1/2, 1, 2, and 4 using trapezoid rule, and using the appropriate Gauss quadrature for n = 6 and 12 quadrature points. Using the formula for the error bounds, obtain an estimate of errors for the two cases.

5. Use a four-level Romberg integration to compute the integral

$$I = \int_{1}^{5} \frac{1}{x^2} dx$$
.

How many quadrature points are required to achieve accuracy of 10^{-6} ?

6. Use Simpson's rule and Gaussian quadrature to compute the multi-dimensional integrals

(a)
$$\int_0^\pi \int_0^\pi (y^2 \sin x + x^2 \cos^2 y) dx dy$$

with n = 4 quadrature points in each direction.

7. Compute the integrals

$$\int_{-\infty}^{\infty} \frac{dx}{1+x^2}$$

(b)
$$\int_0^\infty e^{-x} \sin x dx$$

with n=5 quadrature points, and compute the quadrature error in each case.

Chapter 5

Explicit Discretizations

In this chapter we consider explicit discretizations of space- and time-derivatives. In such discretizations we can express directly a derivative at one grid point in terms of function values at adjacent grid points (spatial discretizations) or in terms of previous time levels (temporal discretizations). This, in turn, implies that there is no implicit coupling, and thus there is no matrix inversion involved but instead simple daxpy type operations.

The material in this chapter is relatively easy to program both on serial as well as on parallel computers. It is appropriate for demonstrating fundamental concepts of discretization as well as primary constructs of the C++ language and of the MPI library. Specifically, we will demonstrate the use of loops, arrays, functions and passing functions to functions. In addition to presenting MPI_Send and MPI_Recv implementations for finite differences, we also introduce $MPI_Sendrecv$ and $MPI_Sendrecv_replace$ as alternative advanced MPI function calls for parallelizing finite differences discretizations.

5.1 Explicit Space Discretizations

5.1.1 Basics

The formulation of derivatives based on function values on a set of points, which we call the *grid*, dates back to Euler in the beginning of the eighteen century. However, advances have been made since then! In this section, we will formulate ways to compute first and higher order derivatives of a function using discrete data points. The key idea is to use Taylor expansions at a subset of adjacent points of the grid as shown in figure 5.1.

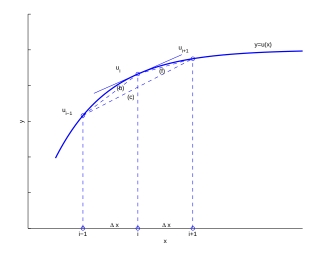


Figure 5.1: A stencil of two or three points is required to define the discrete first derivative: (c) corresponds to central difference; (b) corresponds to backward difference; and (f) corresponds to forward difference.

Assuming that we have an equidistant grid, i.e., the distance $\Delta x \equiv x_{i+1} - x_i$ is constant for every grid point (i), then using Taylor's expansion for the smooth function u(x) around the point (i), we obtain

$$u_{i\pm 1} = u(x \pm \Delta x) = u(x) \pm \Delta x u_x(x) + \frac{\Delta x^2}{2} u_{xx}(x) \pm \dots$$

Here the x subscript denotes differentiation and the i subscript refers to the index of data points; also Δx^2 means $(\Delta x)^2$.

We can now form three types of differences from the above expansions:

•forward:
$$(u_x)_i = \frac{u_{i+1} - u_i}{\Delta x} + \underbrace{\mathcal{O}(\Delta x)}_{-1/2(u_{xx})_i \Delta x}$$

•backward: $(u_x)_i = \frac{u_i - u_{i-1}}{\Delta x} + \underbrace{\mathcal{O}(\Delta x)}_{+1/2(u_{xx})_i \Delta x}$ (upwind •central: $(u_x)_i = \frac{u_{i+1} - u_{i-1}}{2\Delta x} + \underbrace{\mathcal{O}(\Delta x^2)}_{1/6(v_i) \Delta x^2}$

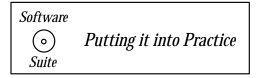
The accuracy of each finite difference approximation depends on the last term. Therefore, the forward and backward differences are first-order, i.e., $\mathcal{O}(\Delta x)$, while the central difference is of second-order, i.e. $\mathcal{O}(\Delta x^2)$. We also note that the backward and forward differences are actually of order $\mathcal{O}(\Delta x^2)$ approximations to derivatives at the half-point $(i \pm 1/2)$, respectively, e.g.,

$$(u_x)_{i+1/2} = \frac{u_{i+1} - u_i}{\Delta x} + \mathcal{O}(\Delta x^2).$$

To obtain a second-order accurate formula for the second derivative we simply add the two Taylor expansions formed at the $(i \pm 1)$ points

$$(u_{xx})_i \approx \frac{u_{i+1} - 2u_i + u_{i-1}}{\Delta x^2} - \frac{1}{12} (u_{xxxx})_i \Delta x^2.$$

The basic subset of grid points, the *stencil*, consists of two or three adjacent points for these discrete derivatives. From the parallel computing standpoint, this implies that *local* computations are involved. In particular, these are **explicit** formulas, in that the derivative at a point is computed in terms of function values at adjacent points in the same stencil. Therefore, there are no other derivatives involved in the discretization besides the derivative at point *i*. In chapter 6, we will see **implicit** formulas in which other derivatives are involved in the discretization.



Explicit formulas are also easy to program. Here we demonstrate with the next C++ example how *first derivatives* can be approximated for both non-periodic and periodic intervals.

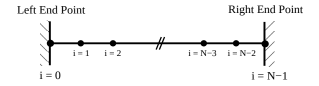


Figure 5.2: Interior and end (boundary) points in an interval.

In this first example, we demonstrate how finite differencing can be done for a non-periodic interval. Notice that we cannot use the central difference formula to calculate an approximation of the derivatives at the end-points because we have no information from outside the interval! Hence, as in this case, we use what are called "one-sided" approximations, which use information only from inside the interval. Observe that this requires us to break the approximation process into three stages:

- 1. Compute the approximation for the interior points (i = 1, ..., N 2).
- 2. Compute the approximation for the left end-point (i = 0).
- 3. Compute the approximation for the right end-point (i = N 1).

For *periodic* boundary conditions, we can use the central approximation for the endpoints; however, care must be taken to properly index the array. Notice that we cannot access the array at the value -1, and hence we must explicitly treat the end-points so that the appropriate values are taken. Once again, we break the computation into three stages:

- 1. Compute the approximation for the interior points (i = 1, ..., N 2).
- 2. Compute the approximation for the left end-point (i = 0).
- 3. Compute the approximation for the right end-point (i = N 1).

Let us point out, at this stage, a small optimization hint that may be of use in the future. Notice in the examples above that at the beginning of the function we explicitly calculate the value of the variable two_invdx , and then continue to use this variable throughout the function. We do this because we observed that the value of dx is not changing inside any of the loops within this function, and hence computing 2.0*dx needs to be done only once. Hence, we can do it up front (at the beginning of the function) instead of repeatedly doing it within the loop. We have chosen to go ahead and do the division because on many machines division is more expensive than multiplication, and then multiplying within the loop instead of dividing is the most efficient choice.

Key Concept

• If something is not dependent on the looping variable, then do not do it in the loop!

5.1.2 Uniform Grids

In the following we show how we can obtain systematically finite difference formulas on one-dimensional grids, assuming that the spacing between the grid points is constant. The objective is to develop formulas for high order approximations for interior points as well as for boundary points.

I. Method of Undetermined Coefficients

The key idea in this method is to consider the Taylor expansions as functions of Δx and obtain equations for the weights (unknown coefficients) by equating the coefficients of powers of Δx . We demonstrate this by an example.

Example: We want to obtain a one-sided, second-order finite difference for $(u_x)_i$, that is

$$(u_x)_i = \frac{au_i + bu_{i-1} + cu_{i-2}}{\Delta x} + \mathcal{O}(\Delta x^2)$$

as shown in the sketch of figure 5.3 in order to handle a boundary condition on the right end of the computational domain. One approach is to employ phantom or ghost nodes outside the domain and construct central difference schemes but this would require extra information about the physics of the problem. The more general approach is to construct one-sided differences extracting information from the interior of the domain.

As before we expand the function at the different points around the point of interest, to

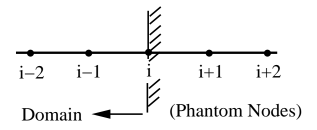


Figure 5.3: Grid at the right boundary of the domain.

obtain:

$$c: u_{i-2} = u_i - 2\Delta x (u_x)_i + 2\Delta x^2 (u_{xx})_i - \frac{(2\Delta x)^3}{6} (u_{xxx})_i + \dots$$

$$b: u_{i-1} = u_i - \Delta x (u_x)_i + \frac{\Delta x^2}{2} (u_{xx})_i - \frac{\Delta x^3}{6} (u_{xxx})_i + \dots$$

$$a: u_i = u_i$$

Upon substitution in the assumed expression for $(u_x)_i$ we obtain

$$(\Delta x)(u_x)_i = au_i + bu_{i-1} + cu_{i-2} + \mathcal{O}(\Delta x^3)$$

= $(a+b+c)u_i - \Delta x(2c+b)(u_x)_i + \frac{\Delta x^2}{2}(4c+b)(u_{xx})_i + \mathcal{O}(\Delta x^3)$

Therefore, solving a 3×3 system we find the values for the coefficients

$$\left\{ \begin{array}{l} a+b+c=0 \\ 2c+b=-1 \\ 4c+b=0 \end{array} \right\} \Rightarrow a=3/2, \ b=-2, \ c=1/2,$$

and the final formula is

$$(u_x)_i = \frac{3u_i - 4u_{i-1} + u_{i-2}}{2\Delta x} + \mathcal{O}(\Delta x^2).$$

This is a second-order *upwind difference* for the first derivative with truncation error $T \approx \frac{\Delta x^2}{3}$.

In general, a first-order derivative at mesh point (i) can be made of order of accuracy (p) by an explicit formula involving (p+1) points. However, this is not true for implicit discretizations (see next chapter), where on a two-point stencil we can obtain second-order accuracy for the first derivative.

II. Difference Operators

The method of difference operators is based on symbolic manipulation. We first define the operators which will be acting on functions assigned to grid points, as follows:

Displacement: $E^n u_i \equiv u_{i+n}$

Forward: $\delta^+ u_i \equiv u_{i+1} - u_i \Rightarrow \delta^+ = E - 1$

Backward: $\delta^- u_i \equiv u_i - u_{i-1} \Rightarrow \delta^- = 1 - E^{-1}$

Half-central: $\delta u_i \equiv u_{i+1/2} - u_{i-1/2} \Rightarrow \delta = E^{1/2} - E^{-1/2}$

Central: $\bar{\delta}u_i \equiv 1/2(u_{i+1} - u_{i-1}) \Rightarrow \bar{\delta} = 1/2(E - E^{-1})$

Average: $\mu u_i \equiv 1/2(u_{i+1/2} + u_{i-1/2}) \Rightarrow \mu = 1/2(E^{1/2} + E^{-1/2})$

Derivative: $Du \equiv \partial u/\partial x$

The symbolic manipulations are straightforward, for example

$$\delta^{+2} = \delta^{+} \delta^{+} = (E - 1)(E - 1) = E^{2} - 2E + 1,$$

and similarly, $\mu\delta = \bar{\delta}$, etc. We can also write the Taylor expansion in operator form by considering the expansion

$$u(x + \Delta x) = u(x) + \Delta x u_x + \frac{\Delta x^2}{2!} u_{xx} + \frac{\Delta x^3}{3!} u_{xxx} + \dots$$

The corresponding operator form is

$$Eu(x) = \left[1 + \Delta x D + \frac{(\Delta x D)^2}{2!} + \frac{(\Delta x D)^3}{3!} + \dots\right] u(x)$$

or

$$Eu(x) = e^{\Delta x D} u(x) \Rightarrow E = e^{\Delta x D} \Rightarrow \boxed{\Delta x D = \ln(E)}.$$
 (5.1)

This last equation is very useful and will be used repeatedly in the following.

• Forward Differences: Starting from equation (5.1) we have

$$\Delta xD = \ln(E) = \ln(1 + \delta^{+}) \text{ because } (\delta^{+} = E - 1)$$

= $\delta^{+} - \frac{\delta^{+2}}{2} + \frac{\delta^{+3}}{3} - \frac{\delta^{+4}}{4} + \dots,$

where the first neglected term gives the truncation error. For example, if we keep two terms:

$$\Delta x(Du)_i = \delta^+ u_i - \frac{\delta^{+2}}{2} u_i$$

$$= (u_{i+1} - u_i) - \frac{1}{2} (E^2 - 2E + 1) u_i$$

$$= (u_{i+1} - u_i) - \frac{1}{2} (u_{i+2} - 2u_{i+1} + u_i)$$

we obtain a one-sided, second-order formula, similar to the one we obtained before using the method of undetermined coefficients:

$$Du_i = (u_x)_i = \frac{-3u_i + 4u_{i+1} - u_{i+2}}{2\Delta x} + \underbrace{\frac{\Delta x^2}{3}(u_{xxx})_i}_{\text{truncation error}}$$

Here the coefficients have the opposite sign compared to the formula before because the above formula is for the left boundary (downwind difference).

• Backward Differences: Similarly, we can obtain formulas for backward differences starting from

$$\Delta xD = \ln(E) = -\ln(1 - \delta^{-})$$

or

$$\ln E = \delta^{-} + \frac{\delta^{-2}}{2} + \frac{\delta^{-3}}{3} + \frac{\delta^{-4}}{4} + \dots$$

• Central Differences: Here we can use either the half-central or the central operators to derive appropriate formulas. To this end,

$$\delta u_i = u_{i+1/2} - u_{i-1/2} = (E^{1/2} - E^{-1/2})u_i$$

$$\Rightarrow \delta = e^{\Delta x D/2} - e^{-\Delta x D/2} = 2\sinh\frac{\Delta x D}{2}$$

$$\Rightarrow \Delta x D = 2\sinh^{-1}\delta/2 = 2\left[\delta/2 - \frac{1}{2\cdot 3}\left(\frac{\delta}{2}\right)^3 + \frac{1\cdot 3}{2\cdot 4\cdot 5}\left(\delta/2\right)^5 - \frac{1\cdot 3\cdot 5}{2\cdot 4\cdot 6\cdot 7}\left(\delta/2\right)^7 + \ldots\right].$$

Therefore, we obtain again

$$\Delta xD = \delta - \frac{\delta^3}{24} + \frac{3\delta^5}{640} - \frac{5\delta^7}{7168} + \dots$$

If we only keep the first term, we derive a second-order formula

$$Du_i = \frac{u_{i+1/2} - u_{i-1/2}}{\Delta x} - \frac{\Delta x^2}{24} (u_{xxx})_i + \dots,$$

however, now we need values at Δx half-integer grid points. This will be suitable for staggered grids where both half-integer and integer grid points are employed.

In order to involve function values at the integer grid points we employ the central operator

$$\bar{\delta} = \frac{1}{2}(E - E^{-1}) = \frac{1}{2}(e^{\Delta x D x} - e^{-\Delta x D}) = \sinh(\Delta x D)$$

$$\Rightarrow \Delta x D = \sinh^{-1} \bar{\delta}$$

$$= (\bar{\delta} - \frac{\bar{\delta}^3}{6} + \frac{3}{2 \cdot 4 \cdot 5} \bar{\delta}^5 + \dots).$$

To achieve second-order accuracy we keep the first term only, i.e.,

$$Du_i = \bar{\delta}/\Delta x = \frac{u_{i+1} - u_{i-1}}{2\Delta x} - \frac{\Delta x^2}{6}(u_{xxx})_i + \dots$$

$$i-2$$
 $i-1$ i $i+1$ $i+2...$

Figure 5.4: A five-point stencil for explicit discretization of the first derivative with fourth-order accuracy.

To achieve fourth-order we need to also keep the term $\bar{\delta}^3$, but this will lead to a relatively long stencil, that is a seven-point stencil because of the index $(i \pm 3)$. In general, a fourth-order accuracy for approximating the first derivative should not require more than a five-point stencil, as shown in figure 5.4. In order to achieve this we require to do some extra work by involving the average operator:

$$\mu^{2} = 1 + \delta^{2}/4 \Rightarrow \mu(1 + \delta^{2}/4)^{-1/2} = 1$$
$$\Rightarrow \mu(1 - \frac{\delta^{2}}{8} + \frac{3\delta^{4}}{128} - \frac{5\delta^{6}}{1024} + \ldots) = 1$$

However, we have already presented the expansion for the half-central operator above, and using it we obtain after multiplying both sides by unity

$$1 \times [\Delta x D] = \left[\delta - \delta^3 / 24 + \frac{3\delta^5}{640} - \frac{5\delta^7}{7168} \right] \times 1$$
$$= \mu (\delta - \frac{1}{3!} \delta^3 + \frac{1^2 2^2}{5!} \delta^5 - \dots)$$
$$= \bar{\delta} (1 - \frac{\delta^2}{3!} + \frac{2^2}{5!} \delta^4 - \frac{2^2 3^2}{7!} \delta^6 + \dots).$$

Therefore, the fourth-order formula for the first derivative on a five-point stencil, see figure 5.4, is

$$(u_x)_i = \frac{-u_{i+2} + 8u_{i+1} - 8u_{i-1} + u_{i-2}}{12\Delta x} + \frac{\Delta x^4}{30} \left(\frac{\partial^5 u}{\partial x^5}\right).$$

• **Higher-Order Derivatives:** To compute higher derivatives we proceed in a similar way except we need to expand symbolically binomials as shown below:

Forward:

$$\left(\frac{\partial^n u}{\partial x^n}\right)_i = D^n u_i = \frac{1}{\Delta x^n} [\ln(1+\delta^+)]^n u_i$$

$$= \frac{1}{\Delta x^n} \left[\delta^{+n} - \frac{n}{2} \delta^{+(n+1)} + \frac{n(3n+5)}{24} \delta^{+(n+2)} - \frac{n(n+2)(n+3)}{48} \delta^{+(n+3)} + \dots \right] u_i$$
 (5.2)

Backward:

$$\left(\frac{\partial^{n} u}{\partial x^{n}}\right)_{i} = -\frac{1}{\Delta x^{n}} \left[\ln(1-\delta^{-})\right]^{n} u_{i}$$

$$= \frac{1}{\Delta x^{n}} \left(\delta^{-} + \frac{\delta^{-2}}{2} + \frac{\delta^{-3}}{3} + \dots\right)^{n} u_{i}$$

$$= \frac{1}{\Delta x^{n}} \left[\delta^{-n} + \frac{n}{2} \delta^{-(n+1)} + \frac{n(3n+5)}{24} \delta^{-(n+2)} + \frac{n(n+2)(n+3)}{48} + \delta^{-(n+3)} + \dots\right] u_{i} \tag{5.4}$$

Central:

$$D^{n}u_{i} = \left(\frac{2}{\Delta x}\sinh^{-1}\frac{\delta}{2}\right)^{n}u_{i} = \frac{1}{\Delta x^{n}}\left[\delta - \frac{\delta^{3}}{24} + \frac{3\delta^{5}}{640} - \frac{5\delta^{7}}{7168} + \ldots\right]^{n}u_{i}$$

$$= \frac{1}{\Delta x^{n}}\delta^{n}\left[1 - \frac{n}{24}\delta^{2} + \frac{n}{64}\left(\frac{22 + 5n}{90}\right)\delta^{4} - \frac{n}{45}\left(\frac{5}{7} + \frac{n - 1}{5} + \frac{(n - 1)(n - 2)}{35}\right)\delta^{6} + \ldots\right]u_{i}$$
(5.5)

Remark: For *n* even, function values at integer grid points are required.

Second Derivative: The formulas for the second derivative are simplified as follows

• Forward:
$$(u_{xx})_i = \frac{1}{\Delta x^2} \left(\delta^{+2} - \delta^{+3} + \frac{11}{12} \delta^{+4} - \frac{5}{6} \delta^{+5} + \ldots \right) u_i$$

• Backward:
$$(u_{xx})_i = \frac{1}{\Delta x^2} \left(\delta^{-2} + \delta^{-3} + \frac{11}{12} \delta^{-4} + \frac{5}{6} \delta^{-5} + \ldots \right) u_i$$

• Central:
$$(u_{xx})_i = \frac{1}{\Delta x^2} \left(\delta^2 - \frac{\delta^4}{12} + \frac{\delta^6}{90} - \frac{\delta^8}{560} + \ldots \right) u_i$$

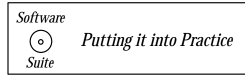
Note that because of symmetry, the central discretization leads to higher-order accuracy by maintaining only the first term.

Example: We can derive useful formulas by keeping only two terms in the expansions, as follows:

• Forward:
$$(u_{xx})_i = \frac{1}{\Delta x^2} (2u_i - 5u_{i+1} + 4u_{i+2} - u_{i+3}) + \frac{11}{12} \Delta x^2 \left(\frac{\partial^4 u}{\partial x^4} \right)$$

- Backward: $(u_{xx})_i = \frac{1}{\Delta x^2} (2u_i 5u_{i-1} + 4u_{i-2} u_{i-3}) \frac{11}{12} \Delta x^2 \left(\frac{\partial^4 u}{\partial x^4} \right)$
- Central: $(u_{xx})_i = \frac{1}{12\Delta x^2}(-u_{i+2} + 16u_{i+1} 30u_i + 16u_{i-1} u_{i-2}) + \frac{\Delta x^4}{90}\left(\frac{\partial^6 u}{\partial x^6}\right)$

The above two one-sided formulas are important in handling boundary points. In the code example below, we implement these formulas for the left boundary, the right boundary, and the interior respectively, assuming the index i runs from left to right.



Just as with the first derivative approximations, we break the approximation process into three stages:

- 1. Compute the approximation for the interior points (i = 1, ..., N 2).
- 2. Compute the approximation for the left end-point (i = 0).
- 3. Compute the approximation for the right end-point (i = N 1).

Here we present the second derivative finite difference approximation for a *non-periodic* interval. Observe that at the end-points we use the four-point one-sided approximation derived above.

As before, for the *periodic* interval approximation, we can use the central approximation for every point, being mindful that we must take special care when computing the end-points to make sure that we do not over-run the arrays.

How do we use these functions? Below we provide codes for testing both the non-periodic and periodic first and second derivative approximations. One thing we would like to point out is our selection of dx. Notice that if we have N points, and that we want points to be uniformly spaced throughout the interval *including the end-points*, then we must chose dx to be the length of the interval divided by (N-1).

```
#include <iostream.h>
#include <iomanip.h>
#include "SCchapter5.h"

double func(double x);
double func_first_der(double x);
double func_second_der(double x);

int main(int argc, char * argv[]){
   const int levels = 10; //number of levels to test
   const double a = 0.0; //left end point of domain
   const double b = 1.0; //right end point of domain
   int i,j,npts;
   double dx,dxp,ux_error,uxx_error;
   double *u,*u_x,*u_xx;

cout << "npts\tError (First Deriv)\tError (Second Deriv)\n";</pre>
```

}

}

```
for(i=2;i<levels+2;i++){
    npts = (int) pow(2,i); //number of grid points is equal
                            // to 2^level
    dx = 1.0/(npts-1); //set dx based on number of points
    // Allocate storage dynamically
    u = new double[npts];
    u_x = new double[npts];
    u_xx = new double[npts];
    for(j=0;j<npts;j++)</pre>
      u[j] = func(j*dx); // set function value
    SO_FirstDeriv_1D (npts,dx,u,u_x); //calc. 1st deriv.
    SO_SecondDeriv_1D (npts,dx,u,u_xx); //calc. 2nd deriv.
    // Computation of the L2 error
    ux_error=0.0;
    uxx_error=0.0;
    for(j=0;j<npts;j++){</pre>
      ux_error += dx*pow((u_x[j]-func_first_der(j*dx)),2);
      uxx_error += dx*pow((u_xx[j]-func_second_der(j*dx)),2);
    }
    cout << setprecision(10) << setiosflags(ios::scientific);</pre>
    cout << npts << "\t" << sqrt(ux_error);</pre>
    cout << "\t" << sqrt(uxx_error) << endl;</pre>
    //Deallocation of dynamic memory
    delete[] u;
    delete[] u_x;
    delete[] u_xx;
  }
double func(double x){
 return(x*x*x*x);
}
double func_first_der(double x){
  return(4*x*x*x);
double func_second_der(double x){
```

```
return(12*x*x);
}
```

After compilation and execution of the program above, the following results are printed by the program.

npts	Error (First Deriv)	Error (Second Deriv)
4	4.4854079320e-01	2.0041110013e+00
8	6.9622902770e-02	2.4294654992e-01
16	1.3089009760e-02	3.6721619875e-02
32	2.7619740848e-03	6.1646753898e-03
64	6.2730612749e-04	1.1069179087e-03
128	1.4891039346e-04	2.1107879261e-04
256	3.6234744984e-05	4.2896354790e-05
512	8.9342897099e-06	9.2915223897e-06
1024	2.2180019754e-06	2.1242917384e-06
2048	5.5255279205e-07	5.0463557743e-07

According to the theory, both of these methods should be second-order accurate. If this were true, then doubling the number of grid points used should lead to a reduction in the error by a factor of four. Notice the convergence rate - just as predicted by the theory! Every time we increase the number of points by a factor of two, the error decreases by about a factor of four, hence we have second-order convergence.

The code below is a slight modification of the one above, in which we now demonstrate the use of the periodic interval functions. Observe that since we have N points, and that we want points to be uniformly spaced throughout the interval including only the left end-point, then we must chose dx to be the length of the interval divided by N. Why only the left end-point? Recall that we want the domain to be periodic. Hence, the last point value should not be the right-hand end of the domain because this value is periodic with the left-hand value. So we chose dx to be the length of the interval divided by N and not by (N-1). The code below demonstrates the use of the periodic differencing functions.

```
#include <iostream.h>
#include <iomanip.h>
#include "SCchapter5.h"

double func(double x);
double func_first_der(double x);
double func_second_der(double x);

int main(int argc, char * argv[]){
  const int levels = 10; //number of levels to test
  const double a = 0.0; //left end point of domain
  const double b = 1.0; //right end point of domain
```

}

}

```
int i,j,npts;
  double dx,dxp,ux_error,uxx_error;
  double *u,*u_x,*u_x;
  cout << "npts\tError (First Deriv)\tError (Second Deriv)\n";</pre>
  for(i=2;i<levels+2;i++){</pre>
    npts = (int) pow(2,i); //number of grid points is equal
                            // to 2^level
    dx = 1.0/(npts); //set dx based on number of points
    // Allocate storage dynamically
    u = new double[npts];
    u_x = new double[npts];
    u_xx = new double[npts];
    for(j=0;j<npts;j++)</pre>
      u[j] = func(j*dx); // set function value
    SO_FirstDeriv_1Dper (npts,dx,u,u_x); //calc. 1st deriv.
    SO_SecondDeriv_1Dper(npts,dx,u,u_xx); //calc. 2nd deriv.
    // Computation of the L2 error
    ux_error=0.0;
    uxx_error=0.0;
    for(j=0;j<npts;j++){</pre>
      ux_error += dx*pow((u_x[j]-func_first_der(j*dx)),2);
      uxx_error += dx*pow((u_xx[j]-func_second_der(j*dx)),2);
    }
    cout << setprecision(10) << setiosflags(ios::scientific);</pre>
    cout << npts << "\t" << sqrt(ux_error) << "\t";</pre>
    cout << << sqrt(uxx_error) << endl;</pre>
    //Deallocation of dynamic memory
    delete[] u;
    delete[] u_x;
    delete[] u_xx;
  }
double func(double x){
  return(sin(2.0*M_PI*x));
```

```
double func_first_der(double x) {
  return(2.0*M_PI*cos(2.0*M_PI*x));
}

double func_second_der(double x) {
  return(-4.0*M_PI*M_PI*sin(2.0*M_PI*x));
}
```

After compilation and execution of the program above, the following results are printed by the program.

npts	Error (First Deriv)	Error (Second Deriv)
4	1.6144558134e+00	5.2880398006e+00
8	4.4288293816e-01	1.4057888067e+00
16	1.1331413699e-01	3.5690383940e-01
32	2.8492869631e-02	8.9570538385e-02
64	7.1335237874e-03	2.2414226421e-02
128	1.7840256502e-03	5.6049069655e-03
256	4.4604671517e-04	1.4013111524e-03
512	1.1151419784e-04	3.5033306453e-04
1024	2.7878706903e-05	8.7583596378e-05
2048	6.9696865656e-06	2.1895920354e-05

Notice the convergence rate - just as predicted by the theory! Just as in the previous example, every time we increase the number of points by a factor of two, the error decreases by about a factor of four, hence we have second-order convergence.

5.1.3 MPI Parallel Implementation of Finite Differences

To implement the above differencing functions in MPI, we will first partition our domain among the processors. Assuming that the domain has been properly partitioned, it is the responsibility of each processor to compute the finite differences necessary for all points contained on that processor. Where do we need MPI? On each subdomain, the end-points of the domain need information that is not resident to the processor, but rather it resides on a different processor. Hence, we must use MPI_Send and MPI_Recv to transmit the information from one processor to another, as demonstrated in figure 5.5.



Below we present our implementation of 1D first derivative finite difference for a non-periodic interval:

• 1D First Derivative Parallel: MPI_Send/MPI_Recv: Non-periodic Interval

```
void SO_FirstDeriv_1DP(int npts, double dx, double *u,
```

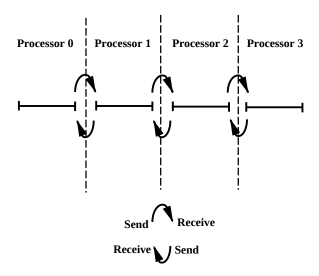


Figure 5.5: MPI Send/Receive pattern for finite differencing across processors.

```
double *u_x, int mynode, int totalnodes){
double two_invdx = 1.0/(2.0*dx);
double mpitemp;
MPI_Status status;
if(mynode == 0)
  u_x[0] = (-3.0*u[0] + 4.0*u[1] - u[2])*two_invdx;
if(mynode == (totalnodes-1))
  u_x[npts-1] = (3.0*u[npts-1] - 4.0*u[npts-2] +
                u[npts-3])*two_invdx;
for(int i=1;i<npts-1;i++)</pre>
  u_x[i] = (u[i+1]-u[i-1])*two_invdx;
if(mynode == 0){
  mpitemp = u[npts-1];
  MPI_Send(&mpitemp,1,MPI_DOUBLE,1,1,MPI_COMM_WORLD);
  MPI_Recv(&mpitemp,1,MPI_DOUBLE,1,1,MPI_COMM_WORLD,
           &status);
  u_x[npts-1] = (mpitemp - u[npts-2])*two_invdx;
else if(mynode == (totalnodes-1)){
  MPI_Recv(&mpitemp,1,MPI_DOUBLE,mynode-1,1,
           MPI_COMM_WORLD, &status);
  u_x[0] = (u[1]-mpitemp)*two_invdx;
```

```
mpitemp = u[0];
    MPI_Send(&mpitemp,1,MPI_DOUBLE,mynode-1,1,
             MPI_COMM_WORLD);
  }
  else{
    MPI_Recv(&mpitemp,1,MPI_DOUBLE,mynode-1,1,
             MPI_COMM_WORLD, &status);
    u_x[0] = (u[1]-mpitemp)*two_invdx;
    mpitemp = u[0];
    MPI_Send(&mpitemp,1,MPI_DOUBLE,mynode-1,1,
             MPI_COMM_WORLD);
    mpitemp = u[npts-1];
    MPI_Send(&mpitemp,1,MPI_DOUBLE,mynode+1,1,
             MPI_COMM_WORLD);
    MPI_Recv(&mpitemp, 1, MPI_DOUBLE, mynode+1, 1,
             MPI_COMM_WORLD, &status);
    u_x[npts-1] = (mpitemp-u[npts-2])*two_invdx;
  }
  return;
}
```

Observe the following cases that need special attention:

- 1. Process 0 contains the left end-point, and hence special one-sided differences must be employed.
- 2. Process (totalnodes 1) contains the right end-point, and hence special one-sided differencing must be employed.
- 3. For any other process, differencing for interior points can be done, and communication with neighbors to obtain information concerning adjacent points should be performed.

Care must be taken to guarantee that at the point that one process is sending information, that another process is ready to receive information. Notice the careful staggering of MPI_Send and MPI_Recv . If two processors were to call MPI_Send at the same time, with messages intended for each other, then we would arrive at what is referred to as a race-condition, and both processors would sit forever waiting for the other to call MPI_Recv !

Key Concept

• Care must be taken to avoid *race-conditions*. Recall that for every Send, there must be a Receive ready to accept the information.

One thing you may notice is that in every case in the example above we are "exchanging" information; that is, every process wants to "swap information" with its adjacent process. We have implemented the above using a combination of sends and receives, however MPI provides us with the functions $MPI_Sendrecv$ and $MPI_Sendrecv_replace$, which can help to automate this process. We now present the function syntax, argument list, usage examples, and remarks for $MPI_Sendrecv$ and $MPI_Sendrecv_replace$.

MPI_Sendrecv/MPI_Sendrecv_replace

Function Call Syntax

```
int MPI_Sendrecv(
         void*
                          sendbuf
                                           in
         int
                          sendcount
                                           in
         MPI_Datatype
                          sendtype
                                           in
                          dest
         int
                                           in
                          sendtag
                                           in
         int
         void*
                          recvbuf
                                           out
         int
                          recvcount
                                           in
                                       /*
         MPI_Datatype
                                           in
                          recvtype
         int
                          source
                                           in
         int
                                           in
                          recvtag
         MPI_Comm
                                           in
                          comm
         MPI_Status*
                          status
                                           out
int MPI_Sendrecv_replace(
         void*
                             buffer
                                            in
         int
                             count
                                            in
         MPI_Datatype
                            sendtype
                                            in
                             dest
                                            in
         int
         int
                            sendtag
                                            in
                             source
         int
                                            in
         int
                             recvtag
                                            in
         MPI\_Comm
                             comm
                                            in
         MPI_Status*
                             status
                                            out
```

Understanding the Argument Lists

- sendbuf starting address of the send buffer.
- sendcount number of elements in the send buffer.
- sendtype data type of the elements in the send buffer.
- dest process rank of destination.

- sendtaq send message tag.
- recvbuf starting address of the receive buffer.
- recvcount number of elements in the receive buffer.
- recvtype data type of the elements in the receive buffer.
- source process rank of source process.
- \bullet sendtag send message tag.
- comm communicator.
- *status* status object.
- buffer starting address of the send/recv buffer (replace case).
- count number of elements in the send/recv buffer (replace case).

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be sent/recv
int process1, process2; //process rank of two
                         // processes to exchange data
int tag1,tag2; // integer message tag
MPI_Status status; // variable to contain status information
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
// Determine datasize, process1, process2
double * sendbuffer = new double[datasize];
double * recvbuffer = new double[datasize];
double * buffer
                  = new double[datasize];
if(mynode == process1){
 // The call below sends the contents of sendbuffer to
       process2 and obtains from process2 data placed
 //
        into recybuffer
```

```
MPI_Sendrecv(sendbuffer,datasize,MPI_DOUBLE,process2,
               tag1, recvbuffer, datasize, MPI_DOUBLE, process2,
               tag2,MPI_COMM_WORLD,&status);
  // The call below will 'swap' the contents of buffer with
       process2; note the corresponding call to
       MPI_Sendrecv_replace below
  //
  MPI_Sendrecv_replace(buffer,datasize,MPI_DOUBLE,process2,
                       tag1,process2,tag2,MPI_COMM_WORLD,
                       &status);
}
if(mynode == process2){
  // The call below sends the contents of sendbuffer to
       process1 and obtains from process1 data placed
  //
       into recybuffer
  MPI_Sendrecv(sendbuffer,datasize,MPI_DOUBLE,process1,
               tag2,recvbuffer,datasize,MPI_DOUBLE,process1,
               tag1,MPI_COMM_WORLD,&status);
  // The call below will 'swap' the contents of buffer
        with process1; note the corresponding call to
  //
        MPI_Sendrecv_replace above
  //
  MPI_Sendrecv_replace(buffer,datasize,MPI_DOUBLE,process1,
                       tag2,process1,tag1,MPI_COMM_WORLD,
                       &status);
}
// At this point, process1 has in its recvbuffer the contents
// of process1's sendbuffer, process2 has in its recvbuffer
// the contents of process1's sendbuffer, and process1
// and process2's buffer arrays have been exchanged with
// each other.
```

Remarks

- MPI_Sendrecv allows us to send to one process and receive from another process (or the same process to which we are sending) in one function call. This type of MPI function is ideal for swapping or shifting information among a collection of processes.
- MPI_Sendrecv_replace allows us the functionality of MPI_Sendrecv with the additional caveat that the information which is being sent is over-written by the received information. MPI guarantees that the "sending" is accomplished before the "receiving".
- For both $MPI_Sendrecv$ and $MPI_Sendrecv_replace$, the sending and receiving processes do not have to be the same. For instance, process one can send data to process two while obtaining data from process zero.
- For $MPI_Sendrecv$, MPI_Recv can be used on the receiving process to obtain the information being sent. Similarly, $MPI_Sendrecv$ can receive data from a process sending using MPI_Send .
- In most cases the *sendtype* and *recvtype* are identical.
- The tag can be any integer between 0-32767.



We implement again the finite differencing function given above, now using $MPI_Sendrecv_replace$ as opposed to MPI_Send/MPI_Recv . Notice that the "race-condition" problem is handled by MPI, and not by us!

Key Concept

• If you find yourself repeating some pattern of MPI_Send and MPI_Recv calls, it is very likely that an MPI function already exists which combines the pattern into one MPI call!

• 1D First Derivative Parallel: MPI_Sendrecv_replace: Non-periodic Interval

```
if(mynode == 0)
    u_x[0] = (-3.0*u[0] + 4.0*u[1] - u[2])*two_invdx;
  if(mynode == (totalnodes-1))
    u_x[npts-1] = (3.0*u[npts-1] - 4.0*u[npts-2] +
                  u[npts-3])*two_invdx;
  for(int i=1;i<npts-1;i++)</pre>
    u_x[i] = (u[i+1]-u[i-1])*two_invdx;
  if(mynode == 0){
    mpitemp = u[npts-1];
    MPI_Sendrecv_replace(&mpitemp,1,MPI_DOUBLE,1,1,1,1,
                         MPI_COMM_WORLD, &status);
    u_x[npts-1] = (mpitemp - u[npts-2])*two_invdx;
  }
  else if(mynode == (totalnodes-1)){
    mpitemp = u[0];
    MPI_Sendrecv_replace(&mpitemp,1,MPI_DOUBLE,mynode-1,
                          1,mynode-1,1, MPI_COMM_WORLD,
                         &status);
    u_x[0] = (u[1]-mpitemp)*two_invdx;
  }
  else{
    mpitemp = u[0];
    MPI_Sendrecv_replace(&mpitemp,1,MPI_DOUBLE,mynode-1,
                          1,mynode-1,1, MPI_COMM_WORLD,
                         &status);
    u_x[0] = (u[1]-mpitemp)*two_invdx;
    mpitemp = u[npts-1];
    MPI_Sendrecv_replace(&mpitemp,1,MPI_DOUBLE,mynode+1,
                          1,mynode+1,1, MPI_COMM_WORLD,
                         &status);
    u_x[npts-1] = (mpitemp-u[npts-2])*two_invdx;
  }
  return;
}
```



In the *software suite* we present a C++ program which uses the MPI differencing functions provided above. It is very similar to the scalar programs shown earlier. Also, we use the MPI_Reduce function, which was discussed earlier, for collecting the error information from all the processors and supplying it to process 0.

We also include several C++/MPI implementations for first and second derivatives on periodic and non-periodic domains using MPI_Send/MPI_Recv and also using $MPI_Sendrecv_replace$.

Variable Coefficient

A situation that arises often in applications is a second-order derivative term with variable coefficient, which is a function of the location and thus it may vary along the grid points. To maintain the desired accuracy in this case we have to symmetrize the discretization as follows:

$$\frac{\partial}{\partial x} \left[\nu(x) \frac{\partial}{\partial x}\right] u_i = \frac{1}{\Delta x^2} \delta^-(\nu_{i+1/2} \delta^+) u_i + \mathcal{O}(\Delta x^2) = \frac{1}{\Delta x^2} \delta^+(\nu_{i-1/2} \delta^-) u_i + \mathcal{O}(\Delta x^2).$$

Thus.

$$\frac{\partial}{\partial x} \left[\nu(x) \frac{\partial}{\partial x} \right] u_i = \frac{\nu_{i+1/2} (u_{i+1} - u_i)}{\Delta x^2} - \frac{\nu_{i-1/2} (u_i - u_{i-1})}{\Delta x^2} + \mathcal{O}(\Delta x^2).$$

Many diffusion problems with space-dependent diffusivity are formulated in this fashion.

5.1.4 Multi-Dimensional Arrays in C++

As the level of complexity of our programs increases, we will increasingly find that single indexed (one-dimensional) arrays do not fulfill all our programming needs. This is not to say that we cannot use single-arrays for doing all of our work, but rather that sometimes the algorithm naturally breaks itself down into what we refer to as multi-dimensional arrays (for example, matrices, which can be considered an $N \times M$ array). Just as with single-dimension arrays, there are two ways to allocate multi-dimensional arrays: statically and dynamically.

Static Allocation of Multi-Dimensional Arrays

Just as with single-indexed arrays, we can allocate multi-dimensional arrays statically as follows: Suppose that we want to declare an array of double which is 20×20 in size. We would use the following array declaration:

double x[20][20];

We have now declared an array x which contains 20×20 doubles, which can be indexed i = 0, 19 in the first index, and j = 0, 19 in the second index. Hence if we wanted item (3,4), we would access x[3][4].

If we want more than two dimensions, we can continue to append dimensions as follows:

```
double x[20][20][20][20];
```

which is a $20 \times 20 \times 20 \times 20$ array! All of the previous rules introduced with single-indexed arrays still apply.

Dynamic Allocation of Multi-Dimensional Arrays

Dynamic allocation of multi-dimensional arrays is very similar to that of single dimension arrays. Suppose, once again, that we now want to dynamically allocate an array of 10×10 elements. This can be done as follows:

```
int npts = 10;
double ** x;

x = new double*[npts];
for(i=0;i<npts;i++)
    x[i] = new double[npts];</pre>
```

And to delete, we would use the following reversal of the process:

```
for(int i=0;i<npts;i++)
   delete[] x[i];
delete[] x;</pre>
```

The key points to get from this are the following:

- For a single dimension array, we used a double*. For a two-dimensional array, we use a double**. For a three-dimensional array, we use a double ***, and so on. This is because at each successive level, we are declaring an "array of arrays". A two-dimensional array is an array of one-dimensional arrays. A three-dimensional array is an array of two-dimensional arrays, which, in turn, are an array of one-dimensional arrays.
- Observe that when declaring a two-dimensional array, we first must declare the array of double* variables, and then we assign to each double* variable an allocation of a single dimensional array. A schematic of this is shown in figure 5.6.
- Creating arrays in this fashion does not create contiguous blocks! This means that what we have shown you here gives the most freedom to the operating system to decide how to fit things into memory.
- Observe that when deleting the dynamic allocation, we must reverse the allocation process. First, we must deallocate the single dimension arrays that we assigned, and then we must delete the original array.

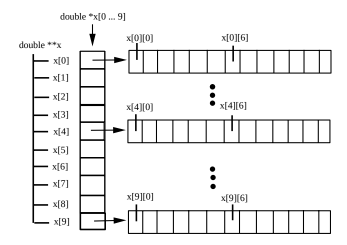


Figure 5.6: Diagram showing conceptual picture of two-dimensional array memory allocation.

Dynamic Allocation of Multi-Dimensional Arrays in Contiguous Blocks

In some instances when using BLAS or MPI it may be necessary to allocate multi-dimensional arrays as a contiguous block of memory, which is different than what was previously shown. This is because both BLAS and MPI act on contiguous blocks of memory. To create a two-dimensional array in which the allocation is a continuous block of memory, allocate the block of memory and index the pointer array ourselves. We will first present a sample code as to how this is accomplished, and then we will comment on this code.

Observe after allocating our indexing array, we allocate a contiguous block of memory equal to the total size that we need and then assign the leading address of that block of memory to the first indexed position of our indexing array. We then loop through the remainder of the indexing array and assign to each position the appropriately offset amount as shown in figure 5.7. With this type of memory allocation we are certain that x[0] points to a contiguous block of memory representing the entire $N \times N$ matrix. Hence, in MPI, if we wanted to send this matrix to another process, we could pass x[0] as the starting pointer and send $N \times N$ doubles to a receiving process.

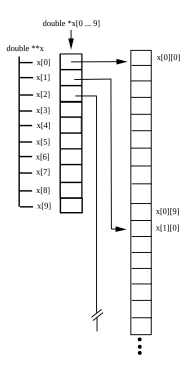


Figure 5.7: Diagram showing conceptual picture of two-dimensional array memory allocation with contiguous memory.

Row-Major Order versus Column-Major Order

Previously we mentioned that there is a difference between "row-major order" and "column-major order". In this section we will explain the differences between these two orderings and explain why they are significant.

Multi-dimensional arrays require us to derive a mapping from the multi-dimensional ordering of the array to a linearly addressable ordering used for accessing memory. In figure 5.8 we present a diagram demonstrating the difference between a row-major ordering and a column-major ordering.

Observe that in a row-major ordering, we map the array row-by-row into the linearly addressed memory, whereas in a column-major ordering we map the array column-by-column. It is important to be mindful row-major ordering and column-major ordering for the following reasons:

- When using BLAS (or ATLAS), by default matrices are assumed to be stored in column-major ordering (consistent with FORTRAN) instead of row-major ordering (consistent with C/C++). You may be required to manually re-order your matrix to accommodate BLAS. Another option provided by BLAS so that manual reordering is not necessary is to specify that you are passing the transpose of the matrix (the transpose of a row-major ordered matrix is a column-major ordered matrix). Care must be taken though when interpreting the results of the BLAS call. The resulting operation will be in column-major order.
- Whether a matrix is stored in column-major order or row-major order determines what is the most efficient way to access the array entries. Suppose we needed to print

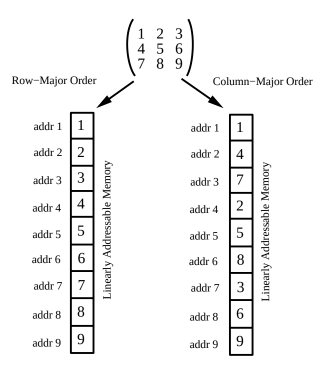


Figure 5.8: Diagram showing a conceptual picture of row-major order versus column-major order for the contiguous allocation of a two-dimensional array. Observe that both orderings require the same quantity of linearly addressable memory, however the distribution of the two-dimensional array elements differ based upon which ordering is used.

the contents of a matrix, and we wanted to access memory in the most efficient way possible. Consider the following looping example:

```
// Optimal for column-major ordering
for(int i=0;i<N;i++)
  for(int j=0;j<N;j++)
     cout << A[j][i] << endl;

// Optimal for row-major ordering
for(int i=0;i<N;i++)
  for(int j=0;j<N;j++)
     cout << A[i][j] << endl;</pre>
```

C++ uses row-major ordering; the latter example shows the optimal means of accessing the contents of the multi-dimensional array A. We design the loops so that the *innermost loop* runs over the *outer-most index*.

5.1.5 Non-Uniform Grids

We often need to compute derivatives on a non-uniform grid which is present because of *steep* gradients in the solution in some regions and smooth variation in other regions. Such solution

behavior is present in boundary layers, but it may also be dictated by the geometry of the problem or because of semi-infinite or infinite computational domains. Here, we present two different ways of handling non-uniform grids, first using *Lagrangian interpolation* and second using *mappings*.

Lagrangian Interpolation and Non-Uniform Grids

So far we have obtained discretizations of the first and second derivatives on uniforms grids. Next, we assume that we have an arbitrary distribution of grid points denoted by $(\alpha_0, \alpha_1, \ldots, \alpha_N)$ as shown in the sketch 5.9. We will look for approximations of the m^{th} -order derivative on a stencil consisting of n points where $n \geq m$. We allow m = 0 which is the degenerate case for interpolation. This method was first proposed by Fornberg, and here we have adopted his notation [37].

Figure 5.9: The grid points are distinct but arbitrarily distributed in the domain.

Let us consider the function f(x) and look for the approximation of the m^{th} -order derivative at the point x_0

$$\frac{d^m f}{dx^m}|_{x=x_0} = \sum_{\nu=0}^n c_{n,\nu}^m f(\alpha_{\nu}) \quad m = 0, 1, \dots, M \\ n = m, m+1, \dots, N$$
 (5.6)

The question is then how to compute $c_{n,\nu}^m$ efficiently for any point in the interval $x \in [\alpha_0, \alpha_N]$. The key idea here is to use *Lagrangian interpolation*.

For simplicity, let us consider the point $x_0 = 0$. We then define

$$F_n(x) \equiv \prod_{k=0}^n (x - \alpha_k),$$

and thus the nth-degree Lagrangian polynomial is

$$h_{\nu}^{n}(x) \equiv \frac{F_{n}(x)}{F_{n}'(\alpha_{\nu})(x - \alpha_{\nu})}; \quad h_{\nu}^{n}(\alpha_{k}) = \delta_{\nu k}.$$

Then we can perform Lagrangian interpolation for f(x), i.e.,

$$f(x) \approx \sum_{\nu=0}^{n} h_{\nu}^{n}(x) f(\alpha_{\nu}). \tag{5.7}$$

Now by comparing equation (5.6) and the m^{th} -derivative of the Lagrangian representation in equation (5.7), we obtain

$$c_{n,\nu}^{m} = \frac{d^{m}}{dx^{m}} h_{\nu}^{n}(x)|_{x=0}$$
.

Inversely, the nth-degree polynomial $h_{\nu}^{n}(x)$ can be expressed using a Taylor expansion of the form:

$$h_{\nu}^{n}(x) = \sum_{m=0}^{n} \frac{c_{n,\nu}^{m}}{m!} x^{m}.$$

The next step is to obtain recurrence formulas for $h_{\nu}^{n}(x)$ so that they can be computed efficiently, and subsequently compute $c_{n,\nu}^{m}$ recursively. To this end,

$$F_n(x) = (x - \alpha_n) F_{n-1}(x) F'_n(x) = (x - \alpha_n) F'_{n-1}(x) + F_{n-1}(x).$$

This recursive function calling is implemented in the example below. We assume that the array alpha[i] contains the grid coordinates as shown in figure 5.9.

```
double F(int n, double x, double *alpha){
  double answer;

if(n==0)
  answer = x - alpha[n];
  else
  answer = (x-alpha[n])*F(n-1,x,alpha);

return answer;
}
```

The recursion for the Lagrangian polynomials is then obtained from:

•
$$\nu \neq n$$
: $h_{\nu}^{n}(x) = \frac{F_{n}(x)}{F'_{n}(\alpha_{\nu})(x - \alpha_{\nu})} = \frac{(x - \alpha_{n})F_{n-1}(x)}{(\alpha_{\nu} - \alpha_{n})F'_{n-1}(\alpha_{\nu})(x - \alpha_{\nu})}$

$$= \frac{x - \alpha_{n}}{\alpha_{\nu} - \alpha_{n}} \frac{F_{n-1}(x)}{F'_{n-1}(\alpha_{\nu})(x - \alpha_{\nu})}$$

$$= \frac{x - \alpha_{n}}{\alpha_{\nu} - \alpha_{n}} h_{\nu}^{n-1}(x),$$

•
$$\nu = n$$
: $h_n^n(x) = \frac{F_{n-1}(x)}{F_{n-1}(\alpha_n)} = \frac{F_{n-2}(\alpha_{n-1})}{F_{n-1}(\alpha_n)} (x - \alpha_{n-1}) h_{n-1}^{n-1}(x)$.

By equating coefficients in the expansion for $h_{\nu}^{n}(x)$ we obtain:

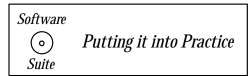
$$x_0 = 0 \begin{cases} n \neq \nu : & c_{n,\nu}^m = \frac{1}{\alpha_n - \alpha_\nu} (\alpha_n c_{n-1,\nu}^m - m c_{n-1,\nu}^{m-1}) \\ \text{and} \\ n = \nu : & c_{n,n}^m = \frac{F_{n-2}(\alpha_{n-1})}{F_{n-1}(\alpha_n)} (m c_{n-1,n-1}^{m-1} - \alpha_{n-1} c_{n-1,n-1}^m) . \end{cases}$$

We can also use

$$\sum_{\nu=0}^{n} c_{n,\nu}^{m} = \begin{cases} 1, m = 0 \\ 0, m \neq 0 \end{cases},$$

however, this formula should not be used because it may induce round-off error.

Note that for $x_0 \neq 0$ we replace $\begin{cases} \alpha_n \to (\alpha_n - x_0) \\ \alpha_{n-1} \to (\alpha_{n-1} - x_0) \end{cases}$ and that there is no restriction on x_0 coinciding with any α_{ν} .



The code below computes the weights in the *Fornberg method* and gives us the opportunity to introduce a triple array, which can be viewed as an array of two-dimensional arrays. Notice that we use a double ***! The indexing of this array is just as we discussed earlier.

```
void FornbergWeights(double xi, double *x, int m, int n,
                      double ***C){
  int i,j,k,mn;
  double C1,C2,C3;
  C[0][0][0] = 1.0;
  C1 = 1.0;
  for(j=1; j \le n; j++) {
    if(j < m)
      mn = j;
    else
      mn = m;
    C2 = 1.0;
    for(k=0;k<=(j-1);k++){
      C3 = x[j]-x[k];
      C2 = C2*C3;
      if(j \le m) C[j][j-1][k] = 0.;
      C[0][j][k] = (x[j]-xi)*C[0][j-1][k]/C3;
      for(i=1;i<=mn;i++)
        C[i][j][k] = ((x[j]-xi)*C[i][j-1][k]-i*
                      C[i-1][j-1][k])/C3;
    }
```

Remark 1: The triple indexing of the double*** array C follows the same rules as previously discussed for double indexed arrays. A diagram showing a conceptual picture of the three-dimensional array is given in figure 5.10. The first index i of C[i][j][k] denotes which double** array is selected. The second index j of C[i][j][k] denotes which double* array is selected. The third index k of C[i][j][k] denotes which double is selected.

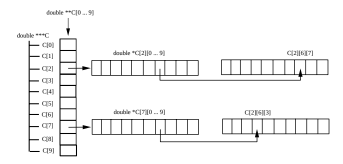


Figure 5.10: Diagram showing conceptual picture of three-dimensional array memory allocation.

Remark 2: Pay careful attention to the required size of the array C. It is the calling functions responsibility to allocate appropriate space for C. The routine above does allocate memory; it assumes that the three-dimensional array C has already been declared. By examining the loops within the routine above, it is possible to deduce a required size for C given the parameters m and n.

Mappings and Non-Uniform Grids

It is common practice to transform non-uniform grids to uniform grids (if possible), and then on the uniform grid to compute the discrete derivatives. This is due to the fact that the formal accuracy (e.g., second-order) of the scheme is maintained on the uniform grid whereas only first-order accuracy can be achieved on the original (non-uniform) grid. However, the actual accuracy depends strongly on the transformation between the non-uniform grid $[x]_i$ and the uniform grid $[\xi]_i$, and thus it is not always clear which is the better grid to compute discrete derivatives. We analyze this next, following the work of Hoffman [58].

Let us assume that the two grids are related by the transformation

$$x = \phi(\xi) \Rightarrow \xi = \phi^{-1}(x). \tag{5.8}$$

On the uniform grid (ξ_i) the first derivative of the function u(x) is

$$u_x = u_{\xi} \phi_x^{-1},$$

and a central difference approximation at point (i) is

$$(u_x)_i^{\xi} = (\phi_x^{-1})_i \frac{u_{i+1} - u_{i-1}}{\xi_{i+1} - \xi_{i-1}}.$$

From Taylor expansion we can also obtain that

$$(u_x)_i^{\xi} = (\phi_x^{-1})_i (u_{\xi})_i + \mathcal{O}(\Delta \xi^2).$$
 (5.9)

It is now clear from equation (5.9) that the approximation of the first derivative on a uniform grid is of second-order. On the other hand, a central difference formula for the first derivative on a uniform grid produces a first-order accuracy approximation of the form

$$(u_x)_i^x = (u_x)_i + \frac{1}{2}(u_{xx})_i(\Delta x_+ - \Delta x_-) + \mathcal{O}(\Delta x_\pm^2), \tag{5.10}$$

where $\Delta x_+ \equiv x_{i+1} - x_i$ and $\Delta x_- \equiv x_{i+1} - x_i$. Thus if Δx_{\pm} is halved by doubling the number of grid points, then the error decreases only by a factor of two, which corresponds to first-order accuracy. However, it is also possible to redistribute the points during refinement so that by doubling the number of grid points the term $(\Delta x_+ - \Delta x_-)$ quarters. In this case, second-order accuracy is achieved even on a non-uniform grid. A procedure to obtain this was derived by Hoffman [58] by expanding the coordinate transformation $x = \phi(\xi)$ in a Taylor series

$$\Delta x_{+} - \Delta x_{-} = \phi_{\xi\xi} \Delta \xi^{2} + \mathcal{O}(\Delta \xi^{4}).$$

This equation suggests that if we half the grid spacing on the uniform grid, then the effective spacing on the non-uniform grid is quartered, and thus we recover second-order type behavior for the approximation error. The corresponding truncation errors on the two grids are also related by

$$(u_x)_i^x = (u_x)_i^\xi + \mathcal{O}(\Delta \xi^2).$$

In the special case that the transformation $\phi(\xi)$ is a polynomial of second-order or less, the two derivative approximations are identical because the truncation error in the above equation is identically zero.

Similar conclusions can be drawn for higher derivatives. For example, for the second derivative the difference in the approximation on the two grids is zero if the transformation is a first-order polynomial.

Some commonly used mappings are:

• Exponential Stretching:

$$x = L(e^{a\xi} - 1)$$

with the purpose of increasing resolution around x = 0, where L is the length of the domain and $\xi \in [0, 1]$. The parameter a determines the exact point distribution.

• Semi-Infinite Domain:

- 1. Algebraic mapping: $x = L \frac{1+\xi}{1-\xi}, \ x \in [0,\infty] \to \xi \in [-1,1]$
- 2. Logarithmic mapping: $x = -L \ln \left(\frac{1-\xi}{2} \right), \ [0, \infty] \to [-1, 1]$

• Infinite Domain:

- 1. Exponential mapping: $x = L \tanh^{-1}, \xi \ x \in [-\infty, \infty] \to \xi \in [-1, 1]$
- 2. Algebraic mapping: $x = L \frac{\xi}{\sqrt{1-\xi^2}}, \quad x \in [-\infty, \infty] \to \xi \in [-1, 1]$

5.1.6 One-Dimensional Boundary Value Problem

Next we apply the discretization of the second derivative for the one-dimensional boundary value problem of the form

$$\begin{cases} \theta'' = -\sin 2\pi x \\ \theta_0 = \theta_N = 0 \end{cases}$$

subject to homogeneous boundary conditions. The exact solution is obtained analytically

$$\Theta(x) = \frac{1}{4\pi^2} \sin 2\pi x.$$

The difference equation is obtained by discretizing the second derivative and apply the equation at the node (i) neglecting the truncation error:

$$\frac{1}{\Delta x^2} (\theta_{i-1} - 2\theta_i + \theta_{i+1}) = -\sin 2\pi x_i, \quad \theta_0 = \theta_N = 0.$$

The equivalent differential equation is obtained by including on the right-hand-side of the differential equation the truncation terms, i.e.,

$$\hat{\theta}'' = -\sin 2\pi x - \underbrace{\frac{\Delta x^2}{12} \hat{\theta}^{(iv)} + \dots}_{\text{truncation error}}.$$

We note that $T_i \to 0$ as $\Delta x \to 0$, and therefore our discretization is *consistent* as we recover the original differential equation when the discretization parameter approaches zero. This condition is called *consistency* and it is the key to convergence for boundary value problems (derived from the Lax theorem [77]).

We can now proceed to solve the difference equation analytically by assuming a solution of the form:

$$\theta_i = A \sin 2\pi i \Delta x \ (i = 0, N \Rightarrow \theta_i = 0).$$

Therefore,

or

$$\frac{A}{\Delta x^2} [\sin 2\pi (i-1)\Delta x - 2\sin 2\pi i\Delta x + \sin 2\pi (i+1)\Delta x] = -\sin 2\pi i\Delta x,$$

$$\frac{A}{\Delta x^2} \sin 2\pi i\Delta x (\cos 2\pi \Delta x - 2 + \cos 2\pi \Delta x) = -\sin 2\pi i\Delta x$$

$$\Rightarrow A = \frac{\Delta x^2}{2(1 - \cos 2\pi \Delta x)} = \frac{\Delta x^2}{2[1 - (1 - \frac{(2\pi)^2 \Delta x^2}{2} + \frac{(2\pi)^4 \Delta x^4}{24}) + \dots]}$$

$$= \frac{\Delta x^2}{(2\pi)^2 \Delta x^2 - \frac{(2\pi)^4 \Delta x^4}{12} + \dots}$$

$$= \frac{1}{4\pi^2 \left[1 - \frac{(2\pi)^2 \Delta x^2}{12} + \dots\right]}$$

$$= \frac{1}{4\pi^2} \left[1 + \frac{(2\pi)^2 \Delta x^2}{12} + \dots\right].$$

Finally, the solution of the difference equation is

$$\theta_i = \underbrace{\frac{1}{4\pi^2} \sin 2\pi x_i}_{\Theta \text{ exact}} + \frac{\Delta x^2}{12} \sin 2\pi x_i + \dots$$

We have shown directly that the approximation of the second derivative with second-order accuracy leads to a solution which is second-order accurate.

To solve the equivalent differential equation we use perturbation expansions taking Δx^2 as a small parameter:

$$\hat{\theta} = \hat{\theta}_0 + \Delta x^2 \hat{\theta}_2 + \Delta x^4 \hat{\theta}_4 + \dots$$

By matching successive powers of Δx we obtain:

$$\Delta x^{0} \colon \begin{array}{l} \hat{\theta}_{0}^{"} = -\sin 2\pi x \\ \hat{\theta}_{0}(0) = \hat{\theta}_{0}(1) = 0 \end{array} \right\} \Rightarrow \hat{\theta}_{0} = \Theta = \frac{1}{4\pi^{2}} \sin 2\pi x$$

$$\Delta x^{2} \colon \begin{array}{l} \hat{\theta}_{2}^{"} = -\frac{1}{12} \hat{\theta}_{0}^{(iv)} = -\frac{(2\pi)^{4}}{12(4\pi^{2})} \sin 2\pi x \\ \hat{\theta}_{2}(0) = \hat{\theta}_{2}(1) = 0 \end{array} \right\} \Rightarrow \hat{\theta}_{2} = \frac{1}{12} \sin 2\pi x$$

$$\hat{\theta}(x) = \frac{1}{4\pi^{2}} \sin 2\pi x + \frac{\Delta x^{2}}{12} \sin 2\pi x + \dots$$

so,

 $v(x) = \frac{1}{4\pi^2} \sin 2\pi x + \frac{1}{12} \sin 2\pi x + \dots$ Notice that $\hat{\theta}(x_i) = \theta_i$ from above in other words $\hat{\theta}(x)$ the sol

Notice that $\hat{\theta}(x_i) = \theta_i$, from above, in other words $\hat{\theta}(x)$, the solution of the equivalent differential equation, collocates the solution at the nodes.

• Thus, the numerical solution is an "exact" solution at the nodes of the corresponding equivalent differential equation but not of the differential equation!

5.1.7 Multi-Dimensional Discretizations

Similar to multi-dimensional interpolation and integration, multi-dimensional differentiation is obtained by *splitting directions*, i.e. differentiating one direction engaging points only in that direction while fixing the other direction, in a Cartesian fashion. For example, we can use Taylor expansions again to obtain discrete derivatives in a five-point stencil or "molecule" (see figure 5.11).

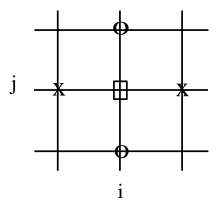


Figure 5.11: Sketch of the five point molecule.

At the marked points we obtain:

$$\times (u_x)_{ij} = \frac{u_{i+1,j} - u_{i-1,j}}{2\Delta x} - \frac{\Delta x^2}{12} \left(\frac{\partial^4 u}{\partial x^4}\right)$$

$$O(u_{yy})_{ij} = \frac{u_{i,j+1} - 2u_{ij} + u_{i,j-1}}{\Delta y^2} - \frac{\Delta y^2}{12} \left(\frac{\partial^4 u}{\partial y^4}\right).$$

Similarly, we can obtain the discrete Laplacian

$$\nabla^2 u_{ij} = (u_{xx} + u_{yy})_{ij} = \underbrace{\frac{u_{i-1,j} - 2u_{ij} + u_{i+1,j}}{\Delta x^2} + \frac{u_{i,j-1} - 2u_{ij} + u_{i,j+1}}{\Delta y^2}}_{\nabla^2_+ + \mathcal{O}(\Delta x^2, \Delta y^2)}$$
(5.11)

where we have introduced the Cartesian discrete Laplacian operator ∇_+^2 , and thus in symbolic form we can write

$$\nabla_+^2 u_{ij} = \left(\frac{\delta_x^2}{\Delta x^2} + \frac{\delta_y^2}{\Delta y^2}\right) u_{ij} ,$$

where δ_x and δ_y are the central difference operators defined in section 5.1.2.

In the example below we present a C++ code that implements the Laplacian 5-point molecule for the interior points. We assume that the boundary values of the differentiation

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are handled elsewhere in the code. The key thing to notice in this example is how easily the finite difference formulas translate into C++ code once you have a handle on how arrays work. In this example, we assume that we have already allocated double arrays for both u and u_xx_yy ; we merely need to index the arrays properly to obtain the desired result.

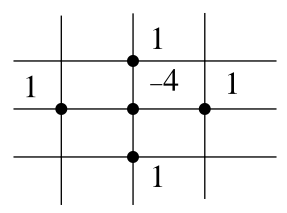


Figure 5.12: Five-point molecule and corresponding weights for a uniform grid.

The above formula can be simplified for **equidistant grids** in both x- and y-directions, that is for $\Delta x = \Delta y$ we obtain the simple molecule of figure 5.12. The numbers at the grid points are the corresponding weights in the discrete Laplacian, and thus

$$\nabla_{+}^{2} u_{ij} = \frac{u_{i-1,j} + u_{i+1,j} - 4u_{ij} + u_{i,j-1} + u_{i,j+1}}{\Delta x^{2}}.$$

Also, using symbolic notation we can derive the consistent second-order discrete Laplacian with a **variable coefficient**:

$$\nabla \cdot (\nu \nabla u)_{ij} = \frac{1}{\Delta x^2} \delta_x^- (\nu_{i+1/2,j} \delta_x^+) u_{ij} + \frac{1}{\Delta y^2} \delta_y^- (\nu_{i,j+1/2} \delta_y^+) u_{ij} + \mathcal{O}(\Delta x^2, \Delta y^2).$$

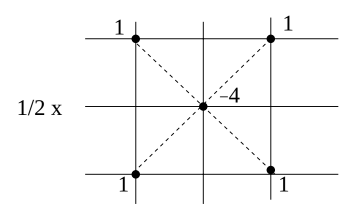


Figure 5.13: Weights for the diagonal splitting in a uniform grid.

An alternative approach to the Cartesian direction splitting we followed to derive the discrete operator ∇_+^2 is to use **diagonal splitting**. In this case we derive the following discrete Laplacian operator

$$\nabla_x^2 u_{ij} = \left[\frac{(\mu_y \delta_x)^2}{\Delta x^2} + \frac{(\mu_x \delta_y)^2}{\Delta y^2} \right] u_{ij}$$

$$= \frac{1}{4\Delta x^2} [(E_y + E_y^{-1} + 2)(E_x + E_x^{-1} - 2)] u_{ij}$$

$$+ \frac{1}{4\Delta y^2} \left[(E_x + E_x^{-1} + 2)(E_y + E_y^{-1} - 2) \right] u_{ij},$$

which for $\Delta x = \Delta y$ has the simpler form

$$\Rightarrow \nabla_x^2 = \frac{1}{2\Delta x^2} [E_x E_y + E_x^{-1} E_y^{-1} + E_x^{-1} E_y + E_x E_y^{-1} - 4].$$

The corresponding stencil is shown diagrammatically in figure 5.13 with the appropriate weights.

However, there is a problem with diagonal splitting as it leads to two independent stencils as shown in figure 5.14, which in turn will produce *odd-even oscillations* in the odd-numbered and even-numbered grid points.

To overcome this difficulty we define a new operator that combines both approaches, i.e., Cartesian and diagonal splitting, using appropriate weights. To this end, we write

$$\nabla_{\star}^{2} u_{ij} = (a \nabla_{\perp}^{2} + b \nabla_{\tau}^{2}) u_{ij}; \quad a+b=1.$$

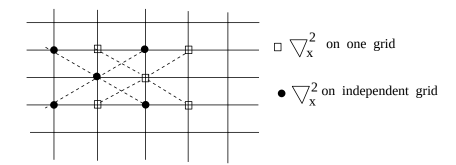


Figure 5.14: Decoupling of the discrete Laplacian operator induced by diagonal splitting.

If we now substitute for ∇_+^2 and ∇_x^2 and also a=1-b, assuming also a uniform grid $(\Delta x = \Delta y)$, we obtain

$$\nabla_*^2 = \frac{1}{\Delta x^2} (\delta_x^2 + \delta_y^2) + \frac{b}{\Delta x^2} [(\mu_y \delta_x)^2 + (\mu_x \delta_y)^2 - (\delta_x^2 + \delta_y^2)]$$
$$= \frac{1}{\Delta x^2} \left\{ (\delta_x^2 + \delta_y^2) + \frac{b}{2} \left[2\delta_x^2 (\mu_y^2 - 1) + 2\delta_y^2 (\mu_x^2 - 1) \right] \right\}$$

but

$$2\delta_x^2(\mu_y^2 - 1) = 2(E_x + E_x^{-1} - 2) \left[\frac{1}{4} (E_y + E_y^{-1} + 2) - 1 \right]$$
$$= \frac{1}{2} (E_x + E_x^{-1} - 2) (E_y + E_y^{-1} - 2)$$
$$= \frac{1}{2} \delta_x^2 \delta_y^2.$$

By symmetry the second term is the same, and therefore we have:

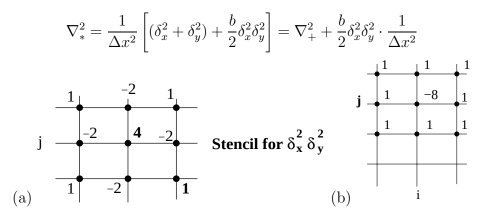


Figure 5.15: (a) Stencil for the product $\delta_x^2 \delta_y^2$. (b) Stencil for b = 2/3.

It is clear that we can now construct a family of two-dimensional stencils for the discrete Laplacian by selecting different values for $b \in (0,1)$. In figure 5.15 we first construct the

weights for the product $\delta_x^2 \delta_y^2$, and subsequently we construct a stencil for b = 2/3. This stencil is similar to a nine-node quadrilateral finite element [59].

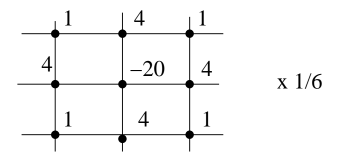


Figure 5.16: Stencil for the Dahlquist-Bjorck stencil.

Another useful construction is the **Dahlquist-Bjorck stencil** corresponding to b = 1/3 (see figure 5.16). In this case we get

$$\nabla_*^2 u_{ij} = \nabla^2 u_{ij} + \frac{\Delta x^2}{12} \left[\underbrace{\frac{\partial^4 u}{\partial x^4} + \frac{\partial^4 u}{\partial y^2} + 6 \cdot \frac{1}{3} \frac{\partial^4 u}{\partial x^2 \partial y^2}}_{\text{truncation error}} \right]$$

From this construction and from the definitions above we can derive

$$\nabla_*^2 u_{ij} = \nabla^2 u_{ij} + \frac{\Delta x^2}{12} \nabla^4 u.$$

This form suggests that in solving the eigenvalue problem $\nabla^2 u = \lambda u$ the truncation error is $-\frac{\Delta x^2}{12}(\lambda^2 u)$, and thus by ejecting its opposite on the right-hand-side as follows

$$\nabla_*^2 u_{ij} = (\lambda + \frac{\lambda^2 \Delta x^2}{12})u$$

we obtain a discretization with a fourth-order truncation error although the discrete operators were all of the second-order. This is called the *method of corrected differences*, or the *booster* method, and it is used often in scientific computing to enhance the formal accuracy of the method.

We can proceed similarly to discretize **mixed derivatives**

$$\frac{\partial^2 u}{\partial x \partial y}.$$

Using symbolic notations and operators defined previously we obtain

$$u_{xy} = \frac{1}{\Delta x \Delta y} \mu_x \delta_x \left[\left(1 - \frac{\delta_x^2}{6} \right) + \mathcal{O}(\Delta x^4) \right] \mu_y \delta_y \left[\left(1 - \frac{\delta_y^2}{6} \right) + \mathcal{O}(\Delta y^4) \right] u_{ij}.$$

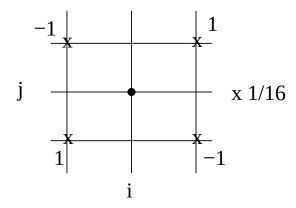


Figure 5.17: Stencil and corresponding weights for a mixed second derivative.

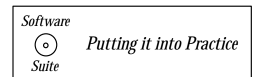
From this we can construct the second-order accurate stencil

$$u_{xy} = \frac{1}{\Delta x \Delta y} \mu_x \delta_x \mu_y \delta_y u_{ij} + \mathcal{O}(\Delta x^2, \Delta y^2),$$

which is shown also diagrammatically in figure 5.17. More explicitly

$$(u_{xy})_{ij} = \frac{1}{4\Delta x \Delta y} \left[u_{i+1,j+1} - u_{i+1,j-1} - u_{i-1,j+1} + u_{i-1,j-1} \right] + \mathcal{O}(\Delta x^2)$$

and we see that the (ij) point is not employed, i.e., there is no element on the diagonal corresponding to u_{ij} . This will lead to algebraic systems with loss of diagonal dominance and thus substantial computational complexity.



The C++ function example below is an implementation of the above cross-type molecule. Note that for writing the function, it is better to expand the symbolic operators first, which leads to an immediate translation into C++ array-indexing. Once again, the key thing to notice in this example is how easily the finite difference formulas translate into C++ code once you have a handle on how arrays work. In this example, we assume that we have already allocated double arrays for both u and u_xy , we merely need to index the arrays properly to obtain the desired result.

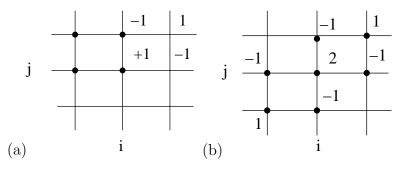


Figure 5.18: Stencils for forward (a) and symmetric (b) differentiation of a mixed derivative.

In order to restore diagonal dominance we proceed as follows. We first observe that in the above formula for the mixed derivative if we omit the average operator μ we loose an order of accuracy, i.e.,

$$(u_{xy})_{ij} = \frac{1}{4\Delta x \Delta y} (\mu_x \delta_x \delta_y^+) u_{ij} + \mathcal{O}(\Delta x^2, \Delta y).$$

Therefore, a first-order accurate stencil in both $\Delta x, \Delta y$ is:

$$(u_{xy})_{ij} = \frac{1}{\Delta x \Delta y} \delta_x^+ \delta_y^+ u_{ij} + \mathcal{O}(\Delta x, \Delta y).$$

This is a *forward differencing* for the mixed derivative, see figure 5.18(a). Similarly, we can construct a backward difference stencil. We can then combine the two stencils to produce a symmetric stencil, see figure 5.18(b), with a non-zero coefficient in the diagonal. This has the form

$$(u_{xy})_{ij} = \frac{1}{2\Delta x \Delta y} [\delta_x^+ \delta_y^+ + \delta_x^- \delta_y^-],$$

and this symmetric stencil is now of second-order on a uniform grid.

5.2 Explicit Time Discretizations

In this section we apply finite difference operations to discretize the time variable in timedependent differential equations. To this end, we consider the initial value problem (IVP)

$$\frac{dU}{dt} = \mathcal{F}(U, t), \tag{5.12}$$

with prescribed initial conditions $U(t = 0) = U_0$. If $\mathcal{F}(U)$ only, that is the right-hand-side does not depend explicitly on time, then this is an *autonomous* ordinary differential equation for which very efficient algorithms can be constructed. In order to analyze the stability of various discretizations it is convenient to employ the corresponding eigenvalue model problem

$$\frac{dU}{dt} = \lambda U, \quad \lambda \in \mathbf{C}, \quad Re(\lambda) < 0, \tag{5.13}$$

that is λ is a complex number with negative real part.

5.2.1 Multi-Step Schemes

In this discretization there are several time steps involved, and the discrete analog of equation (5.12) has the general form

$$\frac{U^{n+1} - U^{n-k}}{(k+1)\Delta t} = \mathcal{F}(t, U^{n+1}, U^n, U^{n-1}, \dots) \quad \text{for } k \ge 0,$$
(5.14)

where $U^n \equiv U(n\Delta t)$. An *implicit scheme* corresponds to right-and-side \mathcal{F} that depends on U^{n+1} , otherwise it is *explicit*. A multi-step scheme has \mathcal{F} depending on several previous time levels of the solution, i.e. U^n, U^{n-1}, U^{n-2} , etc. The simplest member of the class of discretization derived from equation (5.14) is the **Euler-forward**, which is one-sided backward finite difference, i.e.

$$\frac{U^{n+1} - U^n}{\Delta t} = \mathcal{F}(t, U^n) \Rightarrow U^{n+1} = U^n + \Delta t \mathcal{F}(t, U^n).$$

Another example is the **leap-frog scheme**, which is a two-step explicit scheme. For the model eigenvalue problem it has the form:

$$\frac{U^{n+1} - U^{n-1}}{2\Delta t} = \lambda U^n.$$

Clearly, it is of second-order accuracy since the first-derivative is represented by central differencing.

More generally, **multi-step schemes** can be either explicit or implicit. The general formula for a *k*-step scheme is:

$$\sum_{j=0}^{k} a_j U^{n+1-j} = \Delta t \sum_{j=0}^{k} \beta_j \mathcal{F}^{n+1-j},$$

where $\alpha_0 = 1$ for normalization, and $\beta_0 = 0$ corresponds to an explicit scheme. The physical analog is a "recursive digital filter."

Example: The Adams Family

The first member of the family is the Euler-forward scheme. The other two (explicit) members are:

- Second-order Adams-Bashforth: $\frac{U^{n+1}-U^n}{\Delta t} = \frac{3}{2}\mathcal{F}^n \frac{1}{2}\mathcal{F}^{n-1}$.
- Third-order Adams-Bashforth: $\frac{U^{n+1}-U^n}{\Delta t} = \frac{23}{12}\mathcal{F}^n \frac{16}{12}\mathcal{F}^{n-1} + \frac{5}{12}\mathcal{F}^{n-2}$.

Multi-steps schemes require lower order schemes in the first few steps; this may affect their accuracy. In the C++ code below we implement first-, second- and third-order Adams-Bashforth schemes. As mentioned previously, we are confronted with the decision between optimality and generality. In this case, we choose to be general; that is, we have created a function which uses a *switch* statement to interpret which order Adams-Bashforth scheme the user wants to employ.

One additional thing we would like to point out in this example is the use of the default within the *switch* statement. Notice in the example below that the default is Euler-forward, which is also the case one option. Instead of printing an error and terminating if someone were to ask for an Adams-Bashforth scheme which were of higher order than three, we default to first-order. These types of decisions are constantly being made by programmers: how will the function respond if given unexpected input? In this case, if order is not given as either one, two or three, then the action is still well defined: the function will "default" to first-order integration.

Software

O Putting it into Practice
Suite

Below we present a C++ function for accomplishing time integration using the Adams-Bashforth methods. This function takes as arguments the order, the old value, the time step and the saved right-hand-sides (stored in an array called *RHS*).

```
double AdamsBashforth(int order, double u_old,
                      double dt, double * RHS){
  double answer;
  switch(order){
  case 1: /* 1st Order Adams-Bashforth -- Euler-forward */
    answer = u_old + dt*RHS[0];
    break;
  case 2: /* 2nd Order Adams-Bashforth */
    answer = u_old + dt*(1.5*RHS[0] - 0.5*RHS[1]);
    break:
  case 3: /* 3rd Order Adams-Bashforth */
    answer = u_old + dt*((23./12.)*RHS[0] - (4./3.)*
             RHS[1] + (5./12.)*RHS[2]);
    break;
  default: /* default is Euler-forward */
    answer = u_old + dt*RHS[0];
  }
  return answer;
}
```

Remark 1: Observe that we use a *switch* statement to differentiate between the different cases. We have written the *switch* statement so that the default case is Euler-forward, which is first order.

Remark 2: Observe that this function can easily be modified to handle an array of points to integrate in time (such as would be used for time marching a finite difference scheme).

Key Concept

• As programmers, we are always confronted with the following balance: Generality versus Optimality.

5.2.2 Convergence: Consistency and Stability

There exists a connection between consistency and convergence for boundary value problems, as we have briefly discussed in the previous section. This condition, first proposed by Lax, is necessary but not sufficient for initial value problems, and we need to introduce the concept of stability, as we will do next. However, we first need to show that the explicit multistep schemes are *consistent*, and to this end we need to consider the *equivalent differential equation* for the initial value problem given by equation (5.12). For simplicity, let us consider the Euler-forward scheme. From Taylor's expansion we obtain:

$$\left(\frac{dU^n}{dt} - \mathcal{F}^n\right) = \Delta t \left(-\frac{1}{2}\frac{d^2U^n}{dt^2}\right) + \Delta t^2 \left(-\frac{1}{6}\frac{d^3U^n}{dt^3}\right) + \dots$$

Clearly, it satisfies the consistency requirement as the right-hand-side is zero as $\Delta t \to 0$, and thus we recover the original initial value problem. We can also obtain the accuracy of the method by examining the *truncation error*, which is the forcing term in the equivalent differential equation. From the leading error term in the above equation we see that the Euler-forward scheme is first-order.

Unlike boundary value problems (BVP) for which consistency is sufficient to prove convergence, in initial value problems (IVP) we also need to prove *stability*. This is explained graphically in figure 5.19. With time integration there may be an accumulation of error which will eventually lead to catastrophic instabilities!

Next, we introduce the concept of stability by defining the error as

$$\epsilon \equiv U_{ex}^n - U^n.$$

We define

• General stability if

$$|\epsilon^n| < f(t_n),$$

where $t_n = n\Delta t$. In other words, errors are bounded at a fixed time by a function that may depend on time! $(t_n = \text{fixed}, \Delta t \to 0, n \to \infty)$.

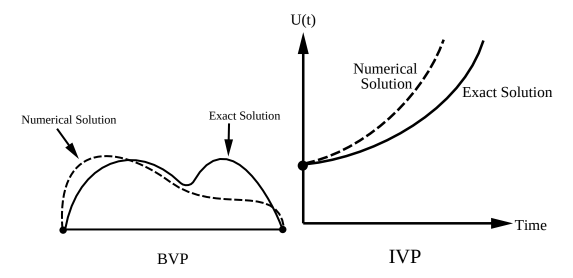


Figure 5.19: Comparison of boundary value (BVP) and initial value (IVP) problems.

• Absolute stability if

$$|\epsilon^{n+1}| \le |\epsilon^n|.$$

Here the equal sign implies *weak* stability whereas the non-equal sign implies *strong* stability. The interpretation here is that for absolute stability all components of error should be uniformly bounded.

Example: Let us examine the stability of the Euler-forward scheme for the eigenvalue model problem $\frac{dU}{dt} = \lambda U, \lambda < 0$.

Then, the error satisfies: $\epsilon^{n+1} = \epsilon^n (1 + \lambda \Delta t)$. But

$$|1 + \lambda \Delta t| < |1 - \lambda \Delta t| < e^{-\lambda \Delta t}$$

thus

$$|\epsilon^{(n+1)}| < e^{-\lambda \Delta t} |\epsilon^n| \le (e^{-\lambda \Delta t})^n |\epsilon^0| \Rightarrow |\epsilon| < e^{-\lambda (n\Delta t)} |\epsilon^1|.$$

According to the above definition for *general stability*, this scheme is then stable! However, consider the counter example for the IVP:

$$\frac{dU}{dt} = -U, \quad U(0) = 1$$

with $U_{ex}=e^{-t}$. Then, for $\Delta t=3$ we obtain $U^{n+1}=-2U^n$, and we produce the sequence:

$$(1, -2, 4, -8, 16...)$$

which obviously does not converge to e^{-t} for t = 12!

General stability is limited in practice, and instead we need to apply the concept of absolute stability. Proceeding as above we need to bound the error, so we impose

$$|1 + \lambda \Delta t| \le 1 \Rightarrow \Delta t \le -2/\lambda.$$

Notice that we have violated this condition in our example by taking $\Delta t = 3$.

The key points to remember are:

- General stability is useful for *short-time* integration only.
- Stability is the property of a difference equation and not of the differential equation.

Lax's Equivalence Theorem: If a (linear) difference equation is consistent with a differential equation and stable, its solution will converge as $\Delta t \to 0$ to that of the differential equation.

This condition is sufficient as well as necessary [77]. The equivalence theorem of Lax shows that in order to analyze an initial value problem, two tasks have to be performed:

- (1) Analysis of the consistency condition: This leads to determining the order of accuracy of the scheme and the corresponding truncation error.
- (2) Analysis of the stability condition: This leads to determining the frequency distribution of the error (e.g., eigenvalue-type analysis).

Remark: A stronger version of this theorem was developed by Dahlquist [20]. It applies to both linear and nonlinear initial value problems, and it states that a *multi-step* scheme is convergent *if and only if* it is consistent and stable. The Lax equivalence theorem, which was formulated for partial differential equations, requires linearity.

5.2.3 Stability and Characteristic Polynomials

A very effective method for studying the stability of time discretizations is to examine the properties of their corresponding characteristic polynomials. To this end, we propose a construction by associating two polynomials $\rho(z)$ and $\sigma(z)$ to the k-step scheme

$$\sum_{j=0}^{k} \alpha_j U^{n+1-j} = \Delta t \sum_{j=0}^{k} \beta_j \mathcal{F}^{n+1-j}.$$

For the left-hand-side we have

$$\rho(z) = \sum_{j=0}^{k} \alpha_j z^{k-j},$$

and for the right-hand-side we have

$$\sigma(z) = \sum_{j=0}^{k} \beta_j z^{k-j}.$$

The degree of the polynomial $\rho(z)$ is k, and the degree of the polynomial $\sigma(z) \leq k$, with the equal sign valid for *implicit* schemes (see chapter 6). We can normalize the coefficients by imposing that

$$\sum_{j=0}^{k} \beta_j = 1.$$

Examples:

- The *Leap-frog* scheme: $\rho(z) = \frac{z^2-1}{2}$; $\sigma(z) = z$, is explicit since $\sigma(z)$ is a linear polynomial but $\rho(z)$ is quadratic.
- The Crank-Nicolson scheme, which is an implicit scheme, see section 6.2, has: $\rho(z) = z 1$; $\sigma(z) = \frac{1}{2}(z + 1)$. Both polynomials are linear.

Based on the characteristic polynomials several properties of multi-step schemes can be deduced, and the interested reader should consult the book by Gear [42].

Rational Approximation Theorem: A multi-step scheme with $\sigma(1) \neq 0$ has order of accuracy p if and only if:

As $z \to 1$

$$\frac{\rho(z)}{\sigma(z)} = \log z + \mathcal{O}((z-1)^{p+1})$$

$$= [(z-1) - \frac{1}{2}(z-1)^2 + \frac{1}{3}(z-1)^3 - \dots]$$

$$+ \mathcal{O}((z-1)^{p+1}).$$

It is a *consistent* scheme if and only if

$$\begin{cases} \rho(1) = 0 & (\Rightarrow p \ge 0) \\ \rho'(1) = \sigma(1) & (\Rightarrow p \ge 1). \end{cases}$$

Example 1: The Crank-Nicolson is a consistent scheme since

$$\sigma(1) = \frac{1}{2}(z+1)|_{z=1} \neq 0, \rho(1) = (z-1) = 0, \rho'(1) = 1 = \sigma(1).$$

To examine its accuracy we consider the ratio:

$$\frac{\rho(z)}{\sigma(z)} = \frac{z-1}{\frac{1}{2}(z+1)} = \frac{z-1}{1+\frac{1}{2}(z-1)} = (z-1)\left[1 - \frac{z-1}{2} + \frac{(z-1)^2}{4} - \dots\right]$$
$$= (z-1) - \frac{(z-1)^2}{2} + \frac{(z-1)^3}{4} - \dots$$

Note that the last term is not equal to $\frac{(z-1)^3}{3}$ (as in the \log expansion), and thus it represents the truncation error. Therefore, we conclude that p=2 and the Crank-Nicolson scheme is second-order accurate.

Example 2: Let $\rho(z) = z^2 - z$ and k = 2, then

$$\sigma(z) = \frac{z^2 - z}{\log z} + \mathcal{O}((z - 1))^3) = \frac{z(z - 1)}{(z - 1) - \frac{1}{2}(z - 1)^2 + \frac{1}{3}(z - 1)^3} + \mathcal{O}((z - 1)^3)$$

$$= \frac{z}{1 - \frac{1}{2}(z - 1) + \frac{1}{3}(z - 1)^{2} \dots} + \mathcal{O}((z - 1)^{3})$$

$$= z \left[1 + \frac{1}{2}(z - 1) - \frac{1}{3}(z - 1)^{2} + \frac{1}{4}(z - 1)^{4} \right] + \mathcal{O}(\dots)$$

$$= \frac{5}{12}z^{2} + \frac{8}{12}z - \frac{1}{12} = \sigma(z).$$

Thus, we obtain the third-order (implicit) Adams-Moulton scheme (see also section 6.2).

Example 3: Consider a multi-step formula based purely on extrapolation of previous values, e.g.,

$$U^{n+1} = 2U^n - U^{n-1}$$

for which $\rho(z) = (z-1)^2$ and $\sigma(z) = 0$. We have that $\rho(1) = 0$ and $\rho'(1) = \sigma(1) = 0$. Notice that we need $\sigma(1) \neq 0$ to satisfy the rational approximation theorem. Obviously, such a formula cannot converge because we do not use at all any differential equation!

Root Condition for Stability: A multi-step scheme is stable if and only if all the roots of $\rho(z)$ satisfy $|z| \le 1$, and any root with |z| = 1 is simple.

Proof. The polynomial $\rho(z)$ has a total of k roots for which we can write (if they are distinct):

$$U^n = z^n$$
.

where on the left-hand-side n denotes superscript while on the right-hand-side it denotes an exponent. For stability we require that the entire sequence

$$\{U^n, n = 1, 2, \dots, \}$$

be bounded, and thus $|z| \leq 1$. However, for multiple roots, say of multiplicity m, we have the roots

$$U^n = nz^n, n^2z^n, \dots, n^{m-1}z^n.$$

In this case the requirement for boundness of the sequence U^n is |z| < 1.

An alternative proof can be formulated based on matrices as follows:

$$\begin{bmatrix} U^{n+2-k} \\ U^{n+3-k} \\ \vdots \\ U^{n+1} \end{bmatrix} = \begin{bmatrix} 0 & 1 & & \\ & 0 & 1 & \\ & & \ddots & \ddots & \\ & & & 1 \\ -\alpha_k & & -\alpha_2 & -\alpha_1 \end{bmatrix} \begin{bmatrix} U^{n+1-k} \\ U^{n+2-k} \\ \vdots \\ U^n \end{bmatrix}$$

or

$$\mathbf{U}^{n+1} = \mathbf{A} \mathbf{U}^n.$$

Notice that the eigenvalues of **A** are roots of the polynomial $\rho(z)$. Thus, for boundness of \mathbf{U}^{n+1} , we require that the eigenvalues of **A** be smaller than 1 and those equal to 1 be simple. A rigorous proof requires to transform **A** to its Jordan canonical form, e.g. see [42].

Example 1: Let us consider an Adams-Bashforth scheme of k-order:

$$\rho(z) = z^k - z^{k-1} = z^{k-1}(z-1)$$

with roots $\{1, 0, 0, \ldots\}$. Therefore, this scheme is stable.

Example 2: We can construct the most accurate two-step explicit scheme corresponding to p = 3 from

$$U^{n+1} = -4U^n + 5U^{n-1} + \Delta t(4\mathcal{F}^n + 2\mathcal{F}^{n-1}).$$

Therefore, $\rho(z) = z^2 + 4z - 5$ with roots $\{1, -5\}$, which according to the root condition it corresponds to an unstable scheme! Note that this scheme violates the **first Dahlquist Stability Barrier** theorem, see section 6.2.1, that states that the order of accuracy p of any explicit multi-step scheme (k steps) cannot exceed the number of steps, i.e. p < k.

Example 3: Similarly, we can construct a three-step scheme from

(2)
$$U^{n+1} = -\frac{3}{2}U^n + 3U^{n-1} - \frac{1}{2}U^{n-2} + 3\Delta t \cdot f^n$$

with corresponding polynomial

$$\rho(z) = 2z^3 + 3z^3 - 6z + 1 = (z - 1)(2z^2 + 5z - 1),$$

which has the roots 1.0, 0.186 and -2.686. Therefore, this scheme is unstable!

Example 4: Based on the root condition, we can also prove that all one-step methods of the form

$$U^{n+1} = U^n + \Delta t \mathcal{F}^n$$

are stable because $\rho(z) = z - 1$. Therefore, the root is 1 and simple, and thus the root condition for stability is satisfied.

Remark: We note here that in all the examples above stability was interpreted according to the general definition. In the following, we introduce the requirements for absolute stability. According to the **Second Dahlquist Stability Barrier** theorem, see section 6.2.1, an explicit multi-step scheme cannot be absolutely stable for arbitrary values of the time step Δt .

Stability Regions and Absolute Stability

Let us consider the linear ordinary differential equation

$$\frac{dU}{dt} = \lambda U + q(t), \quad \lambda \in \mathbf{C}, Re(\lambda) < 0.$$

The corresponding error, ϵ^n , at the time level $(n\Delta t)$ satisfies the equation

$$\sum_{j=0}^{k} (\alpha_j - \Delta t \lambda \beta_j) \epsilon^{k-j} = 0,$$

independently of the inhomogeneity q(t).

To compute the solution of the above difference equation, we introduce $\epsilon^n = z^n$ (where n on the left-hand-side denotes superscript but on the right-hand-side denotes an exponent), and thus:

$$\sum_{j=0}^{k} (\alpha_j - \Delta t \lambda \beta_j) z^{k-j} = 0.$$

Therefore, the z are roots of the polynomial

$$\Pi(z) \equiv \rho(z) - \lambda \Delta t \cdot \sigma(z) = 0,$$

which we denote as z_i .

If all the roots are distinct, then the general solution for the error ϵ^n is:

$$\epsilon^n = \sum_{j=1}^k c_j z_j^n,$$

(assuming k roots). If z_i is an m-fold root, then the term

$$(c_j + c_{j+1}n + c_{j+2}n^2 + \ldots + c_{j+m-1}n^{m-1})z_j^m$$

is present. Therefore, for any $|z_j| \ge 1$, disturbances are amplified irrespective of Δt , and the corresponding scheme is unstable.

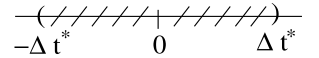


Figure 5.20: Sketch showing the neighborhood around 0.

In general, the roots depend on Δt , i.e., $z_j = z_j(\lambda \Delta t)$. Therefore, if

$$|z_j(0)| < 1 \Rightarrow \exists \Delta t^* : |z_j(\lambda \Delta t)| \le 1$$

in the neighborhood around 0 with radius Δt^* (see sketch in figure 5.20). Expanding $z_j(\lambda \Delta t)$ in a Taylor series around 0 we obtain:

$$z_j(\lambda \Delta t) = z_j(0) + \mathcal{O}(\Delta t \lambda).$$

Thus,

$$|z_i^n(\lambda \Delta t)| \le |z_i(0) + C\Delta t|^n \le |1 + C\Delta t|^n \le e^{Cn\Delta t} \le e^{Ct}$$

i.e., if $|z_j(0)| \leq 1$ then the roots of $\Pi(z)$ are less than 1, which implies stability. This explains also how the roots of $\rho(z) = \Pi(0)$ are related to stability of the scheme (see previous theorem on root condition). For *strong stability* we require that the roots of $\Pi(z)$ are ≤ 1 in absolute value, i.e., we have to find:

$$\lambda \Delta t : |z_i(\lambda \Delta t)| \le 1$$
,

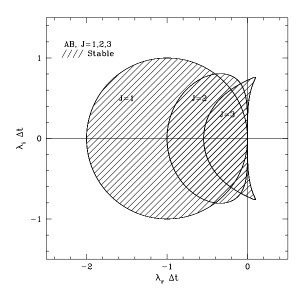


Figure 5.21: Stability diagrams for the first three members of the (explicit) Adams-Bashforth family.

and this constraint determines the *regions of absolute stability* of a time discretization scheme. We illustrate this in the following examples.

Example 1 - Euler-forward: We first construct the polynomial

$$\Pi_{EF} = (z-1) - \lambda \Delta t \cdot 1 = z - (1 + \lambda \Delta t),$$

and obtain the roots : $z = 1 + \lambda \Delta t$. We now require that

$$-1 \leq 1 + \lambda t \leq 1 \Rightarrow |\lambda \Delta t - (-1)| \leq 1 \Rightarrow$$

$$\Delta t \le -2/\lambda$$
, $\lambda \in \mathbf{C}$, $Re(\lambda) < 0$.

Note that for physical stability, we have also considered that $\lambda \Delta t \leq 0$ for the exact modes to decay. The region of stability in the complex plane is shown in figure 5.21 for J=1. Note that it touches the imaginary axis (which represents advection) only at one point, and thus this scheme is only marginally stable for advection problems.

Example 2 - Leap-frog: We construct the polynomial

$$\Pi_{LF} = z^2 - 2\lambda \Delta tz - 1,$$

and obtain the two roots whose product is $z_1 \cdot z_2 = 1$. Therefore, we have that

$$|z_1| \le 1, |z_2| \ge 1.$$

For stability the only possibility is $|z_1| = |z_2| = 1$, and thus the region of stability is the region on the imaginary axis between (-i, i). Therefore, the leap-frog scheme is a good candidate for

problems with purely imaginary eigenvalues, e.g., advection in a periodic domain, discretized by a non-dissipative scheme. Even slight dissipation, however, will introduce real eigenvalues which will lead to instabilities.

In general, to obtain the stability regions of multi-step schemes we construct the curve

$$\lambda \Delta t = \frac{\rho(z)}{\sigma(z)},$$

with $z = e^{i\theta}$, $\theta \in [0, 2\pi]$ in the complex $\lambda \Delta t$ plane. Using this approach we have obtained the stability regions for the (explicit) Adams family and the results are shown in figure 5.21. Note that the third-order scheme has stability region with a substantial intersection with the imaginary (advection) axis.

5.2.4 Runge-Kutta Methods

Unlike the Adams family, Runge-Kutta methods, developed by Runge (1895), Heun (1900) and Kutta (1901), are a single-step methods, and in particular they are multi-stage methods. The basic idea is to create a weighted sum of corrections ΔU^k to the solution at several stages within the same time step, i.e.,

$$U^{n+1} = U^n + C_1 \Delta U^1 + C_2 \Delta U^2 + C_3 \Delta U^3 + \dots$$

The coefficients C_k are determined by matching the above expansion with the corresponding expansion by Taylor series. Other parameters are also introduced to accommodate different stages, e.g.,

$$\Delta U^{1} = \Delta t \mathcal{F}^{n}(t^{n}, U^{n}) \quad (Euler-forward)$$

$$\Delta U^{2} = \Delta t \mathcal{F}(t^{n} + \alpha \Delta t, U^{n} + \beta \Delta U^{1})$$

:

Runge-Kutta methods tend to be *non-unique* due to the large number of parameters introduced. In modern versions the objective has been to minimize storage levels as shown in the following examples. For hyperbolic systems it is important to produce Runge-Kutta schemes which are total variation diminishing (TVD) or total variation bounded (TVB) – see for example [50]. TVD and TVB schemes do not allow any spurious oscillation to appear in the numerical solution, and this is a highly desirable property. In the following, we present representative algorithms of some of the most popular versions.

(1) Second-Order (RK2) with Two Levels of Storage

Set:

$$X = U^n$$
$$Y = \mathcal{F}(U, t^n)$$

Compute:

$$X = X + \alpha \cdot \Delta t Y$$

$$Y = aY + \mathcal{F}(X, t^n + \alpha \Delta t)$$

Update:

$$U^{n+1} = X + \frac{\Delta t}{2\alpha} Y$$

where $a = -1 + 2\alpha - \alpha^2$

Note that for $\alpha = 1/2$ we obtain the modified Euler method and for $\alpha = 1$ we obtain the classical Heun method.

(2) Classical Fourth-Order (RK4)

Compute:

$$X_1 = \mathcal{F}(U^n, t^n)$$

$$X_2 = \mathcal{F}(U^n + \frac{1}{2}X_1\Delta t, t^n + \frac{1}{2}\Delta t)$$

$$X_3 = \mathcal{F}(U^n + \frac{1}{2}X_2\Delta t, t^n + \frac{1}{2}\Delta t)$$

$$X_4 = \mathcal{F}(U^n + X_3\Delta t, t^n + \Delta t)$$

Update:

$$U^{n+1} = U^n + \frac{1}{6}\Delta t[X_1 + 2X_2 + 2X_3 + X_4]$$

An alternative implementation requires only three levels of storage, as follows (due to

Blum [9]).

Stage I:
$$X = U^n; Y = X; Z = \mathcal{F}(X, t^n)$$

Stage II:
$$X = X + \frac{1}{2}\Delta tZ; Y = Z; Z = \mathcal{F}(X, t^n + \frac{1}{2}\Delta t)$$

Stage III:
$$X=X+\frac{1}{2}\Delta t(Z-Y); Y=\frac{1}{6}Y; Z=\mathcal{F}(X,t^n+\frac{1}{2}\Delta t)$$
$$-\frac{1}{2}Z$$

Stage IV:
$$X = X + \Delta t Z$$
; $Y = Y - Z$; $Z = \mathcal{F}(X, t^n + \Delta t) + 2Z$

Update:

$$U^{n+1} = X + \Delta t \left(Y + \frac{1}{6}Z \right)$$

We provide below a C++ implementation of these algorithms. Notice that we are reusing two previously discussed concepts: that of passing a function to a function, and

Software



Putting it into Practice

that of using static declaration of an array. In this function, we pass the *rkfunc* function (which is the right-hand-side of our ODE) into the Runge-Kutta function so that it can be evaluated (in this case, being evaluated four times). We store the intermediate stages within the statically allocated array *hold*, which we explicitly declare to contain four values since there are four stages to this method.

One key issue to point out is that it would be perfectly legitimate for us to dynamically allocate the space for the array *hold*. However, in this case, it is more efficient to allocate the space for *hold* statically at the beginning of the function. A convenience of static allocation in this case is that deallocation is handled automatically! Recall that there is no need to explicitly deallocate the static declaration. All of this is possible because we know ahead of time the number of items that *hold* will need to contain (in this case, four items), and hence we can explicitly declare how much memory we will need to properly implement this function.

(3) Autonomous ODE

It is defined by the right-hand-side being

$$\mathcal{F} = \mathcal{F}(U)$$

only.

The following version is due to Jameson et al. [60], and it produces a Runge-Kutta method of order p with only 3 levels of storage.

Set:
$$X = U^n$$

For
$$k = p, 1, -1$$

$$X = U^n + \frac{1}{k}\Delta t \mathcal{F}(X)$$

End For

Update: $U^{n+1} = X$

Software



Putting it into Practice

The algorithm presented above is implemented in the example code below. Here we show how this method easily translates into a C++ function. We want to draw your attention to one particular item within this function: notice the looping index. In previous examples, we have started at a value for the incrementor, and have consistently added to that value. In this example, we decrement! Careful attention should be placed on the stopping condition. Notice that we want the value of k to start at order and to decrement to k = 1 inclusive.

WARNING

Programmer Beware!

• Think carefully about your loop ending condition!

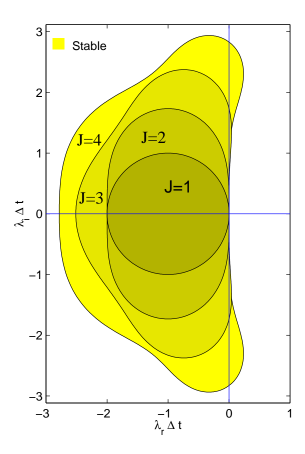


Figure 5.22: Stability diagrams for the Runge-Kutta methods of order one to four.

5.2.5 Stability of Runge-Kutta Methods

As an example let us consider the equation

$$\frac{dU}{dt} = \lambda U$$

and examine the stability of RK4. We compute:

$$X_1 = \lambda U^n$$

$$X_2 = \lambda (U^n + \frac{1}{2}\lambda U^n \Delta t)$$

$$\vdots$$

and therefore

$$U^{n+1} = U^n + \frac{1}{6}\Delta t[X_1 + X_2 + 2X_3 + X_4]$$

or

$$U^{n+1} = U^n \left[1 + \lambda \Delta t + \frac{\lambda^2 \Delta t^2}{2} + \frac{\lambda^3 \Delta t^3}{6} + \frac{\lambda^4 \Delta t^4}{24} \right]$$

The growth factor is then

$$G = \left[1 + \ldots + \frac{\lambda^4 \Delta t^4}{24}\right]$$

and we require

$$|G| \leq 1$$
.

By setting $\mu \equiv \lambda \Delta t$, we solve:

$$1 + \mu + \frac{\mu^2}{2} + \frac{\mu^3}{6} + \frac{\mu^4}{24} = e^{i\theta}, \ \theta \in [0, 2\pi]$$

and determine (numerically) $\mu(\theta)$. In figure 5.22 we plot the stability regions of the Runge-Kutta methods. We notice that unlike the multi-step methods, in Runge-Kutta methods the stability regions increase with order!

5.3 Homework Problems

- 1. Use the method of undetermined coefficients to obtain a third-order finite difference approximation for an one-sided (upwind-type) first and second (one-dimensional) derivatives.
- 2. Use the half-central and the central difference operators to derive approximations of second- and fourth-order of accuracy approximations for the fourth (one-dimensional) derivative.
- 3. Finite differencing as we have presented it can be formulated as a matrix multiplication operation. Each explicit finite difference operator yields a differentiation matrix which can be formed during a pre-processing stage before computations are accomplished. Each time differentiation is needed, pre-multiplication by the differentiation matrix yields an approximation of the derivative.
 - (a) Formulate an one-dimensional second-order first derivative differentiation matrix and an one-dimensional second-order second derivative differentiation matrix assuming a periodic domain.
 - (b) What is the bandwidth of the matrix?
 - (c) Write C++ functions which generate these matrices (individual functions per matrix).
 - (d) Write a C++ function which implements differentiation by matrix multiplication. The function should take as input the size of the differentiation matrix, the differentiation matrix (as a double ** array), an array containing the values on which to operate and an array for storing the result.
 - (e) Verify your routines by comparing results with the functions $SO_FirstDeriv_1Dper$ and $SO_SecondDeriv_1Dper$.

 Demonstrate second-order convergence as shown in the text.
- 4. Write a C++ function to implement a five-point stencil for explicit discretization of the first derivative. Assume that the domain is periodic. Verify that you obtain fourth-order convergence.
- 5. Write an MPI program using $MPI_Sendrecv$ which implements a parallel version of the previous function just described. For number of processors P=2,4, and 6, verify that you obtain the same convergence rate as the previous function.
- 6. Write an MPI program using $MPI_Sendrecv_replace$ which implements a parallel version of the previous function just described. Compared with the previous program, how many MPI function calls are necessary? For number of processors P=2,4, and 6, verify that you obtain the same convergence rate as the previous program.
- 7. Obtain numerical solutions to the the ordinary differential equation

$$\frac{dy}{dt} = 2\sqrt{y}$$

in the interval [0, 1], with initial conditions:

$$y = 0.1$$
; $y = 0.01$; $y = 0.001$; $y = 0.0001$; $y = 0$.

Compare the Runge-Kutta scheme of second-order to the Adams-Bashforth scheme of second-order. What do you observe?

8. (Period doubling and chaos)

Sometimes numerical instabilities appear as numerical chaos, similar to the physical response that we encounter in dynamical systems. This is demonstrated here with the ordinary differential equation

$$\frac{dy}{dt} = 10y(1-y), \quad y(0) = 0.1.$$

- (a) Solve analytically the above problem and show that $y(t) \to 1$ as $t \to \infty$.
- (b) Show that using the Euler-forward method with step side $h = \Delta t$ we obtain

$$y^{n+1} = (1+10h)y^n - 10h(y^n)^2, \quad y^0 = 0.1.$$

- (c) For h = 0.18, 0.23, 0.25 and 0.3 show that the first 40 iterations of the Euler-forward scheme appear to: converge to 1; jump between 1.18 and 0.69; jump between 1.23, 0.54, 1.16 and 0.70; and display no discernible pattern, respectively for all cases. Produce a plot y^n versus h for n = 1001 to 2000 and h = 0.1 to 1.
- (d) Approximate now the solution with the fourth-order Runge-Kutta scheme and compute the first 60 iterations using h = 0.3. Repeat with h = 0.325 and h = 0.35. Which values of h, if any, give the correct approximation to the solution and why?

Note: The transitions from convergence to jumping between two numbers, to four numbers, and so on, are called *period doubling* while the phenomenon exhibited when h = 0.3 is *chaos*!

- 9. Obtain the stability curves of figures 5.21 and 5.22 for the Adams-Bashforth and the Runge-Kutta methods, respectively.
- 10. (Evolution of an ecosystem)

In this problem we model and solve competition between different species in an attempt to quantify Darwin's thoughts! Vito Volterra, an Italian mathematician, was the first to do this in the 1920s.

Consider an ecological system that contains a predator and a prey species. Prey, whose concentration is denoted by x, multiplies autonomously but is consumed by the predator whose concentration is denoted by y. The evolution of the population of the prey follows the ODE

$$\frac{dx}{dt} = ax - Axy, (5.15)$$

where a is positive birth rate constant (determined by the societal norms), and A is another positive constant that expresses how often predator catches prey. The evolution of the predator, on the other hand, follows the differential equation.

$$\frac{dy}{dt} = -by + A\epsilon xy,\tag{5.16}$$

where b is a death constant due to starvation, and ϵ is a share constant that indicates how many individual predators are necessary to eat up one individual prey.

The two equations above comprise the Lotka-Volterra system. There are two obvious steady state solutions (fixed points), denoted by (x^*, y^*) . The trivial one $(x^*, y^*) = (0, 0)$, and a nontrivial one, which is found by setting the right-hand-side of the above equations equal to zero and solving the resulting system of nonlinear algebraic equations for x and y.

Consider a system with $\epsilon=1,\ a=0.400,\ b=0.450,\ A=0.50$. Calculate the nontrivial steady concentrations (x^*,y^*) , and then integrate the Lotka-Volterra system using the *explicit third-order Adams-Bashforth method* subject to the initial conditions $x(t=0)=0.10;\ y(t=0)=0.20$. Carry the integration for a long enough time so that you can assess the asymptotic behavior of the system at very long times. Plot x(t) versus y(t) and discuss the trajectories of this solution as $t\to\infty$.

11. (Stochastic ODE)

Consider the first-order linear ODE:

$$\frac{dy}{dt} = -ky \quad \text{with } y(t=0) = y_0 \text{ and } t \in [0,T], \qquad (5.17)$$

where k is a stochastic process of second order such that:

$$k = \overline{k} + \nu (t, \omega). \tag{5.18}$$

Here \overline{k} is the mean value of k and $\nu\left(t,\omega\right)$ represents a random variable depending on time and random space. We assume that the probability distribution function (PDF) remains the same for all k's and takes the form of a Gaussian distribution with constant variance. The PDF of k at time t is:

$$f(k(t)) = \frac{1}{\sigma\sqrt{2\pi}}e^{-\frac{1}{2}\frac{\left(k(t)-\overline{k}\right)^2}{\sigma^2}},$$
(5.19)

where \overline{k} is the mean value of k and σ^2 is the variance of k.

Random values of k, generated in time every Δt_s , can be mutually independent, partially correlated or fully correlated. To this end, we consider the following cases:

Case 1: The PDF of the solution for the mutually independent case takes the form:

$$f(y(t)) = \frac{1}{\sigma u \sqrt{2\pi t \Delta t_s}} e^{-\frac{1}{2} \frac{\left(\ln \frac{y}{y_0} - \overline{k}t\right)^2}{\sigma^2 t \Delta t_s}}$$

$$(5.20)$$

Case 2: The PDF of the solution for the partially correlated case takes the form:

$$f(y(t)) = \frac{1}{\sigma \Delta t_s S y \sqrt{2\pi}} e^{-\frac{1}{2} \frac{\left(\ln \frac{y}{y_0} - \overline{k}t\right)^2}{(\sigma \Delta t_s S)^2}}, \quad \Delta t_s \ll T$$
 (5.21)

and

$$S = \left(N\frac{(1+C)}{(1-C)} - 2C\frac{(1-C^N)}{(1-C)^2}\right)^{\frac{1}{2}}$$
 (5.22)

where $C = e^{-\frac{\Delta t_s}{A}}$, $N = \frac{T}{\Delta t_s}$ and A is the correlation length of the random process.

Case 3: In the fully correlated case, the random process k becomes a random variable and the PDF of the solution takes the form:

$$f(y(t)) = \frac{1}{\sigma t y \sqrt{2\pi}} e^{-\frac{1}{2} \frac{\left(\ln \frac{y}{y_0} - \overline{k}t\right)^2}{(\sigma t)^2}}$$

$$(5.23)$$

An easy way to write the *first moment* (i.e., mean) in a general form for all cases is to express it in a logarithmic form:

$$\ln\left(\frac{E(y)}{y_0}\right) = -\overline{k}t + \frac{1}{2}\sigma^2\Gamma\tag{5.24}$$

where $\Gamma = t\Delta t_s$ for Case 1, $\Gamma = t^2$ for Case 3 and $\Gamma = \Delta t_s \left(t \frac{(1+C)}{(1-C)} - 2C\Delta t_s \frac{(1-C^N)}{(1-C)^2}\right)$ for Case 2.

Second moment (i.e., variance) for all cases can be obtained from the expressions of the mean.

$$E(y^2) = (e^{\sigma^2 \Gamma} - 1) E^2(y)$$
(5.25)

where $\Gamma = t\Delta t_s$ for Case 1, $\Gamma = t^2$ for Case 3 and $\Gamma = t\Delta t_s \left(t\frac{(1+C)}{(1-C)} - 2C\Delta t_s \frac{(1-C^N)}{(1-C)^2}\right)$ for Case 2.

- (a) Use a multi-step or a multi-stage ODE solver of various orders to obtain solutions and corresponding errors for the *mean* and *variance* response corresponding to $\bar{k} = 0; \sigma = 1$, and final time of integration T = 1. To do this you need to follow a **Monte Carlo** approach where $\nu(t,\omega)$ is obtained from a Gaussian PDF corresponding to $\sigma = 1$. The correlation length for Case 2 is assumed to be A = 0.1.
- (b) Does the accuracy of the numerical solutions you obtained increases with the formal solver of the time-stepping scheme you used as expected? Explain.
- 12. We presented in the text a function for time marching using Adams-Bashforth one single ODE. Modify the function to handle a system of ODEs.

- (a) What changes to the arguments must occur?
- (b) Which is more efficient, looping over the switch or placing loops within the switch statement? Verify your answer with an example and corresponding with timings.

Chapter 6

Implicit Discretizations

In this chapter we consider implicit discretizations of space- and time-derivatives. Unlike the explicit discretizations presented in the previous chapter, here we express a derivative at one grid point in terms of function values as well as derivative values at adjacent grid points (spatial discretization) or in terms of previous and current time levels (temporal discretization). This, in turn, implies that there is implicit coupling, and thus matrix inversion is required to obtain the solution.

The material of this chapter serves to introduce solutions of tridiagonal systems and correspondingly parallel computing of sparse linear systems using MPI. We also introduce two new MPI functions: $MPI_Barrier$, used for synchronizing processes, and MPI_Wtime , used for obtaining the wallclock timing information.

6.1 Implicit Space Discretizations

The discretizations we present here are appropriate for any order spatial derivative involved in a partial differential equation, but they are particularly useful when high-order accuracy and locality of data is sought. The explicit finite differences could also lead to high accuracy but at the expense of long stencils, and this, in turn, implies coupling involving many grid points and consequently a substantial communications overhead. On the contrary, the implicit finite differences employ very compact stencils and guarantee locality, which is the key to success of any parallel implementation. We only consider discretizations on uniform (i.e., equidistant) grids, as we assume that a mapping of the form presented in the previous chapter is always available to transform a non-uniform grid to a uniform one. We also present discretizations only for one-dimensional grids since multi-dimensional discretizations are accomplished using directional splitting, as before.

6.1.1 Difference Operators

We can employ the same difference operators defined in the previous chapter to obtain compact formulas for first- and higher order derivatives.

First Derivative

We begin with the expansion

$$\Delta xD = \mu(\delta - \frac{\delta^3}{3!} + \frac{1^2 2^2}{5!} \delta^5 - \dots),$$

and for fourth-order accuracy we truncate as follows

$$\Delta xD = \mu \delta (1 - \frac{\delta^2}{6}) + \mathcal{O}(\Delta x^5).$$

What makes the implicit approach different than the explicit one is the treatment of the operator in parentheses for the first term on the right-hand-side. To this end, we approximate it as a geometric series expansion employing the so-called *Padé approximation* to obtain

$$\Delta x D \approx \frac{\mu \delta}{1 + \frac{\delta^2}{6}} + \mathcal{O}(\Delta x^5),$$

which we re-write as

$$(1 + \delta^2/6)D = \frac{\mu\delta}{\Delta x} + \mathcal{O}(\Delta x^4). \tag{6.1}$$

We can further simplify the term on the right-hand-side

$$\mu \delta u_i = \mu [u_{i+1/2} - u_{i-1/2}] = \frac{1}{2} [u_{i+1} + u_i - u_i - u_{i-1}],$$

and therefore the right-hand-side of equation (6.1) is equal to

RHS =
$$\frac{1}{2} \frac{u_{i+1} - u_{i-1}}{\Delta x} + \mathcal{O}(\Delta x^4)$$
.

Similarly, we can work on the left-hand-side of equation (6.1):

$$\left(1 + \frac{\delta^2}{6}\right) Du_i = \left[(u_x)_i + \frac{1}{6} (E + E^{-1} - 2)(u_x)_i \right]
= (u_x)_i + \frac{1}{6} \left[(u_x)_{i+1} + (u_x)_{i-1} - 2(u_x)_i \right]
= \frac{1}{6} \left[(u_x)_{i+1} + 4(u_x)_i + (u_x)_{i-1} \right].$$

Upon substitution in equation (6.1), we obtain

$$\frac{1}{6}[(u_x)_{i+1} + 4(u_x)_i + (u_x)_{i-1}] = \underbrace{\frac{u_{i+1} - u_{i-1}}{2\Delta x}}_{\text{2nd-order}} + \mathcal{O}(\Delta x^4). \tag{6.2}$$

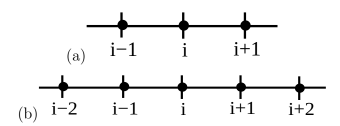


Figure 6.1: (a) Three-point stencil for implicit discretization of the first derivative with fourth-order accuracy. (b) Corresponding five-point stencil for explicit discretization.

Equation (6.2) indicates that we need to employ a three-point stencil (figure 6.1(a)) to discretize the first derivative with fourth-order accuracy. In contrast to obtain fourth-order accuracy with explicit discretization we need to employ a five-point stencil as follows (see figure 6.1(b))

$$(u_x)_i = \frac{-u_{i+2} + 8u_{i+1} - 8u_{i-1} + u_{i-2}}{12\Delta x} + \frac{\Delta x^4}{30} \frac{\partial^5 u}{\partial x^5}.$$

The graphical comparison of figure 6.1 of the two stencils involved in the implicit and explicit discretization indicates the locality of the computations in implicit approaches.

This derivation of the first derivative on a three-point stencil can be represented by the general scheme

$$\beta(u_x)_{i-1} + \alpha(u_x)_i + \beta(u_x)_{i+1} = \frac{u_{i+1} - u_{i-1}}{2\Delta x} + \mathcal{O}(\Delta x^p)$$

and can be extended to a longer stencil to match any order (p). In other words, on the left-hand-side we have a linear symmetric coupling of the derivative approximated at the points of the grid and on the right-hand-side we have a lower order (here second-order) approximation. In the next section we will discuss methods for computing the weights (coefficients α, β and γ) more systematically. Notice that equation (6.2) is implicit and requires the solution of a tridiagonal linear system.

One of the useful properties of implicit discretizations as we have discussed above is its *compactness*. This is demonstrated clearly with the **two-point stencil** for discretizing the first derivative. We begin again with

$$\Delta xD = \ln E = \ln(1 + \delta^+)$$

or

$$\Delta xD = \delta^{+} - \frac{\delta^{+2}}{2} + \underbrace{\frac{\delta^{+3}}{3} - \frac{\delta^{+4}}{4}}_{\mathcal{O}(\Delta x^{3})} + \dots$$

$$= \delta^{+} \left(1 - \frac{\delta^{+}}{2}\right) + \mathcal{O}(\Delta x^{3})$$

$$= \delta^{+} \frac{1}{1 + \frac{\delta^{+}}{2}} + \mathcal{O}(\Delta x^{3}),$$

where in the last equation we employ the geometric series approximation. We then proceed as follows

$$\left(1 + \frac{\delta^{+}}{2}\right)D = \frac{\delta^{+}}{\Delta x} + \mathcal{O}(\Delta x^{2}),$$

or

$$\{(u_x)_i + \frac{1}{2}[(u_x)_{i+1} - (u_x)_i]\} = \frac{u_{i+1} - u_i}{\Delta x} + \mathcal{O}(\Delta x^2),$$

which leads to a second-order formula for the first derivative, i.e.,

$$\frac{1}{2}[(u_x)_{i+1} + (u_x)_i] = \frac{u_{i+1} - u_i}{\Delta x} + \mathcal{O}(\Delta x^2).$$
(6.3)

This formula is also implicit but one-sided, and it involves the solution of a linear bidiagonal system.

Second Derivative

The method to compute second or higher order derivatives is similar as above. The starting point here is to employ the appropriate expansion given by the expansion of $D^n u_i$, see equations (5.2,5.4,5.5), and subsequently to use Padé approximation in the truncation of the series. For example, we can apply this approach to obtain a fourth-order accuracy approximation for the second derivative using central differencing, as follows

$$(u_{xx})_{i} = \frac{1}{\Delta x^{2}} \left[\delta^{2} - \frac{\delta^{4}}{12} + \frac{\delta^{6}}{90} - \frac{\delta^{8}}{560} + \dots \right] u_{i}$$

$$= \frac{\delta^{2}}{\Delta x^{2}} \left[1 - \frac{\delta^{2}}{12} \right] u_{i} + \mathcal{O}(\Delta x^{4})$$

$$= \frac{1}{\Delta x^{2}} \frac{\delta^{2} u_{i}}{1 + \delta^{2}/12} + \mathcal{O}(\Delta x^{4}).$$

From this we obtain

$$\left(1 + \frac{\delta^2}{12}\right)(u_{xx})_i = \frac{1}{\Delta x^2}\delta^2 u_i + \mathcal{O}(\Delta x^4),$$

which leads to the formula

$$\frac{1}{12}[(u_{xx})_{i+1} + 10(u_{xx})_i + (u_{xx})_{i-1}] = \frac{1}{\Delta x^2}[u_{i+1} - 2u_i + u_{i-1}] + \mathcal{O}(\Delta x^4). \tag{6.4}$$

We thus obtained fourth-order accuracy on the three-point stencil unlike the explicit discretization which leads to second-order accuracy on the same stencil. The price to pay here is that we need to obtain the derivative values at the grid points by solving the tridiagonal system

$$\frac{1}{12} \begin{bmatrix} \ddots & \ddots & O \\ 1 & 10 & 1 \\ O & \ddots & \ddots \end{bmatrix} [(u_{xx})_i]$$

with a known right-hand-side given by the function values at the grid points.

6.1.2 Method of Undetermined Coefficients

We now present a more general approach to constructing compact implicit finite difference schemes on uniform grids for first-, second-, third-, and fourth-order derivatives. It is based on the method of undetermined coefficients that we first encountered in the previous chapter, see section 5.1. However, unlike the explicit constructions, in the implicit approach we employ two different grids. Here, we follow the derivations of Lele [68].

First Derivative

5/7 Stencil:

Figure 6.2: Two grids employed for the implicit discretization of the first derivative.

We consider two grids and appropriate corresponding stencils for the discretization expressed by the following equation

$$\beta(u_x)_{i-2} + \alpha(u_x)_{i-1} + (u_x)_i + \alpha(u_x)_{i+1} + \beta(u_x)_{i+2}$$

$$= c\frac{u_{i+3} - u_{i-3}}{6\Delta x} + b\frac{u_{i+2} - u_{i-2}}{4\Delta x} + a\frac{u_{i+1} - u_{i-1}}{2\Delta x}.$$

The key idea here is to approximate the left-hand-side of this equation on the grid and corresponding five-point stencil shown in figure 6.2 (upper) and the right-hand-side on the seven-point stencil of figure 6.2 (lower). In particular, on the former we assign derivative values at the grid points whereas on the latter we form explicit central differences on substencils formed by the entire stencil. These explicit differences are of lower order (here second-order) accuracy.

The unknowns on the above equation are the weight coefficients on both sides, i.e., a total of five unknowns

$$\{\alpha, \beta, c, b, a\}.$$

Note that if $\beta=0$ then the computational complexity of obtaining the derivative values at the grid points is equivalent to solving a tridiagonal system, whereas if $\beta \neq 0$ then we have to solve a pentadiagonal system. Therefore, β defines the implicit stencil related to the right-hand-side. Similarly, the extent of the right-hand-side stencil is dictated by the coefficients a, b and c, and this selection is important for parallel computations as it dictates locality of data.

The method to obtain these unknown coefficients is similar to the one presented in the previous chapter, and it based on matching the Taylor expansions for $(u_x)_{i\pm 1}, u_{i\pm 1}$, etc. The difference, however, is that now we need to expand both the function as well as the first derivative around the central point (i). The first unmatched coefficient will give the truncation error.

Based on this matching we obtain the following constraints:

$$a + b + c = 1 + 2\alpha + 2\beta, \ \mathcal{O}(\Delta x^{2})$$

$$a + 2^{2}b + 3^{2}c = 2\frac{3!}{2!}(\alpha + 2^{2}\beta), \ \mathcal{O}(\Delta x^{4})$$

$$a + 2^{4}b + 3^{4}c = 2\frac{5!}{4!}(\alpha + 2^{4}\beta), \ \mathcal{O}(\Delta x^{6})$$

$$a + 2^{6}b + 3^{6}c = 2\frac{7!}{6!}(\alpha + 2^{6}\beta), \ \mathcal{O}(\Delta x^{8})$$

$$\vdots$$

We first consider **Tri-Diagonal Schemes** corresponding to $(\beta = 0)$ and requiring (5N) operations, where N is the number of the grid points in the grid. In such triagonal schemes we can have either 3/5 stencils or 3/7 stencils corresponding to c = 0 and $c \neq 0$, respectively. Here the first number (3) refers to the bandwidth of the stencil of the first grid, and the second number (5 or 7) to the bandwidth of the stencil of the second grid.

In the 3/5 stencil, see figure 6.3, we employ only two equations, and thus the truncation error is $\mathcal{O}(\Delta x^4)$. We also have that

$$c = 0 \Rightarrow a = \frac{2}{3}(\alpha + 2), \quad b = \frac{1}{3}(4\alpha - 1).$$

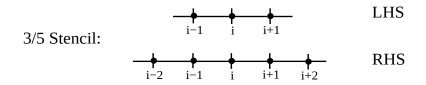


Figure 6.3: The 3/5 dual stencil for discretizing the first derivative.

Specifically, the truncation error is

$$T'_{3/5} = \frac{4}{5!} (3\alpha - 1) \Delta x^4 \left(\frac{\partial^5 u}{\partial x^5} \right)_i,$$

which is a function of the free parameter α that can be chosen arbitrarily.

Therefore, we can construct an α -family of schemes, with some typical members of the α -family corresponding to:

- $\alpha = 1/4 \Rightarrow b = 0$, and therefore this is a 3/3 stencil, the most compact scheme of the family. It is often called the *Padé scheme*, and it is of $\mathcal{O}(\Delta x^4)$ accuracy although it uses an identical three-point stencil as its explicit counterpart of order $\mathcal{O}(\Delta x^2)$.
- $\alpha = 1/3$. This choice leads to cancellation of the leading term in the truncation error, and thus it produces an $\mathcal{O}(\Delta x^6)$ scheme.
- $\alpha \to 0$. Here we recover the explicit central difference of $\mathcal{O}(\Delta x^2)$.

To construct a 3/7 stencil we employ three constraints and obtain schemes of (at least) sixth-order accuracy. The corresponding stencils are shown in figure 6.4.

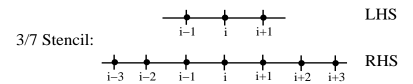


Figure 6.4: The 3/7 dual stencil for discretizing the first derivative.

The corresponding coefficients are obtained from

$$a = \frac{1}{6}(\alpha + 9);$$
 $b = \frac{1}{15}(32\alpha - 9);$ $c = \frac{-3\alpha + 1}{10},$

so again we have an α -family of schemes. We can obtain eighth-order accuracy, i.e., $\mathcal{O}(\Delta x^8)$, by setting $\alpha = 3/8$ which zeroes out the leading term in the truncation error. This scheme provides the highest order of the α -family.

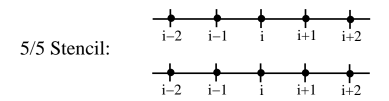
Next, we consider **Penta-Diagonal Schemes** corresponding to $(\beta \neq 0)$ and requiring (11N) operations, where N is the total number of grid points. This reflects the computational complexity of inverting a penta-diagonal matrix using Gaussian elimination, see section 9.1.

A 5/5 stencil, see figure 6.5, can be obtained by setting c = 0 with corresponding sixth-order accuracy. All coefficients are then defined in terms of α as follows:

$$a = \frac{2}{9}(8 - 3\alpha); \quad \beta = \frac{-1 + 3\alpha}{12}; \quad b = \frac{-17 + 57\alpha}{18},$$

and the corresponding truncation error is

$$T'_{5/5} = \frac{4}{7!} (9\alpha - 4)\Delta x^6 \left(\frac{\partial^7 u}{\partial x^7}\right).$$



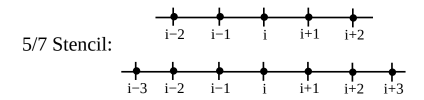


Figure 6.5: Stencils for penta-diagonal schemes in approximating the first derivative.

Similarly, on a 5/7 stencil, see figure 6.5, where $c \neq 0$, we can obtain eighth-order accuracy from the set of coefficients

$$a = \frac{12 - 7\alpha}{6}$$
; $b = \frac{568\alpha - 183}{150}$; $\beta = \frac{-3 + 8\alpha}{20}$; $c = \frac{9\alpha - 4}{50}$.

We can even obtain *tenth-order* accuracy for the proper choice of α :

$$T'_{5/7} = \frac{144}{9!} (2\alpha - 1) \Delta x^8 \frac{\partial^9 u}{\partial x^9},$$

if we set $\alpha = 1/2$. This is the highest possible order of accuracy in this family.

We note that in contrast the corresponding explicit difference scheme would require an eleven-point stencil!

Second Derivative

Here we employ a general dual symmetric stencil of the 5/7 type as shown in figure 6.6.

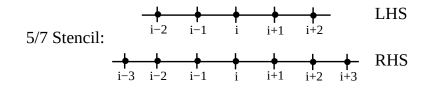


Figure 6.6: A 5/7 dual stencil for implicit discretization of the second derivative.

The general form of the discretization is

$$\beta(u_{xx})_{i-2} + \alpha(u_{xx})_{i-1} + (u_{xx})_i + \alpha(u_{xx})_{i+1} + \beta(u_{xx})_{i+2}$$

$$= a \frac{u_{i+1} - 2u_i + u_{i-1}}{\Delta x^2} + b \frac{u_{i+2} - 2u_i + u_{i-2}}{4\Delta x^2} + c \frac{u_{i+3} - 2u_i + u_{i-3}}{9\Delta x^2}.$$

We use Taylor expansions for the terms on both sides, and upon substitution in the above equation we obtain the constraints:

$$a + b + c = 1 + 2\alpha + 2\beta, \quad \mathcal{O}(\Delta x^{2})$$

$$a + 2^{2}b + 3^{2}c = \frac{4!}{2!}(\alpha + 2^{2}\beta), \quad \mathcal{O}(\Delta x^{4})$$

$$a + 2^{4}b + 3^{4}c = \frac{6!}{4!}(\alpha + 2^{4}\beta), \quad \mathcal{O}(\Delta x^{6})$$

$$a + 2^{6}b + 3^{6}c = \frac{8!}{6!}(\alpha + 2^{6}\beta), \quad \mathcal{O}(\Delta x^{8})$$

$$a + 2^{8}b + 3^{8}c = \frac{10!}{8!}(\alpha + 2^{8}\beta), \quad \mathcal{O}(\Delta x^{10})$$

$$\vdots$$

or in general form, for truncation error up to (Δx^p) :

$$a + 2^{p-2}b + 3^{p-2}c = \frac{p!}{(p-2)!}(\alpha + 2^{p-2}\beta), \quad p = 4, 6, \dots$$

As before, we can now construct either a 3/5 and a 5/7 stencil, the former corresponding to one-parameter family of schemes while the latter to a two-parameter family.

For the 3/5 stencil we have that $c = \beta = 0$ and for fourth-order accuracy we employ the first two constraints to obtain

$$a = \frac{4}{3}(1 - \alpha);$$
 $b = \frac{1}{3}(-1 + 10\alpha),$

with the truncation error

$$T_{3/5}'' = -\frac{4}{6!}(11\alpha - 2)\Delta x^4 \frac{\partial^6 u}{\partial x^6}.$$

We thus have an α -family of schemes with typical members:

• $\alpha = 1/10$; this leads to b = 0 which corresponds to the classical 3/3 Padé scheme.

- $\alpha = 2/11$; this leads to truncation error of order $\mathcal{O}(\Delta x^6)$.
- $\alpha \to 0$, this recovers the explicit central difference scheme.

The longer 5/7 stencil leads to a two-parameter family for schemes of sixth-order accuracy $\mathcal{O}(\Delta x^6)$, with free parameters α and β . Specifically, the truncation error has the form

$$T_{5/7}'' = -\frac{8}{8!}(9 - 38\alpha + 214\beta)\Delta x^6 \frac{\partial^8 u}{\partial x^8},$$

and the coefficients of the right-hand-side are

$$a = \frac{6 - 9\alpha - 12\beta}{4}$$
; $b = \frac{-3 + 24\alpha - 6\beta}{5}$; $c = \frac{2 - 11\alpha + 124\beta}{20}$.

If we require accuracy $\mathcal{O}(\Delta x^8)$, then we will employ four constraints while the unknowns are five, and thus a *one-parameter family* of schemes is obtained, the α -family. The corresponding coefficients are:

$$\beta = \frac{38\alpha - 9}{214}; \ a = \frac{696 - 1191\alpha}{428}; \ b = \frac{2454\alpha - 294}{530},$$

with truncation error

$$T_{5/7}'' = \frac{899\alpha - 334}{2696400} \Delta x^8 \frac{\partial^{10} u}{\partial x^{10}}.$$

Therefore, we can obtain very high-order accuracy, i.e., $\mathcal{O}(\Delta x^{10})$, for $\alpha = 334/899!$

Third Derivative

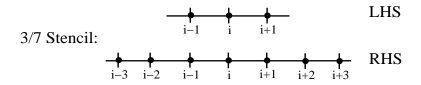


Figure 6.7: A 3/7 stencil for implicit discretization of the third derivative.

In order to approximate the third derivative with fourth-order accuracy on a 3/7 stencil, see figure 6.7, we need second-order approximations at both the integer and half grid points. On the integer grid points we have

$$(u_{xxx})_i = \frac{1}{2\Delta x^3}(u_{i+2} - 2u_{i+1} + 2u_{i-1} - u_{i-2}) - \frac{1}{4}\Delta x^2(u^{(iv)}),$$

while on the half grid points we have

$$(u_{xxx})_i = \frac{1}{\Delta x^3} (u_{i+3/2} - 3u_{i+1/2} + 3u_{i-1/2} - u_{i-3/2}) - \frac{\Delta x^2}{8} (u^{(v)}).$$

We then employ the equation

$$\alpha(u_{xxx})_{i-1} + (u_{xxx})_i + \alpha(u_{xxx})_{i+1} = b \frac{u_{i+3} - 3u_{i+1} + 3u_{i-1} - u_{i-3}}{8\Delta x^3} + a \frac{u_{i+2} - 2u_{i+1} + 2u_{i-1} - u_{i-2}}{2\Delta x^3}$$

to obtain the coefficients

$$a = 2; \quad b = 2\alpha - 1,$$

with truncation error

$$T_{3/7}^{"'} = \frac{42}{7!} (16\alpha - 7) \Delta x^4 \frac{\partial^7 u}{\partial x^7}.$$

We can now deduce some special members of this α -family:

- For $\alpha = 1/2$, we obtain b = 0, which shows that the 3/5 is the most compact $\mathcal{O}(\Delta x^4)$ scheme.
- For $\alpha = 7/16$ the leading term in the truncation error is zero, and thus we obtain $\mathcal{O}(\Delta x^6)$, which gives the highest possible accuracy for this family.

Fourth Derivative

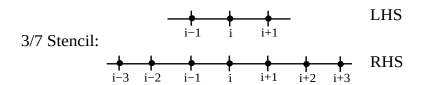


Figure 6.8: A 3/7 stencil for the implicit discretization of the fourth derivative.

From the many possible variations, here we present only the 3/7 stencil shown in figure 6.8. The corresponding matching of Taylor expansions leads to

$$\alpha u_{i-1}^{iv} + u_i^{iv} + \alpha u_{i+1}^{iv} = b \frac{u_{i+3} - 9u_{i+1} + 16u_i - 9u_{i-1} + u_{i-3}}{6\Delta x^4} + a \frac{u_{i+2} - 4u_{i+1} + 6u_i - 4u_{i-1} + u_{i-2}}{\Delta x^4}.$$

For a fourth-order accurate scheme we obtain the coefficients:

$$a = 2(1 - \alpha); \quad b = 4\alpha - 1,$$

with the best members

- For $\alpha = 1/4$, we obtain the most compact (3/5) stencil of $\mathcal{O}(\Delta x^4)$.
- For $\alpha = 7/26$, the leading term of the truncation error is zero and we obtain accuracy of $\mathcal{O}(\Delta x^6)$.

Boundary Conditions

Next we obtain one-sided formulas for implicit discretization of the first- and second derivative on a five-point stencil as shown in figure 6.9.

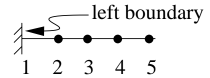


Figure 6.9: A five-point stencil for implicit discretization of the first and second derivatives.

For the **first derivative**, the equation generating the coefficients is

$$(u_x)_1 + \alpha(u_x)_2 = \frac{1}{\Delta x}(au_1 + bu_2 + cu_3 + du_4),$$

and depending on the required accuracy we can construct a specific formula or a family of schemes, as follows:

- $\mathcal{O}(\Delta x^2)$: $a = -\frac{3+\alpha+2d}{2}$; b = 2+3d; $c = \frac{1-\alpha+6d}{2}$.
- $\mathcal{O}(\Delta x^3)$: $a = -\frac{11+2\alpha}{6}$; $b = \frac{6-\alpha}{2}$; $c = \frac{2\alpha-3}{2}$; $d = \frac{2-\alpha}{6}$.
- $\mathcal{O}(\Delta x^4)$: $\alpha = 3$, a = -17/6; b = 3/2; d = -1/6.

For the **second derivative** we obtain similarly as before:

$$(u_{xx})_1 + 11(u_{xx})_2 = \frac{1}{\Delta x^2}(13u_1 - 27u_2 + 15u_3 - u_4) + \frac{\Delta x^3}{12} \left(\frac{\partial^5 u}{\partial x^5}\right)_1$$

which is of third-order accuracy. It is instructive, to compare this with the corresponding explicit one-sided formula:

$$(u_{xx})_1 = \frac{1}{\Delta x^2} \left(\frac{35}{12} u_1 - \frac{26}{3} u_2 + \frac{19}{2} u_3 - \frac{14}{3} u_4 + \frac{11}{12} u_5 \right) + \frac{5}{6} \Delta x^3 \left(\frac{\partial^5 u}{\partial x^5} \right)_1.$$

We see that the truncation error of the explicit difference is *ten times* larger than the corresponding third-order compact scheme!

Remark: We note here that in applications it is often possible to mix the order of discretization accuracy in the interior and at the boundary with the latter being of lower order. This may be required for stability of the approximation, e.g. in hyperbolic systems, or for computational efficiency, e.g. in elliptic problems. Theoretical justification for this can be obtained by a theorem, e.g. see [65].

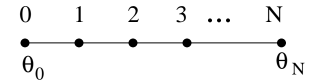


Figure 6.10: Grid for the one-dimensional boundary value problem.

6.1.3 One-Dimensional Boundary Value Problem

Let us revisit the one-dimensional boundary value problem (BVP) that we first encountered in section 5.1.6:

$$\begin{cases} \theta'' = q(x) \\ \theta_0 = \theta_N = 0 \end{cases}$$

We will apply different discretizations to this simple problem on the grid of figure 6.10 in order to illustrate the computational complexity involved in each approach.

We start with explicit second-order, $\mathcal{O}(\Delta x^2)$, differencing:

$$\frac{1}{\Delta x^2}(\theta_{i+1} - 2\theta_i + \theta_{i-1}) = q_i, \quad i = 1, \dots, N - 1.$$

We can cast these equations into matrix form as follows - recall that each equation is a row in the matrix:

$$\begin{bmatrix}
1 \\
1/\Delta x^2 & -2/\Delta x^2 & 1/\Delta x^2 & 0 \\
\vdots & \vdots & \vdots \\
0 & 1/\Delta x^2 & -2/\Delta x^2 & 1/\Delta x^2 \\
1
\end{bmatrix}
\begin{bmatrix}
\theta_0 \\
\theta_1 \\
\vdots \\
\theta_{N-1} \\
\theta_N
\end{bmatrix} = \begin{bmatrix}
q_0 \\
q_1 \\
\vdots \\
q_{N-1} \\
q_N
\end{bmatrix}$$

We can also write this in compact form

$$\mathbf{A}\,\theta = \mathbf{q}$$

where \mathbf{A} is a sparse tridiagonal matrix.

We now employ $matrix\ condensation$ to impose Dirichlet boundary conditions for this system. Assuming that the columns of matrix \mathbf{A} are denoted by \mathbf{a}_i then

$$\sum_{i=1}^{N-1} \theta_i \mathbf{a}_i = \mathbf{q} - \theta_0 \mathbf{a}_0 - \theta_N \mathbf{a}_N.$$

This is an efficient approach as it effectively reduces the rank of the matrix we need to invert.

Remark: Note here that it will be extremely inefficient to obtain the solution by multiplying the right-hand-side by the inverse of matrix \mathbf{A} , as the inverse is a full matrix! This can be

heuristically justified by a physical analog. Consider, for example, that we solve a steadystate heat equation on a rod, which is governed by the above one-dimensional BVP. Let us now denote the columns of \mathbf{A}^{-1} by \mathbf{b}_{i}^{-1} ; then the solution can be written as

$$\theta = \mathbf{A}^{-1}\mathbf{q} \Rightarrow \theta = \sum_{i=0}^{N} q_i \mathbf{b}_i^{-1}.$$

Now consider that we apply a discrete unit heat source at node (i), i.e.,

$$q_i = 1; \quad q_j = 0, \quad j \neq i,$$

then:

$$\theta = \mathbf{b}_i^{-1}$$
.

In other words the elements of the column-vector \mathbf{b}^{-1} are the temperature values at the nodes due to a unit heat source at node i. Thus, they should be *non-zero* because of the ellipticity property and the fact that we imposed zero values at the end-points.

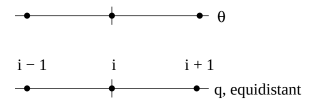


Figure 6.11: Dual 3/3 stencil for discretization of the second-derivative in the BVP.

Next we employ a *compact scheme* on a 3/3 stencil (figure 6.11) for the discretization of the one-dimensional BVP. We could write down directly the discretization but here we repeat the derivation of the formulas using Taylor expansions and the method of undetermined coefficients. To this end, we have to expand

$$\theta_{i\pm 1} = \theta_i \pm \Delta x^2 \theta_i' + \frac{\Delta x^2}{2} \theta_i'' \pm \frac{\Delta x^4}{24} \theta_i^{iv} + \dots$$

and use the model

$$\alpha\theta_{i-1} + \beta\theta_i + \alpha\theta_{i+1} = aq_{i-1} + bq_i + aq_{i+2} + T_i.$$

Notice here that instead of expanding the second derivative we expand the right-hand-side. However, we have that

$$\theta'' = q \Rightarrow q_{i\pm 1} = q_i \pm \Delta x q_i' + \frac{\Delta x^2}{2} \theta^{(iv)} + \mathcal{O}(\Delta x^4).$$

Proceeding as before, we obtain the coefficients

$$\alpha = \frac{1}{\Delta x^2}; \quad \beta = -2/\Delta x^2$$

$$a = \frac{1}{12}; \quad b = \frac{5}{6}.$$

Thus, upon substitution in the BVP we obtain

$$\frac{1}{\Delta x^2}(\theta_{i-1} - 2\theta_i + \theta_{i+1}) = \frac{1}{12}(q_{i-1} + 10q_i + q_{i+1}) + \mathcal{O}(\Delta x^4).$$

This rather simple example helps us to understand what is the difference between the *explicit* and the *implicit* discretization. Specifically, here following the implicit discretization we have effectively modified the right-hand-side (RHS) through a *mass matrix*, that is we have distributed the forcing around the node of interest, just like in finite element methods (e.g., see [59]). Therefore, for the RHS we have:

$$RHS: \frac{1}{12} \begin{bmatrix} & & & & \\ & \ddots & \ddots & O \\ & 1 & 10 & 1 \\ O & & \ddots & \ddots \\ & & Mass\ Matrix \end{bmatrix} \begin{bmatrix} \vdots \\ q_{i-1} \\ q_i \\ q_{i+2} \\ \vdots \end{bmatrix}$$

Note that what we have obtained by expanding the RHS in this particular BVP is identical to the most compact scheme of $\mathcal{O}(\Delta x^4)$ for $\alpha = 1/10$, i.e.,

$$\frac{1}{10} (\underbrace{\theta_{xx}}_{q_{i-1}})_{i-1} + (\underbrace{\theta_{xx}}_{q_i})_i + \frac{1}{10} (\underbrace{\theta_{xx}}_{q_{i+1}})_{i+1} = \frac{12}{10} \frac{\theta_{i+1} - 2\theta_i + \theta_{i+1}}{\Delta x^2} + \mathcal{O}(\Delta x^4).$$

This formula is exactly what we have derived previously in equation (6.4).

6.1.4 Thomas Algorithm for Tridiagonal Systems

Three-point stencils lead to second-order accuracy and fourth-order accuracy for explicit and implicit discretizations of second-order boundary value problems, respectively. As we have seen from the example above, solution of such BVPs reduces to solving the linear system

$$\mathbf{A}\mathbf{x} = \mathbf{q}$$

where the matrix **A** is tridiagonal if the boundary conditions are Dirichlet. In the following, we demonstrate how to solve this system using the *Thomas algorithm*, which is a special case of Gaussian elimination, see also section 9.1. This method consists of three main steps:

- The LU decomposition of matrix **A**, that is its factorization into a lower triangular matrix **L** and an upper triangular matrix **U**. Note that this factorization maintains the bandwidth, and therefore both matrices **L** and **U** are bidiagonal.
- The forward substitution where the matrix L is involved, and
- The final backward substitution, where the matrix **U** is involved.

More specifically, we have:

$$\begin{bmatrix} a_1 & c_1 & & & & \\ b_2 & a_2 & c_2 & 0 & & \\ & b_3 & a_3 & c_3 & & \\ & \ddots & \ddots & \ddots & \\ & 0 & & b_N & a_N \end{bmatrix} = \begin{bmatrix} 1 & & & & \\ \ell_2 & 1 & & 0 & & \\ & \ell_3 & 1 & & \\ & & \ddots & \ddots & \\ & 0 & & \ell_N & 1 \end{bmatrix} \begin{bmatrix} d_1 & u_1 & & & \\ & d_2 & u_2 & 0 & & \\ & & d_3 & u_3 & & \\ & & & 0 & \ddots & \ddots & \\ & & & & d_N \end{bmatrix}$$

Step 1: (LU Decomposition) A = LU.

We determine the elements of matrices ${\bf L}$ and ${\bf U}$ in three stages, separating the end-points from the interior points as follows:

$$d_1 = a_1, \quad u_1 = c_1$$

$$i^{th} \begin{cases} \ell_i d_{i-1} = b_i \Rightarrow \ell_i = b_i / \ell_i d_{i-1}, & (N \ multiplications) \\ \ell_i u_{i-1} + d_i = a_i \Rightarrow d_i = a_i - \ell_i u_{i-1} & (N \ multiplications, N \ additions) \end{cases}$$

$$N^{th} \begin{cases} \ell_N d_{N-1} = b_N \Rightarrow \ell_N = b_N / d_{N-1} \\ \ell_N u_{N-1} + d_N = a_N \Rightarrow d_N = a_N - \ell_N u_{N-1} \end{cases}$$

Therefore, from the above we see that the total *computational complexity* for an LU decomposition of a tridiagonal matrix corresponds to 2N multiplications and N additions.

Step 2: (Forward Substitution) Ly = q The intermediate vector y is determined from

$$\begin{bmatrix} 1 & & & & \\ \ell_2 & 1 & & & \\ & \ell_3 & 1 & & \\ & & \ddots & \ddots & \\ & & & \ell_N & 1 \end{bmatrix} \begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ \vdots \\ y_N \end{bmatrix} = \begin{bmatrix} q_1 \\ q_2 \\ q_3 \\ \vdots \\ q_N \end{bmatrix} \Rightarrow \begin{cases} y_1 = q_1 \\ \ell_i y_{i-1} + y_i = q_i \Rightarrow y_i = q_i - \ell_i y_{i-1} \end{cases}$$

Here the operation count is N multiplications and N additions.

Step 3: (Backward Substitution) Ux = y

In the final step we have

$$\begin{bmatrix} d_1 & u_1 & & & & & \\ & d_2 & u_2 & & & & \\ & & d_3 & u_3 & & & \\ & 0 & & \ddots & \ddots & \\ & & & & d_N \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \\ \vdots \\ x_N \end{bmatrix} = \begin{bmatrix} y_1 \\ y_2 \\ y_3 \\ \vdots \\ y_N \end{bmatrix}$$

and the final solution is obtained from:

$$\begin{cases} x_N = y_N/d_N \\ d_i x_i + u_i x_{i+1} = y_i \Rightarrow x_i = (y_i - u_i x_{i+1})/d_i, & i = N - 1, \dots, 1 \end{cases}$$

The corresponding operation count is 2N multiplications and N additions.

We can now summarize the operation count:

- 1) LU: 2N multiplications, N additions
- 2) Forward: N multiplications, N additions
- 3) Backward: 2N multiplications, N additions

Total: 5N multiplications, 3N additions

Remark: It can be shown that the above algorithm will always converge if the tridiagonal system is diagonally dominant, i.e.,

$$|a_k| \ge |b_k| + |c_k|, k = 2, ..., N - 1$$

 $|a_1| > |c_1| \text{ and } |a_N| > |b_N|.$

Also, if a, b, c are matrices instead of scalars we will have a *block-tridiagonal* system and the same algorithm can be applied.

WARNING

Programmer Beware!

• Think carefully about indexing. Simple mistakes can cause major headaches!

Software



Putting it into Practice

Below we present a serial C++ implementation of the Thomas algorithm presented above.

```
y = new double[N];
/* LU Decomposition */
d[0] = a[0];
u[0] = c[0];
for(i=0;i<N-2;i++){
  l[i] = b[i]/d[i];
  d[i+1] = a[i+1] - l[i]*u[i];
  u[i+1] = c[i+1];
}
1[N-2] = b[N-2]/d[N-2];
d[N-1] = a[N-1] - 1[N-2]*u[N-2];
/* Forward Substitution [L][y] = [q] */
y[0] = q[0];
for(i=1;i<N;i++)
  y[i] = q[i] - 1[i-1]*y[i-1];
/* Backward Substitution [U][x] = [y] */
x[N-1] = y[N-1]/d[N-1];
for(i=N-2;i>=0;i--)
  x[i] = (y[i] - u[i]*x[i+1])/d[i];
delete[] 1;
delete[] u;
delete[] d;
delete[] y;
return;
```

}

Common Programming Trick: Notice that every time that we call the above routine we must allocate and deallocate memory. Suppose that we are calling this routine over and over, and using the same size allocation each time. We are wasting a lot of time just allocating and deallocating! What can be done? One common trick is to use static. When a variable is declared static, it is allocated each once (in the static part of a program's memory), and remains throughout the duration of the program. Hence, if you declare a pointer variable as static within the routine, and allocate an array of memory the first time that the routine is called, then you can dispense with allocating/deallocating each time. This is demonstrated in the following modified code.

```
if(1 == NULL){
  l = new double[N];
  u = new double[N];
  d = new double[N];
  y = new double[N];
}
/* LU Decomposition */
d[0] = a[0];
u[0] = c[0];
for(i=0;i<N-2;i++){
  l[i] = b[i]/d[i];
  d[i+1] = a[i+1] - l[i]*u[i];
  u[i+1] = c[i+1];
}
1[N-2] = b[N-2]/d[N-2];
d[N-1] = a[N-1] - 1[N-2]*u[N-2];
/* Forward Substitution [L][y] = [q] */
y[0] = q[0];
for(i=1;i<N;i++)</pre>
  y[i] = q[i] - 1[i-1]*y[i-1];
/* Backward Substitution [U][x] = [y] */
x[N-1] = y[N-1]/d[N-1];
for(i=N-2;i>=0;i--)
  x[i] = (y[i] - u[i]*x[i+1])/d[i];
return;
```

}

Remark: In the example we show above, the pointer variables are declared static, and are initialized to NULL the first time that the routine is run. Since the pointer is NULL, the memory is allocated the first time the routine is run; however, for all subsequent calls, the value of the pointer l is not NULL (it contains some address value), and hence memory is not allocated. Of course, in this implementation we have assumed that the value of N is always less than or equal to the first value of N passed to this routine. More complex schemes can be devised to make allocate/deallocate only when the size changes. This methodology is a common trick - valid with respect to the language, but despised by many as unclean programming!

Key Concept

• You need not recompute things that do not change!

Instead of using static allocations, one preferred way of increasing code re-use is to move the memory allocation outside of the Thomas Algorithm routines and to break the algorithm into two functions:

- 1. ThomasAlgorithmLU accomplishes the LU decomposition of the matrix **A**. This routine needs to be called only once per matrix **A**.
- 2. ThomasAlgorithmSolve accomplishes the forward and back substitution. This routine needs to be called every time the right-hand-side value **b** changes.

The memory allocation is moved outside of these functions; the calling function is responsible for memory allocation. We now present both the functions described above.

```
void ThomasAlgorithmLU(int N, double *b, double *a, double *c,
                       double *1, double *u, double *d){
  int i;
  /* LU Decomposition */
  d[0] = a[0];
  u[0] = c[0];
  for(i=0;i<N-2;i++){
    l[i] = b[i]/d[i];
    d[i+1] = a[i+1] - l[i]*u[i];
    u[i+1] = c[i+1];
  }
  1[N-2] = b[N-2]/d[N-2];
  d[N-1] = a[N-1] - 1[N-2]*u[N-2];
  return;
}
void ThomasAlgorithmSolve(int N, double *1, double *u, double *d,
                           double *x, double *q){
  int i;
  double *y = new double[N];
  /* Forward Substitution [L][y] = [q] */
  y[0] = q[0];
  for(i=1;i<N;i++)
```

```
y[i] = q[i] - 1[i-1]*y[i-1];

/* Backward Substitution [U][x] = [y] */
x[N-1] = y[N-1]/d[N-1];
for(i=N-2;i>=0;i--)
    x[i] = (y[i] - u[i]*x[i+1])/d[i];

delete[] y;
return;
}
```

Remark: Notice that the function which accomplishes the forward solve and the back solve does not require the matrix arrays a, b and c; it requires only the l, u and d arrays which contain the LU decomposition of \mathbf{A} . Once the LU decomposition has been accomplished, and if the matrix \mathbf{A} is not needed for any other purpose, the arrays a, b, and c can be deallocated.

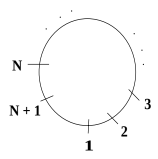


Figure 6.12: Domain for solving the steady heat equation in a ring.

Thomas Algorithm for Periodic Tridiagonal Systems

The boundary value problem we considered in the example above employed Dirichlet boundary conditions, but often *periodic boundary conditions* are required. This could be a case for example, where an infinite domain is simulated or the physics of the problem dictates it, as in solving an elliptic problem on a ring (see figure 6.12). In this case, despite the sparsity of the matrix resulting from the discretization and its almost tridiagonal form everywhere, the bandwidth is actually equal to the order of the matrix in the form shown below:

$$\begin{bmatrix} a_1 & c_1 & & & b_1 \\ b_2 & a_2 & c_2 & 0 & & \\ & \ddots & \ddots & \ddots & \\ & & \ddots & \ddots & c_N \\ c_{N+1} & 0 & & b_{N+1} & a_{N+1} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ \vdots \\ x_N \\ x_{N+1} \end{bmatrix} = \begin{bmatrix} q \end{bmatrix}$$

where we assume that b_1 and c_{N+1} are coefficients corresponding to the periodic boundary conditions, (e.g., equal to 1 in the example above).

We can solve this system by first "condensing" the matrix, that is eliminating the last row and the last column, to arrive at:

$$\begin{bmatrix}
a_1 & c_1 \\
b_2 & a_2 & c_2 & 0 \\
& \ddots & \ddots & \\
0 & b_N & a_N
\end{bmatrix}
\begin{bmatrix}
x_1 \\
x_2 \\
\vdots \\
x_N
\end{bmatrix} = \mathbf{q} - \begin{bmatrix}
b_1 \\
0 \\
\vdots \\
0 \\
c_N
\end{bmatrix} x_{N+1}.$$

Now we use the linear property and propose a superposition of the form

$$\mathbf{x} = \mathbf{x}^{(1)} + \mathbf{x}^{(2)} \cdot x_{N+1} \,,$$

where $x^{(1)}$ and $x^{(2)}$ are solutions of the tridiagonal "condensed" system with N unknowns, i.e.,

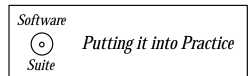
$$\mathbf{A}^{c}\mathbf{x}^{(1)} = \mathbf{q}$$
 $\mathbf{A}^{c}\mathbf{x}^{(2)} = \begin{bmatrix} -b_1 \\ 0 \\ \vdots \\ 0 \\ -c_N \end{bmatrix}$.

We finally compute x_{N+1} from the last equation in the original system by back substitution, i.e.,

$$c_{N+1}(x_1^{(1)} + x_{N+1}x_1^{(2)}) + b_{N+1}(x_N^{(1)} + x_{N+1}x_N^{(2)}) + a_{N+1}x_{N+1} = q_{N+1}$$

and we solve for x_{N+1} :

$$x_{N+1} = \frac{q_{N+1} - c_{N+1} x_1^{(1)} - b_{N+1} x_N^{(1)}}{a_{N+1} + c_{N+1} x_1^{(2)} + b_{N+1} x_N^{(2)}}.$$



Below we present a serial C++ implementation of the Thomas Algorithm for *periodic* systems. As was discussed above, the Thomas Algorithm for periodic systems requires us to accomplish LU solves on *condensed systems*. Note that we accomplish this by re-using the Thomas algorithm functions that we previously presented on the condensed system.

```
void ThomasAlgorithm_per(int N, double *b, double *a, double *c,
                         double *x, double *q){
  int i;
  double *x1,*x2,*q2;
  x1 = new double[N-1];
  x2 = new double[N-1];
  q2 = new double[N-1];
  /* Prepare secondary q */
  for(i=0;i<N-1;i++)
   q2[i] = 0.0;
  q2[0] = -b[N-1];
  q2[N-2] = -c[N-2];
  ThomasAlgorithm(N-1,b,a,c,x1,q);
  ThomasAlgorithm(N-1,b,a,c,x2,q2);
  x[N-1] = (q[N-1] - c[N-1]*x1[0] - b[N-2]*x1[N-2])/
    (a[N-1] + c[N-1]*x2[0] + b[N-2]*x2[N-2]);
  for(i=0;i<N-1;i++)
   x[i] = x1[i] + x2[i]*x[N-1];
  delete[] x1;
  delete[] x2;
  delete[] q2;
}
```

Key Concept

• Code re-use is important. If you have already invested the time to make sure that a routine is well-written and correctly implemented, then you can use the routine as a component in new routines.

6.1.5 Parallel Algorithm for Tridiagonal Systems

In seeking a parallelization strategy for solving triagonal systems, we will once again examine the structure of the LU decomposition as we did in formulating the Thomas algorithm. By exploiting the recursive nature of the LU decomposition, we will devise a *full-recursive-doubling*

procedure for solving for the unknown LU coefficients. For a more detailed description of the algorithm which follows we refer the reader to [24].

As before, we seek an LU decomposition of the tridiagonal matrix A as follows:

$$\begin{bmatrix} a_1 & c_1 & & & & \\ b_2 & a_2 & c_2 & 0 & & \\ b_3 & a_3 & c_3 & & \\ & \ddots & \ddots & \ddots & \\ 0 & & b_N & a_N \end{bmatrix} = \begin{bmatrix} 1 & & & & \\ \ell_2 & 1 & & 0 & & \\ & \ell_3 & 1 & & \\ & & \ddots & \ddots & \\ 0 & & \ell_N & 1 \end{bmatrix} \begin{bmatrix} d_1 & u_1 & & & \\ d_2 & u_2 & 0 & & \\ & d_3 & u_3 & & \\ & & 0 & \ddots & \ddots & \\ & & & d_N \end{bmatrix}.$$

Upon examination of the expression above, we see that we can formulate recurrence relations for the unknown coefficients d_i , u_i , and l_i as follows:

$$a_1 = d_1 \tag{6.5}$$

$$c_i = u_i \tag{6.6}$$

$$c_{j} = u_{j}$$
 (6.6)
 $a_{k} = d_{k} + l_{k}u_{k-1}$ (6.7)

$$b_k = l_k d_{k-1} (6.8)$$

where j = 1, ..., N and k = 2, ..., N. Given equation (6.6), we can immediately solve for all the unknown coefficients u_i . To solve for d_i and l_i , we rely on the recursive nature of these equations. Substituting equations (6.6) and (6.8) into equation (6.7) and rearranging terms yields the following rational recursion relationship for the unknown coefficient d_i :

$$d_{j} = a_{j} - l_{j}u_{j-1}$$

$$= a_{j} - \frac{b_{j}}{d_{j-1}}u_{j-1}$$

$$= \frac{a_{j}d_{j-1} - b_{j}c_{j-1}}{d_{j-1} + 0}.$$

We can then inductively solve for the all coefficients d_i , and use this information along with equation 6.8 to solve for l_i .

To parallelize this procedure, we make use of a full-recursive-doubling procedure on the sequence of 2×2 matrices given by:

$$\mathbf{R}_0 = \left[\begin{array}{cc} a_0 & 0 \\ 1 & 0 \end{array} \right]$$

and

$$\mathbf{R}_j = \left[\begin{array}{cc} a_j & -b_j c_{j-1} \\ 1 & 0 \end{array} \right]$$

for j = 1, ..., N. Using the Möbius transformations

$$\mathbf{T}_i = \mathbf{R}_i \mathbf{R}_{i-1} \dots \mathbf{R}_0$$

we have that

$$d_{j} = \frac{\begin{pmatrix} 1 \\ 0 \end{pmatrix}^{t} \mathbf{T}_{j} \begin{pmatrix} 1 \\ 1 \end{pmatrix}}{\begin{pmatrix} 0 \\ 1 \end{pmatrix}^{t} \mathbf{T}_{j} \begin{pmatrix} 1 \\ 1 \end{pmatrix}}.$$

To explain how this information can be used for parallelization, we will examine a specific example. Suppose that we are given a tridiagonal matrix A of size 40, and that we want to solve the problem using 8 processes. Assume that all processes have a copy of the original matrix A. We first partition the matrix such that each process is responsible for five rows: process P_0 is responsible for rows 0-4, P_1 is responsible for rows 5-9, etc. We then accomplish the following steps:

- 1. On each process P_i form the matrices \mathbf{R}_k , where k corresponds to the row indices for which the process is responsible, and ranges between k_min and k_max .
- 2. On each process P_i form the matrix $\mathbf{S}_i = \mathbf{R}_{k_max} \mathbf{R}_{k_max-1} \dots \mathbf{R}_{k_min}$.
- 3. Using the full-recursive-doubling communication pattern as given in table 6.1, distribute and combine the S_i matrices as given in table 6.2.
- 4. On each process P_j calculate the local unknown coefficients d_k $(k_min \le k \le k_max)$ using local R_k and matrices obtained from the full-recursive-doubling.
- 5. For processes P_0 through P_6 , send the local d_{k_max} to the process one process id up (i.e., P_0 sends to P_1 ; P_1 sends to P_2 ; etc.).
- 6. On each process P_i calculate the local unknown coefficients l_k $(k_min \le k \le k_max)$ using the local d_k values and the value obtained in the previous step.
- 7. Distribute the d_j and l_j values across all processes so that each process has all the d_j and l_i coefficients.
- 8. On each process P_i perform a local forward and backward substitution to obtain the solution.

Software

Putting it into Practice

$$\begin{array}{ccccc} \underline{\operatorname{Stage}\ 1} & \underline{\operatorname{Stage}\ 2} & \underline{\operatorname{Stage}\ 3} \\ \overline{P_0 \to P_1} & \overline{P_0 \to P_2} & \overline{P_0 \to P_4} \\ P_1 \to P_2 & P_1 \to P_3 & P_1 \to P_5 \\ P_2 \to P_3 & P_2 \to P_4 & P_2 \to P_6 \\ P_3 \to P_4 & P_3 \to P_5 & P_3 \to P_7 \\ P_4 \to P_5 & P_4 \to P_6 \\ P_5 \to P_6 & P_5 \to P_7 \\ P_6 \to P_7 \end{array}$$

Table 6.1: Full-recursive-doubling communication pattern. The number of stages is equal to the $\log_2 M$ where M is the number of processes. In this case, M=8 and hence there are three stages of communication.

$\underline{\text{Process}}$	Stage 0	Stage 1	Stage 2	Stage 3
P_0	$\overline{S_0}$	' <u></u>		<u> </u>
P_1	S_1	S_1S_0		
P_2	S_2	S_2S_1	$S_2S_1S_0$	
P_3	S_3	S_3S_2	$S_3S_2S_1S_0$	
P_4	S_4	S_4S_3	$S_4S_3S_2S_1$	$S_4 S_3 S_2 S_1 S_0$
P_5	S_5	S_5S_4	$S_5S_4S_3S_2$	$S_5S_4S_3S_2S_1S_0$
P_6	S_6	S_6S_5	$S_6S_5S_4S_3$	$S_6S_5S_4S_3S_2S_1S_0$
P_7	S_7	S_7S_6	$S_7S_6S_6S_4$	$S_7 S_6 S_5 S_4 S_3 S_2 S_1 S_0$

Table 6.2: Distribution and combination pattern of the S_j matrices for each stage. The interpretation of the table is as follows: Given the communication pattern as given in table 6.1, in stage one P_0 sends S_0 to P_1 , which P_1 combines with its local S_1 to form the product S_1S_0 . Similarly in stage one, P_1 sends S_1 to P_2 , etc. In stage two, P_0 sends S_0 to P_2 , which P_2 combines with its local product S_2S_1 to form $S_2S_1S_0$. Similarly P_1 sends S_1S_0 to P_3 which is then combined on P_3 to form $S_3S_2S_1S_0$. In stage three, the final communications occur such that each process j stores locally the product $S_jS_{j-1}...S_0$.

We now present a parallel Thomas algorithm function which uses the full-recursiveprocedure discussed above. This function assumes that the MPI initialization has already been accomplished by the calling function, and it requires that the number of processes used is a power of two. It takes as input its process id number, the total number of processes being used, the size of the matrix, the matrix \mathbf{A} stored in the arrays a, b, and c as before, and the right-hand-side vector \mathbf{q} stored in the array q. The output of this function on all processes is the solution vector contained within the array x. We first present the function definition and then present some remarks on the code.

```
void ThomasAlgorithm_P(int mynode, int numnodes, int N, double *b,
                       double *a, double *c, double *x, double *q){
  int i,j,k,i_global;
  int rows_local,local_offset;
  double S[2][2],T[2][2],s1tmp,s2tmp;
  double *1,*d,*v;
  MPI_Status status;
  l = new double[N];
  d = new double[N];
  y = new double[N];
  for(i=0;i<N;i++)
    l[i] = d[i] = y[i] = 0.0;
  S[0][0] = S[1][1] = 1.0;
  S[1][0] = S[0][1] = 0.0;
  rows_local = (int) floor(N/numnodes);
  local_offset = mynode*rows_local;
  // Form local products of R_k matrices
  if(mynode==0){
    s1tmp = a[local_offset]*S[0][0];
    S[1][0] = S[0][0];
    S[1][1] = S[0][1];
    S[0][1] = a[local_offset] *S[0][1];
    S[0][0] = s1tmp;
    for(i=1;i<rows_local;i++){</pre>
      s1tmp = a[i+local_offset]*S[0][0] -
              b[i+local_offset-1]*c[i+local_offset-1]*S[1][0];
      s2tmp = a[i+local_offset]*S[0][1] -
              b[i+local_offset-1]*c[i+local_offset-1]*S[1][1];
      S[1][0] = S[0][0];
      S[1][1] = S[0][1];
      S[0][0] = s1tmp;
```

```
S[0][1] = s2tmp;
  }
}
else{
  for(i=0;i<rows_local;i++){</pre>
    s1tmp = a[i+local_offset]*S[0][0] -
            b[i+local_offset-1]*c[i+local_offset-1]*S[1][0];
    s2tmp = a[i+local_offset]*S[0][1] -
            b[i+local_offset-1]*c[i+local_offset-1]*S[1][1];
    S[1][0] = S[0][0];
    S[1][1] = S[0][1];
    S[0][0] = s1tmp;
    S[0][1] = s2tmp;
 }
}
// Full-recursive doubling algorithm for distribution
for(i=0; i<=log2(numnodes);i++){</pre>
  if(mynode+pow(2,i) < numnodes)</pre>
    MPI_Send(S,4,MPI_DOUBLE,int(mynode+pow(2,i)),0,
             MPI_COMM_WORLD);
  if (mynode-pow(2,i)>=0) {
    MPI_Recv(T,4,MPI_DOUBLE,int(mynode-pow(2,i)),0,
             MPI_COMM_WORLD,&status);
    s1tmp = S[0][0]*T[0][0] + S[0][1]*T[1][0];
    S[0][1] = S[0][0]*T[0][1] + S[0][1]*T[1][1];
    S[0][0] = s1tmp;
    s1tmp = S[1][0]*T[0][0] + S[1][1]*T[1][0];
    S[1][1] = S[1][0]*T[0][1] + S[1][1]*T[1][1];
    S[1][0] = s1tmp;
  }
}
//Calculate last d_k first so that it can be distributed,
//and then do the distribution.
d[local\_offset+rows\_local-1] = (S[0][0] + S[0][1])/
                                (S[1][0] + S[1][1]);
if(mynode == 0){
  MPI_Send(&d[local_offset+rows_local-1],1,MPI_DOUBLE,
           1,0,MPI_COMM_WORLD);
}
else{
  MPI_Recv(&d[local_offset-1],1,MPI_DOUBLE,mynode-1,0,
           MPI_COMM_WORLD,&status);
  if(mynode != numnodes-1)
```

```
MPI_Send(&d[local_offset+rows_local-1],1,MPI_DOUBLE,
            mynode+1,0,MPI_COMM_WORLD);
}
// Compute in parallel the local values of d_k and l_k
if(mynode == 0){
 1[0] = 0;
 d[0] = a[0];
 for(i=1;i<rows_local-1;i++){</pre>
   l[local_offset+i] = b[local_offset+i-1]/
                       d[local_offset+i-1];
   d[local_offset+i] = a[local_offset+i] -
                       l[local_offset+i]*c[local_offset+i-1];
 }
  1[local_offset+rows_local-1] = b[local_offset+rows_local-2]/
                                d[local_offset+rows_local-2];
}
else{
  for(i=0;i<rows_local-1;i++){</pre>
   l[local_offset+i] = b[local_offset+i-1]/
                       d[local_offset+i-1];
   d[local_offset+i] = a[local_offset+i] -
                       l[local_offset+i]*c[local_offset+i-1];
 }
 1[local_offset+rows_local-1] = b[local_offset+rows_local-2]/
                                d[local_offset+rows_local-2];
}
if(mynode>0)
 d[local_offset-1] = 0;
// Distribute d_k and l_k to all processes
double * tmp = new double[N];
for(i=0;i<N;i++)
  tmp[i] = d[i];
MPI_Allreduce(tmp,d,N,MPI_DOUBLE,MPI_SUM,MPI_COMM_WORLD);
for(i=0;i<N;i++)</pre>
  tmp[i] = 1[i];
MPI_Allreduce(tmp,1,N,MPI_DOUBLE,MPI_SUM,MPI_COMM_WORLD);
delete[] tmp;
```

```
if(mynode ==0){
    /* Forward Substitution [L][y] = [q] */
    y[0] = q[0];
    for(i=1;i<N;i++)
        y[i] = q[i] - l[i]*y[i-1];

    /* Backward Substitution [U][x] = [y] */
    x[N-1] = y[N-1]/d[N-1];
    for(i=N-2;i>=0;i--)
        x[i] = (y[i] - c[i]*x[i+1])/d[i];
}

delete[] l;
delete[] y;
delete[] d;
return;
}
```

Remark 1: Since we know that we are dealing with 2×2 matrices, we have chosen to allocate the 2×2 S array statically. It is important to note that when static allocation of arrays is used, the memory allocation is contiguous and in row-major order as shown in figure 6.13. We can use the contiguousness of the block of memory to our advantage when using MPI. Since S is stored as one contiguous block in memory, we can send the entire array in one MPI call instead of having to send the array row by row (as in the case where each row was dynamically allocated using the **new** command).

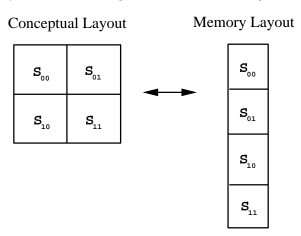


Figure 6.13: Memory layout of the matrix S. The double indexed array S is stored in a contiguous block of memory in row-major order.

Remark 2: Sometimes it becomes advantageous to use the reduction operator to mimic a gathering operation. We pictorially demonstrate how this can be accomplished in figure 6.14. In the code above, we use this trick to gather all the d_i and l_i values across all processors.

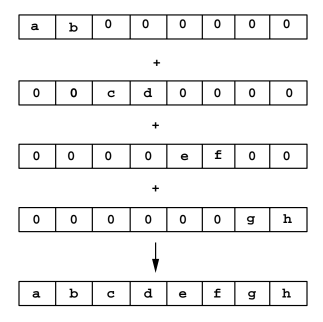


Figure 6.14: How to use the reduction operator to mimic the gathering process. In this example, we have four processes, each of which has two unique items to contribute. On each process, all other entries in the array are zeroed and then a sum is performed. The result is that the data is gathered into one array.

MPI Implementation Issues

In the sections above, we presented a serial and a parallel version of the Thomas algorithm. How can we time our parallel program to examine the speed-up due to adding more processes? MPI provides two functions which allow us to accomplish this task: MPI_Wtime and MPI_Wtick . We will now present for these two functions the function call syntax, argument list explanation, usage example and some remarks.

Function Call Syntax

```
double MPI_Wtime(void);
```

double MPI_Wtick(void);

Understanding the Argument Lists

• MPI_Wtime and MPI_Wtick take no arguments.

Example of Usage

```
int mynode, totalnodes;
double starttime, finaltime, precision;
```

```
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

precision = MPI_Wtick();
starttime = MPI_Wtime();

// Execution of commands here

finaltime = MPI_Wtime();

if(mynode == 0){
   cout << "The execution time was : " << finaltime-starttime;
   cout << " sec. with a precision of " << precision;
   cout << " sec." << endl;
}</pre>
```

Remarks

- These commands are very useful both for determining the parallel speed-up of your algorithm and for determining the components of your program which are using the most time.
- These commands provide to you the wallclock time (the physical time which has elapsed), not specifically the CPU time or communication time.

One question you may ask is how do I know that all the processes are exactly at the same point (assuming that I am doing the timing only on process 0)? MPI provides a function for synchronizing all processes called $MPI_Barrier$. When $MPI_Barrier$ is called, the function will not return until all processes have called $MPI_Barrier$. This functionality allows you to synchronize all the processes, knowing that all processes exit the $MPI_Barrier$ call at the same time. We will now present the function call syntax, argument list explanation, usage example and some remarks.

MPI_Barrier

Function Call Syntax

```
int MPI_Barrier(
MPI_Comm comm /* in */,
```

Understanding the Argument Lists

• comm - communicator

Example of Usage

```
int mynode, totalnodes;

MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

MPI_Barrier(MPI_COMM_WORLD);

// At this stage, all processes are synchronized
```

Remarks

• This command is a useful tool to help insure synchronization between processes. For example, you may want all processes to wait until one particular process has read in data from disk. Each process would call MPI_Barrier in the place in the program where the synchronization is required.

6.2 Implicit Time Discretizations

Unlike the explicit time discretizations of the previous chapter, here the solution at the current level cannot be expressed in terms of information from previous only time levels. Instead, the right-hand-side for the first-order initial value problem (IVP)

$$\frac{dU}{dt} = \mathcal{F}(U,t), \quad U(t=0) = U_0, \tag{6.9}$$

is also evaluated at the *current time level* resulting in an implicit expression for the solution.

The most popular schemes are derived from the θ -family, which are one-step schemes:

$$\frac{U^{n+1} - U^n}{\Delta t} = \theta \mathcal{F}(U^{n+1}, t^{n+1}) + (1 - \theta) \mathcal{F}(U^n, t^n).$$
 (6.10)

- For $\theta = 1$, we obtain the Euler-backward scheme, while
- For $\theta = 1/2$, we obtain the *Crank-Nicolson* scheme.

We note that θ can take any value in the range [0,1], and for $\theta = 0$ we obtain the Euler-forward method, an explicit scheme we encountered in section 5.2.

The θ -family produces a consistent discretization of the IVP we consider, in the form

$$\left(\frac{dU^n}{dt} - \mathcal{F}^n\right) = \Delta t \left(\theta \frac{d\mathcal{F}^n}{dt} - \frac{1}{2} \frac{d^2 U^n}{dt^2}\right) + \Delta t^2 \left(\frac{\theta}{2} \frac{d^2 \mathcal{F}^n}{dt^2} - \frac{1}{6} \frac{d^3 U^n}{dt^3}\right) + \dots$$

As $\Delta t \to 0$, the right-hand-side of the above discretization goes to zero and thus we recover the original initial value problem of equation (6.9). Specifically, for $\theta = 1$ or $\theta = 1/2$ the truncation error (right-hand-side) asymptotes to zero with first- and second-order convergence rate, correspondingly. This convergence rate, which expresses the first- and second-order accuracy of the Euler-backward and Crank-Nicolson scheme, is consistent with the results on spatial discretization, as they correspond to one-sided and central-differencing, respectively.

We now examine the *stability* of these schemes following the definitions established in the previous chapter. For a general θ -scheme we solve the linear eigenproblem

$$\frac{dU}{dt} = \lambda U, \quad Re(\lambda) < 0, \tag{6.11}$$

which is discretized as follows

$$\frac{U^{n+1} - U^n}{\Delta t} = \theta \lambda U^{n+1} + (1 - \theta) \lambda U^n.$$

The solution error also satisfies the above equation and thus

$$\epsilon^{n+1}(1 - \theta \lambda \Delta t) = \epsilon^n[1 + (1 - \theta)\lambda \Delta t]$$

SO

$$\epsilon^{n+1} = \epsilon^n \frac{1 + (1 - \theta)\lambda \Delta t}{1 - \theta \lambda \Delta t}.$$

The scheme is absolutely stable if

$$|1 + (1 - \theta)\lambda \Delta t| \le 1 - \theta \lambda \Delta t, \ (Re(\lambda) < 0)$$

or

or

$$\begin{cases} 0 \ge \lambda \Delta t \ \forall \ \Delta t \ge 0 \\ -1 + \theta \lambda \Delta t \le 1 + (1 - \theta) \lambda \Delta t \\ \Rightarrow \boxed{-(1 - 2\theta)\lambda \Delta t \le 2} \end{cases}$$

- For $0 \le \theta < 1/2 \Rightarrow \Delta t \le \frac{-2}{(1-2\theta)\lambda}$, which implies conditional stability.
- For $\theta \geq 1/2$ we obtain unconditional stability. Therefore, both the Crank-Nicolson $(\theta = 1/2)$ and the Euler-backward $(\theta = 1)$ are unconditionally stable schemes. The Euler-backward scheme converges monotonically and damps high frequency components rapidly whereas the Crank-Nicolson scheme converges in an oscillatory manner.

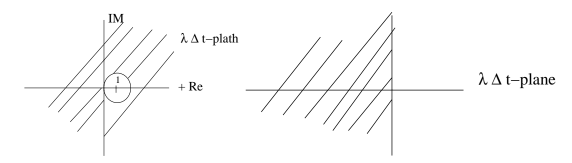


Figure 6.15: Regions of stability for the Euler-backward (left) and Crank-Nicolson (right) schemes.

The stability regions of both the Euler-backward and Crank-Nicolson schemes include the entire half-left plane in the complex plane, and such schemes are referred to as **A-stable**. This region can also be obtained using the theory of characteristic polynomials of section 5.2 in the previous chapter. For example, for the Crank-Nicolson scheme, we construct the polynomial:

$$\Pi_{CN} = (z - 1) - \frac{\lambda \Delta t}{2}(z + 1) = \left(1 - \frac{1}{2}\lambda \Delta t\right)z - \left(1 + \frac{1}{2}\lambda \Delta t\right)$$
$$z = \frac{1 + 1/2\lambda \Delta t}{1 - 1/2\lambda \Delta t} = \frac{2 + \lambda \Delta t}{2 - \lambda \Delta t}.$$

For $Re(\lambda) \leq 0$, we obtain absolute stability in the entire half-left plane as shown in figure 6.15.

Forward versus Backwards Differentiation

In implicit temporal discretizations, it is possible either to construct the backwards difference of the left-hand-side of equation (6.9) or to expand its right-hand-side over several time levels. This is demonstrated in the next two examples:

• Third-order Backwards Differentiation method:

$$\frac{U^{n+1} - \frac{1}{11}(18U^n - 9U^{n-1} + 2U^{n-2})}{\Delta t} = \frac{6}{11}\mathcal{F}^{n+1}.$$

Here the left-hand-side is expanded backwards in time whereas the right-hand-side is evaluated at the current time level. In section 6.2.2 we will see that such schemes have very interesting stability properties.

• Third-order Adams-Moulton method:

$$\frac{U^{n+1} - U^n}{\Delta t} = \frac{5}{12} \mathcal{F}^{n+1} + \frac{8}{12} \mathcal{F}^n - \frac{1}{12} \mathcal{F}^{n-1}.$$

Here, the right-hand-side is formed by evaluating it at the current as well as at previous time levels. Typically, the coefficients of the Adams-Moulton method are smaller than the coefficients of the Adams-Bashforth method that we presented in section 5.2. This, in turn, means that they correspond to a lower truncation error as well as round-off error. Also, for the same accuracy the Adams-Moulton family employs fewer points than the Adams-Bashforth family, but it is implicit and thus computationally more complex.

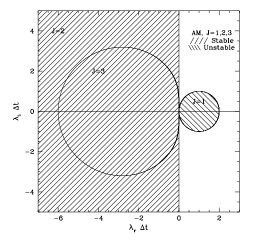


Figure 6.16: Stability diagrams for the first three members of the (implicit) Adams-Moulton family. Note that J = 1 corresponds to the Euler-backward discretization.

The region of stability for the implicit Adams-Moulton can be constructed using the characteristic polymonials, see section 5.2. By comparison of the plots in figure 6.16 with the plots in figure 5.21 we see that the stability regions of the Adams-Moulton family is larger by a factor of about ten than that of the Adams-Bashforth family.

6.2.1 Fundamental Theorems for Multi-Step Methods

We summarize here without proof two basic theorems of multi-step methods originally proposed by Dahlquist [20]. They relate accuracy, stability, and number of steps:

I. First Dahlquist Stability Barrier

The order of accuracy p of a stable k-step (linear) multi-step formula satisfies

$$p \le \left\{ \begin{array}{ll} k+2, & k \text{ even} \\ k+1, & k \text{ odd} \\ k, & \text{explicit form} \end{array} \right\}$$

For example, the four-step Adams-Moulton formula corresponds to sixth-order accuracy. Similarly, the Crank-Nicolson one-step method is of second-order accuracy, and the explicit three-step Adams-Bashforth method is of third-order accuracy. Notice that the highest order accuracy obtainable with a k-step method is 2k, however, such a method is unstable.

II. Second Dahlquist Stability Barrier

The order of accuracy of an explicit A-stable multi-step formula satisfies $p \leq 2$. An explicit multi-step formula cannot be A-stable.

Notice that of all second-order A-stable methods, the one with the smallest truncation error is the Crank-Nicolson scheme. It is also possible to get around the restriction $p \leq 2$ for A-stability by use of Richardson's extrapolation, which is a similar to the Romberg procedure discussed in section 4.2.

6.2.2 Stability of Stiff ODEs

A "stiff" equation has a slowly varying solution combined with rapidly decaying transients. Typically, of interest is the long term (asymptotic) solution, however the initial transients cause severe problems in stability. There are a plethora of physical problems that such behavior is encountered, e.g., in most coupled-domain problems, such as aero-acoustics, combustion, flow-structure interactions, electric circuits, etc. We illustrate this behavior with the following example

$$\frac{dU}{dt} = -1000(U - t^3) + 3t^2, \quad U(0) = 1,$$

which has the exact solution

$$U_{ex}(t) = e^{-1000t} + t^3,$$

plotted in figure 6.17. Notice that the boundary layer term is important only initially.

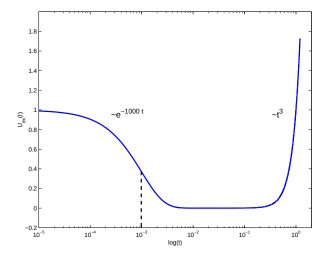


Figure 6.17: Stiff solution of an ordinary differential equation. A boundary layer of thickness 0.001 is present. Notice the logarithmic horizontal scale.

Let us first attempt to employ the Euler-forward explicit scheme

$$U^{n+1} = U^n - 1000\Delta t[U^n - (t^n)^3] + 3\Delta t(t^n)^2$$

with error

$$\epsilon^{n+1} = \epsilon^n (1 - 1000 \Delta t).$$

If

$$\Delta t = 0.1 \Rightarrow \epsilon^{n+1} = 99\epsilon^n$$

thus we loose two decimal points in each time step, yet the scheme is generally stable according to the corresponding definition, i.e.,

$$\epsilon^n = (1 - 1000\Delta t)^n \epsilon^0 < e^{1000\Delta t n} |\epsilon^0|$$

or

$$\epsilon^n < e^{1000t} |\epsilon^0|, \quad \forall \, \Delta t.$$

However, the error bound constant is e^{1000t} , which is so large at t=1 that this discretization is totally impractical even though it is stable!

For useful results we require absolute stability, i.e.,

$$|1 - 1000\Delta t| < 1 \Rightarrow \Delta t = 0.002,$$

that is we require a very small Δt to resolve the transient e^{-1000t} rather than the long-term solution t^3 .

If we now employ the implicit Euler-backward scheme we obtain

$$U^{n+1} = U^n - 1000\Delta t(U^{n+1} - (t^{n+1})^3) + 3\Delta t(t^{n+1})^2$$

with error

$$\epsilon^{n+1} = \epsilon^n / (1 + 1000 \Delta t).$$

k	2	3	4
β_0	2/3	6/11	12/25
α_1	4/3	18/11	48/25
α_2	-1/3	-9/11	-36/25
α_3	_	2/11	16/25
α_4			-3/25

Table 6.3: Weights for the first four stiffly stable schemes.

If $\Delta t = 0.1$ then

$$\epsilon^{n+1} = \epsilon^n/101.$$

Therefore, the error in the Euler-backward scheme decreases as rapidly as it increases for the Euler-forward method. For large values of Δt the boundary layer is inaccurate but we capture the long-term solution. Clearly, an A-stable scheme would solve the stiffness problem, however it is too restrictive. Ideally, for these problems we require (see figure 6.18):

- Absolute stability for the transient behavior.
- General stability for the long-term behavior.

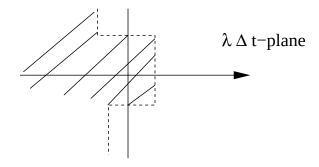


Figure 6.18: Required region of stability for discretizations of stiff equations.

To this end, following the work of Gear [42], we introduce a general stiffly stable scheme of order k^{th} of the form

$$U^{n+1} = \sum_{j=0}^{k} \alpha_{j+1} U^{n-j} + \Delta t \beta_0 \mathcal{F}^{n+1}$$

for the general initial value problem described by equation (6.9). The coefficients are obtained by matching appropriate Taylor expansions and are given in table 6.3.

For stiffly stable schemes *stability studies* are more complicated as the $\rho(z)$ polynomials are non-trivial. For example the third-order scheme corresponds to:

$$\rho(z) = \frac{1}{11}(11z^3 - 18z^2 + 9z - 2)$$

$$\sigma(z) = \frac{6}{11}z^3.$$

They are stable for order $1 \le k \le 6$ and generally unstable for 6 < k < 12 (with some exceptions); they are definitely unstable for $k \ge 12$ (see proof in [53]). Typical stability regions for the first three members of the family are plotted in figure 6.19.

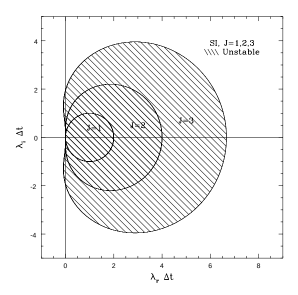


Figure 6.19: Region of stability for stiffly stable schemes for the first three members. Note that the first member corresponds to the Euler-backward scheme.

6.2.3 Second-Order Initial Value Problems

Next we consider second-order initial value problems of the form

$$U_{tt}(t) = \mathcal{F}(t, U(t), U_t(t)), \quad U(0) = U_0, \ U_t(0) = Z_0,$$
 (6.12)

where U_0 and Z_0 are given. Such problems are encountered in flow-structure interactions, wave propagation, etc. They can be reduced to a first-order system by the simple substitution

$$U_t = Z, \quad Z_t = \mathcal{F}(t, U, Z),$$

and thus one of the explicit or implicit algorithms that we have presented previously can be used for the above system.

A particularly successful approach is the **Newmark method**, which is second-order accurate and unconditionally stable for certain values of its parameters. We explain this method further, starting with the Taylor expansion of $U^{n+1} \equiv U(t + \Delta t)$, i.e.,

$$U^{n+1} = U^n + \Delta t U_t^n + \Delta t^2 \left(\beta U_{tt}^{n+1} + (\frac{1}{2} - \beta) U_{tt}^n \right) + \mathcal{O}(\Delta t^3),$$

where β is a parameter that determines the degree of implicit dependence. Upon substitution from equation (6.12), we obtain

$$U^{n+1} = U^n + \Delta t U_t^n + \Delta t^2 \left(\beta \mathcal{F}^{n+1} + \left(\frac{1}{2} - \beta \right) \mathcal{F}^n \right) + \mathcal{O}(\Delta t^2).$$

We also expand similarly the first derivative $U_t^{n+1} \equiv U_t(t+\Delta t)$ and substitute from equation (6.12) to arrive at

$$U_t^{n+1} = U_t^n + \Delta t \left(\gamma \mathcal{F}^{n+1} + (1 - \gamma) \mathcal{F}^n \right) + \mathcal{O}(\Delta t^2),$$

where again γ is a parameter that determines the degree of implicit dependence.

We can now make the substitution

$$Z(t) = U_t(t)$$

to obtain the first-order system

$$U^{n+1} = U^n + \Delta t Z^n + \Delta t^2 \left(\beta \mathcal{F}^{n+1} + (\frac{1}{2} - \beta) \mathcal{F}^n \right) + \Delta t T_u$$

$$Z^{n+1} = Z^n + \Delta t \left(\gamma \mathcal{F}^{n+1} + (1 - \gamma) \mathcal{F}^n \right) + \Delta t T_z$$

The leading terms in the (local) truncation errors T_u and T_z are

$$T_u = \left(\frac{1}{6} - \beta\right) \Delta t^2 U_{ttt}^n + \mathcal{O}(\Delta t^3)$$

$$T_z = \left(\frac{1}{2} - \gamma\right) \Delta t U_{ttt}^n + \mathcal{O}(\Delta t^2).$$

Therefore, it is clear that in order to obtain a second-order accurate scheme we need to choose

$$\gamma = \frac{1}{2} \, .$$

If $\gamma \neq \frac{1}{2}$ then we have a first-order only scheme. The choice of the parameter β will depend on the stability of the scheme. In particular, by considering the corresponding second-order linear oscillator

$$U_{tt} + \omega^2 U = 0,$$

we can obtain the stability requirements which for $\gamma = \frac{1}{2}$ reduce to:

$$\beta \geq \frac{1}{4}$$
.

Finally, we can combine the two equations of the system above to write a single statement for the Newmark scheme for the special case $\gamma = \frac{1}{2}$ and $\beta = \frac{1}{4}$:

$$U^{n+2} - 2U^{n+1} + U^n = \Delta t^2 \left(\frac{1}{4} \mathcal{F}^{n+2} + \frac{1}{2} \mathcal{F}^{n+1} + \frac{1}{4} \mathcal{F}^n \right).$$

In practice, it is more convenient to use the Newmark scheme in a system form instead of the single statement above.

6.2.4 How to March in Time

We summarize here some of the most important features of the explicit and implicit time discretizations, based on which we can decide what specific time-integrator to use in applications:

- The Rung-Kutta method is the easiest to use and is often a good choice, although somewhat inefficient. Low storage and TVB (Total Variation Bounded) schemes are useful in practice. Boundary conditions at intermediate stages may be an issue as they may affect accuracy.
- Multi-step methods (e.g., Adams-Bashforth) are often used in discretizing convective contributions. They require a lot of memory, but they are efficient. The high-order versions exhibit very low dispersion errors. In an unstable multi-step time integration the error has oscillatory form.
- Leap-frog schemes are good for problems with complex eigenvalues.
- **Predictor-corrector** methods are good except when it is necessary to change the step size. It is also easy to estimate time errors. They are less stable than implicit.
- Stiff equations require implicit methods. For efficiency, **stiffly stable** methods (i.e., backwards differentiation) are recommended.
- The truncation error of a k-order stiffly stable method is (1/k) compared to the corresponding Adams-Moulton method.
- Typical instabilities appear in a spontaneous (explosive) way it only takes a few time steps for the solution to blow up!

6.3 Homework Problems

- 1. Consider a 3-point one-dimensional stencil, consisting of the points (i) and $(i \pm 1)$. Construct the highest order difference formula of the second derivative at the point (i), which does not involve any other second-order derivatives. You may include first-order derivatives, however, as well as function values at the three points. Also, compute the truncation error. What is the computational work to compute such a derivative on an N-point grid with periodic boundary conditions?
- 2. Write a C++ program to compute first-order and second-order derivatives with high-accuracy (greater than two) using the implicit formulas. More specifically, design a computer library of one-dimensional implicit formulas according to the text in section 6.1.2. For the families of different orders allow the user to input the proper parameter in order to select different members of the family.
- 3. Solve numerically the elliptic Helmholtz equation in the square domain $[-1, 1] \times [-1, 1]$, i.e.,

$$\nabla^2 U - \lambda U = \sin \frac{\pi x}{2} \sin \frac{\pi y}{2}; \quad \lambda = 1000$$

using the Padé scheme of fourth-order. First use a uniform grid of 50×50 grid points and then an appropriate non-uniform grid of 25×25 grid points and 50×50 grid points. Compare the errors in the three solutions. What do you observe? Also, justify your particular selection of the non-uniform grid.

4. Solve numerically the elliptic Helmholtz equation in the wedge-domain of figure 6.20, i.e.,

$$\nabla^2 U - \lambda U = F(x, y), \quad \lambda = 1$$

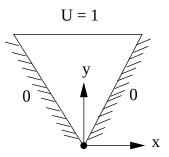


Figure 6.20: Wedge domain.

- (a) First, using second-order finite difference explicit differentiation.
- (b) Second, using fourth-order finite difference implicit differentiation. Consider different functions F(x, y) and plot your results in terms of iso-contours of U(x, y).
- 5. (Sherman-Morrison formula)

Prove that for a non-singular matrix **T**

$$(\mathbf{T} - \mathbf{a}_1 \mathbf{a}_2^T)^{-1} = \mathbf{T}^{-1} - b^{-1} \mathbf{T}^{-1} \mathbf{a}_1 \mathbf{a}_2^T \mathbf{T}^{-1},$$

where

$$b = 1 + \mathbf{a}_2^T \mathbf{T}^{-1} \mathbf{a}_1 \neq 0.$$

Now consider the Thomas algorithm for the problem with periodic boundary conditions (values c and d) and write

$$\mathbf{A} = \mathbf{T} + \mathbf{a}_1 \mathbf{a}_2^T,$$

where **T** is the tridiagonal submatrix and $\mathbf{a}_1 = c\mathbf{e}_1 + d\mathbf{e}_2$ and also $\mathbf{a}_2 = \mathbf{e}_1 + \mathbf{e}_2$.

- (a) Use the Sherman-Morrison formula to obtain the solution to $\mathbf{A}\mathbf{x} = \mathbf{b}$ using only tridiagonal solvers. What is the dominant cost to invert such matrices \mathbf{A} ?
- (b) Write the forward substitution and backward substitution using full-recursive doubling which will reduce the cost to $\mathcal{O}(\log n)$ for the solves.
- 6. Formulate a penta-diagonal solver using the same methodology as used for the Thomas algorithm.
 - (a) Write a C++ function to solve $\mathbf{A}\mathbf{x} = \mathbf{b}$ where \mathbf{A} is penta-diagonal.
 - (b) Solve the one-dimensional heat equation

$$\frac{d^2T}{ds^2} = \sin 2\pi s, \ s \in [0, 1]$$

with u(0) = u(1) = 0.0 using a five-point explicit stencil.

Use more than one grid and compare convergence rates as well as computational times.

- 7. Add timing to the parallel Thomas algorithm function. Using the three-point stencil to approximate second derivatives, formulate the matrix **A** and use it to perform the following tests:
 - (a) For N = 10, 20, 30, and 100 compute the timings when using the serial version.
 - (b) Repeat this experiment using P=2,4,8 processes.
 - (c) What is the parallel speed-up? Does it change depending on the number of points (N) used? Please explain your answer.
- 8. Find the parameter η for which the multi-step implicit time-stepping scheme

$$\frac{U^{n+1} + \eta U^n - (1+\eta)U^{n-1}}{\Delta t} = \frac{1}{2} [-\eta \mathcal{F}^{n+1} + (4+3\eta)\mathcal{F}^n]$$

is of second-order and A-stable discretization to $dU/dt = \mathcal{F}$.

9. Employ a third-order stiffly stable scheme to solve the system

$$\frac{dx}{dt} = -1000x + y$$

$$\frac{dy}{dt} = 999x - 5y$$

assuming that x(0) = 1 and y(0) = 0. Is your solution stable for long-time integration for any Δt ?

10. (Period doubling and chaos)

Consider the homework problem of section 5.3 for period doubling and chaos and use the Crank-Nicolson method with step size h = 0.3, 1 and 5 to obtain numerical solutions. Do you obtain the correct approximation to the solution for all the above values of h? Please justify your answer.

11. Employ the method of undetermined coefficients that we studied in sections 5.1.2 and 6.1.2 for deriving finite differences, to obtain the fourth-order Adams-Bashforth scheme

$$\frac{U^{n+1} - U^n}{\Delta t} = \frac{1}{24} [55\mathcal{F}^n - 59\mathcal{F}^{n-1} + 37\mathcal{F}^{n-2} - 9\mathcal{F}^{n-3}],$$

and also the fourth-order Adams-Moulton scheme

$$\frac{U^{n+1} - U^n}{\Delta t} = \frac{1}{24} [9\mathcal{F}^{n+1} + 19\mathcal{F}^n - 5\mathcal{F}^{n-1} + \mathcal{F}^{n-2}].$$

- 12. Plot the stability region of the fourth-order Adams-Bashforth scheme and compare it with the stability region of the fourth-order Adams-Moulton scheme. What do you observe?
- 13. Plot the stability regions for stiffly-stable schemes of order p=1 to 6 and contrast them against the stability diagrams of Runge-Kutta for order p=1 to 4.
- 14. (Evolution of an ecosystem)

In this problem we revisit the Lotka-Volterra equations for the evolution of an ecosystem that we considered in section 5.3 using explicit methods.

Can you solve this system by using an implicit scheme, e.g. Crank-Nicolson? What do you observe? Please compare and discuss your answers and also comment on the computational complexity of the implicit versus explicit approaches.

Chapter 7

Relaxation: Discretization and Solvers

In this chapter we present discretizations for mixed initial value/boundary value problems (IVP/BVP) as well as relaxation iterative solvers associated with such discretizations. The analogy between iterative procedures and equations of evolution, especially of parabolic type (diffusion), was realized about two centuries ago, but a rigorous connection was not established until the mid 1950s.

In the following, we first consider various mixed discretizations, and subsequently we derive some of the most popular iterative solvers. Our emphasis will be on parallel computing: A good algorithm is not simply the one that converges faster but also the one that is parallelizable. The Jacobi algorithm is such an example; forgotten for years in favor of the Gauss-Seidel algorithm, which converges twice as fast for about the same computational work, it was re-discovered in the last two decades as it is trivially parallelizable, and to-day is used mostly as a preconditioner for multigrid methods. The Gauss-Seidel algorithm, although faster on a serial computer, is not parallelizable unless a special multi-color algorithm is employed, as we explain in section 7.2.4. Based on these two basic algorithms, we present the multigrid method that exploits their good convergence properties but in a smart adaptive way.

On the parallel computing side, we introduce three new commands: MPI_Gather , $MPI_Allgather$, and $MPI_Scatter$. Both MPI_Gather and $MPI_Allgather$ are used for gathering information from a collection of processes. $MPI_Scatter$ is used to scatter data from one process to a collection of processes. In addition to providing syntax and usage information, we present the applicability of the gathering functions in the parallel implementation of the Jacobi method.

7.1 Discrete Models of Unsteady Diffusion

We introduce here unsteady diffusion, a physical process, in order to model relaxation to equilibrium and thus convergence to a steady-state, i.e., a fixed point. Let us consider the one-dimensional *parabolic* equation

$$\begin{cases}
\frac{\partial \Theta}{\partial t} = \kappa \frac{\partial^2 \Theta}{\partial x^2}; & 0 \le x \le 1 \\
\Theta(0, t) = \Theta(1, t) = 0 \\
\Theta(x, 0) = \sin \pi x = \Theta_0(x)
\end{cases}$$
(7.1)

Here we specify both initial conditions as well as boundary conditions since this is a mixed initial value-boundary value problem (IVP/BVP). The dependent variable $\Theta(x,t)$ may represent the temperature and correspondingly κ can be interpreted as the thermal diffusivity. For the sinusoidal initial solution form specified, we can obtain the exact solution using separation of variables, i.e.,

$$\Theta(x,t) = e^{-\kappa \pi^2 t} \sin \pi x.$$

This solution decays to a steady state, which, in this particular case in the absence of any heat sources, is zero. This is shown graphically in the sketch of figure 7.1.

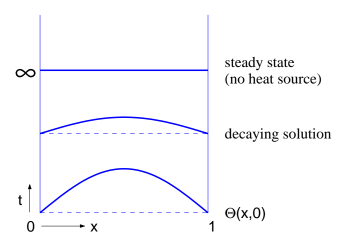


Figure 7.1: The initial solution specified at t=0 decays exponentially to zero (equilibrium state).

We are dealing, therefore, with a continuous diffusion model that produces a solution which relaxes exponentially to the equilibrium state, i.e. the steady-state. Next, we will construct corresponding discrete models.

7.1.1 Temporal and Spatial Discretization

We first discretize the differential equation in time while keeping the spatial derivative continuous. Following a multi-step discretization we obtain

$$\frac{\Theta^{n+1} - \Theta^n}{\theta \Delta t} = \mathcal{F}\left(\frac{\partial^2 \Theta^{n+1}}{\partial x^2}, \frac{\partial^2 \Theta^n}{\partial x^2}, \frac{\partial^2 \Theta^{n-1}}{\partial x^2}, \dots\right).$$

Here \mathcal{F} represents a linear combination of its arguments, and its form is determined by the time-stepping methods as in the discretization of ODEs in sections 5.2 and 6.2. For example, following the θ -method we obtain the semi-discrete equation

$$\frac{\Theta^{n+1} - \Theta^n}{\kappa \Delta t} = \theta \frac{\partial^2 \Theta^{n+1}}{\partial x^2} + (1 - \theta) \frac{\partial^2 \Theta^n}{\partial x^2}.$$

Next, we can discretize each spatial derivative at all required time levels on the right-handside of the equation above, to obtain

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\kappa \Delta t} = \theta \Delta_{xx} \Theta_j^{n+1} + (1 - \theta) \Delta_{xx} \Theta_j^n, \quad j = 1, \dots, N - 1$$
(7.2)

$$\Theta_0^{n+1} = 0 = \Theta_N^{n+1} \tag{7.3}$$

$$\Theta_j^0 = \sin \pi x_j \,. \tag{7.4}$$

Specifically, we have employed the second-order three-point stencil

$$\Delta_{xx}\Theta_j \equiv \frac{1}{\Delta x^2} (\Theta_{j+1} - 2\Theta_j + \Theta_{j-1})$$

to approximate the second derivatives. The notation we adopt is as follows:

- Superscripts denote time levels, and
- Subscripts denote spatial location, unless otherwise stated explicitly.

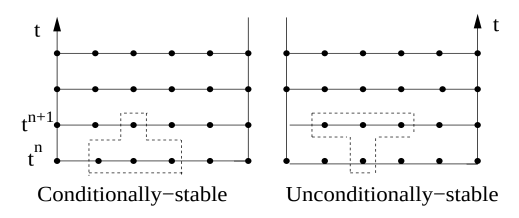


Figure 7.2: Space-time explicit (left) and implicit (right) stencils.

In the above discretization, we can obtain two fundamentally different time-space discretization stencils. Specifically,

- For $\theta = 0$ we obtain a fully parallel but conditionally stable scheme, see figure 7.2 (left).
- For $\theta = 1$, we obtain a partially parallel but unconditionally stable scheme, see figure 7.2 (right).

Next, we will examine in some detail the properties of these and similar *time-space* stencils; we start with accuracy.

7.1.2 Accuracy of Difference Equation

We will denote the convergence rate and correspondingly the accuracy of the time-space stencils using the notation

$$\mathcal{O}(\Delta t^p, \Delta x^q)$$

where p corresponds to temporal accuracy order and q to spatial accuracy order.

We re-write the difference equation in the form

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\kappa \Delta t} - \theta \Delta_{xx} \Theta_j^{n+1} - (1 - \theta) \Delta_{xx} \Theta_j^n = 0,$$

and we expand the time-derivative using Taylor series

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} = \frac{\partial \Theta_j^n}{\partial t} + \frac{1}{2} \Delta t \frac{\partial^2 \Theta_j^n}{\partial t^2} + \mathcal{O}(\Delta t^2).$$

Next, we expand the space-derivatives at time levels n and (n + 1).

$$\Delta_{xx}\Theta_{j}^{n} = \frac{\partial^{2}\Theta_{j}^{n}}{\partial x^{2}} + \underbrace{\frac{\Delta x^{2}}{12} \frac{\partial^{4}\Theta_{j}^{n}}{\partial x^{4}}}_{\mathcal{O}(\Delta x^{2})} + \mathcal{O}(\Delta x^{4})$$

$$\Delta_{xx}\Theta_{j}^{n+1} = \Delta_{xx}\Theta_{j}^{n} + \Delta t \Delta_{xx} \frac{\partial \Theta_{j}^{n}}{\partial t} + \mathcal{O}(\Delta t^{2})$$

$$= \frac{\partial^{2}\Theta^{n}}{\partial x^{2}} + \frac{\Delta x^{2}}{12} \frac{\partial^{4}\Theta_{j}^{n}}{\partial x^{4}} + \Delta t \frac{\partial^{2}}{\partial x^{2}} \frac{\partial \Theta_{j}^{n}}{\partial t} + \mathcal{O}(\Delta t \Delta x^{2}) + \mathcal{O}(\Delta x^{4})$$

Assuming it is permissible to perform the following operation

$$\frac{\partial \Theta}{\partial t} = \kappa \frac{\partial^2 \Theta}{\partial x^2} \Rightarrow \frac{\partial^2 \Theta}{\partial t^2} = \kappa \frac{\partial^2}{\partial x^2} \frac{\partial \Theta}{\partial t} = \kappa \frac{\partial^4 \Theta}{\partial x^4},$$

and also setting $\kappa = 1$ for simplicity, we obtain the difference equation

$$\frac{\partial \Theta_{j}^{n}}{\partial t} + \frac{1}{2} \Delta t \frac{\partial^{4} \Theta_{j}^{n}}{\partial x^{4}} + \mathcal{O}(\Delta t^{2}) - \theta \frac{\partial^{2} \Theta_{j}^{n}}{\partial x^{2}} - \theta \frac{\Delta x^{2}}{12} \frac{\partial^{4} \Theta_{j}^{n}}{\partial x^{4}} - \theta \Delta t \frac{\partial^{4} \Theta_{j}^{n}}{\partial x^{4}} + \mathcal{O}(\Delta t^{2}) + \mathcal{O}(\Delta t^{3})$$

$$-(1 - \theta) \frac{\partial^{2} \Theta_{j}^{n}}{\partial x^{2}} - (1 - \theta) \frac{\partial^{4} \Theta_{j}^{n}}{\partial x^{4}} \frac{\Delta x^{2}}{12} + \mathcal{O}(\Delta x^{4}) + \mathcal{O}(\Delta t \Delta x^{2}) = 0.$$

We can now collect all terms and substitute in the difference equation to obtain the *equivalent* differential equation:

$$\left(\frac{\partial \Theta_{j}^{n}}{\partial t} - \frac{\partial^{2} \Theta_{j}^{n}}{\partial x^{2}}\right) \underbrace{-\Delta t(\theta - 1/2) \frac{\partial^{4} \Theta_{j}^{n}}{\partial x^{4}}}_{T_{i}^{n} = \text{truncation error}} + \underbrace{\frac{\mathcal{O}(\Delta x^{2}) + \mathcal{O}(\Delta t^{2}) + \mathcal{O}(\Delta x^{2}, \Delta t^{2})}{+\mathcal{O}(\Delta t \Delta x^{2}) = 0}}_{\text{H.O.T.}}.$$

(where H.O.T. denotes higher order terms). For *consistency* we require that the truncation error

$$T_i^n \to 0$$
, for Δt , $\Delta x \to 0$,

which is true for this difference equation. The truncation error also reveals the order of accuracy. Specifically:

• For $\theta = 0$ or 1 the truncation error is

$$T \sim \mathcal{O}(\Delta t, \Delta x^2),$$

as expected since this corresponds to the Euler-forward and Euler-backward timestepping scheme, respectively.

• For $\theta = 1/2$ the truncation error is

$$T \sim \mathcal{O}(\Delta t^2, \Delta x^2),$$

which corresponds to a second-order (Crank-Nicolson) time-stepping scheme.

7.1.3 Stability of Difference Equation

The Lax *Equivalence Theorem* for linear partial differential equations (PDEs) states that if a difference equation is consistent with a PDE, i.e.,

$$||T_i^n||_2 \to 0, \quad \Delta t, \Delta x \to 0,$$

then stability of the difference equation is a sufficient and necessary condition for convergence.

We have already seen in section 5.2 that for the difference equation resulting from an ODE of the form

$$\frac{dU}{dt} = \lambda U,$$

we need to march with a time step smaller than a critical value, i.e.

$$\Delta t < \Delta t_c(\lambda),$$

where here in equation (7.1), λ corresponds to the eigenvalues of the Laplacian operator since we can write

$$\frac{\partial^2 \Theta}{\partial^2 x} = \lambda \Theta$$
, and in general $\nabla^2 \Theta = \lambda \Theta$.

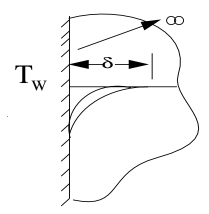


Figure 7.3: Propagation of heat in a semi-infinite medium.

These eigenvalues depend on the spatial discretization parameter Δx , so we expect that for conditional stability we would have

$$\Delta t < \mathcal{F}(\Delta x)$$
.

The specific form of the above right-hand-side will depend on the spatial differential operator.

We can also use physical intuition to justify the constraint $\Delta t < \mathcal{F}(\Delta x)$. To this end, let us consider a semi-infinite plate of thermal diffusivity κ and model the heat propagation through this medium by equation (7.1). By performing dominance balance on this equation assuming a characteristic spacing Δx and characteristic time scale Δt_c we obtain

$$\frac{\Theta}{\Delta t_c} \sim \kappa \frac{\Theta}{\Delta x^2} \Rightarrow \Delta t_c \sim \frac{\Delta x^2}{\kappa}$$
.

In order to resolve that time scale we require that

$$\Delta t < \Delta t_c \Rightarrow \Delta t < C \frac{\Delta x^2}{\kappa}$$

where $C \sim \mathcal{O}(1)$, that is C is a constant of order one. This inequality can be re-written as

$$D \equiv \frac{\kappa \Delta t}{\Delta x^2} < C, \tag{7.5}$$

which simply states that the non-dimensional left-hand-side, termed diffusion number, should be less than a constant of order unity for proper temporal resolution of the heat propagation in this isotropic medium.

In the following, we will demonstrate three different ways of obtaining the exact stability constraint, that is, the value of the constant C. We will start from the more intuitive perturbation method, and we will proceed with the classical von Neumann analysis, and a more general method based on matrices.

Discrete Perturbation Stability Analysis

This method is based on the physics of propagation of perturbation. A discrete perturbation in $\Theta(x,t)$ is introduced into the equation at an arbitrary point, and its effect is followed in time. Stability is indicated if the perturbation dies out [78].

To demonstrate this approach we employ for simplicity the Euler-forward time-stepping scheme. Let us introduce a disturbance ϵ into the difference equation at time level n at grid point i, and follow its evolution in time, i.e.,

$$\frac{\Theta_i^{n+1} - (\Theta_i^n + \epsilon)}{\Delta t} = \underbrace{\kappa}_{\text{thermal diffusivity}} \frac{\Theta_{i+1}^n - 2(\Theta_i^n + \epsilon) + \Theta_{i-1}^n}{\Delta x^2}.$$

Let us also assume that at time level (n) we have $\Theta_i^n = 0$, and thus

$$\frac{\Theta_i^{n+1} - \epsilon}{\Delta t} = -2 \frac{\kappa \epsilon}{\Delta x^2} \,.$$

The initial disturbance at time level (n) is shown schematically in figure 7.4.

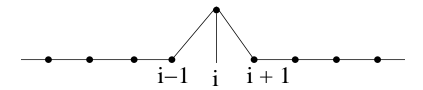


Figure 7.4: Initial disturbance introduced at time level n.

The disturbance distribution at the next time level is determined by

$$\Theta_i^{n+1} = \epsilon (1 - 2D)$$

where D is the diffusion number, see equation (7.5). We note here that Θ_i^{n+1} represents the evolving disturbance (since $\Theta_i^n = 0$) of the initial disturbance ϵ . We now require that

$$\left| \frac{\Theta_i^{n+1}}{\epsilon} \right| \le 1 \Rightarrow -1 \le 1 - 2D \le 1 \tag{7.6}$$

and therefore

$$D < 1$$
.

If we require that no *overshoots* or *undershoots* appear in the solution, that is we want to prevent spurious oscillations, we will need to impose the *same sign* constraint, i.e.,

$$\Theta_i^{n+1}/\epsilon \ge 0$$
,

which in turn leads to a more strict upper limit for the diffusion number, i.e.,

$$D < 1/2$$
.

At the (n+1) time level we can obtain the new perturbation at all grid points from the difference equation: $\Theta_{i+1}^{n+1} = \epsilon D$, thus

$$\begin{array}{lll} \Theta_i^{n+2} & = & \Theta_i^{n+1} + D[\Theta_{i+1}^{n+1} + \Theta_{i-1}^{n+1} - 2\Theta_i^{n+1}] \\ & = & \epsilon(1-2D) + D[\epsilon D + \epsilon D - 2\epsilon(1-2D)] \,, \end{array}$$

and therefore

$$\Theta_i^{n+2} = \epsilon (1 - 4D + 6D^2).$$

We require also at this time level that disturbances do not grow, that is we impose

$$\left| \frac{\Theta_i^{n+2}}{\epsilon} \right| \le 1 \Rightarrow -1 \le 1 - 4D + 6D^2 \le 1,$$

which is satisfied for

$$D < 2/3$$
.

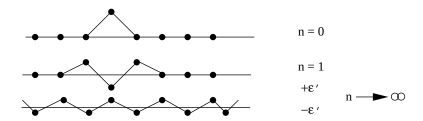


Figure 7.5: Evolution of disturbance at different time levels.

Graphically, the evolution of disturbance for the first few time steps is described in figure 7.5, and after many time steps the asymptotic spreading of a single perturbation approaches a $2\Delta x$ wave. Therefore, the most dangerous disturbance is produced asymptotically and corresponds to an amplitude ϵ' . The difference equation then gives

$$\Theta_i^{n+1} = \epsilon' + D(-\epsilon' - 2\epsilon' - \epsilon') = \epsilon'(1 - 4D).$$

For stability we require that

$$\left| \frac{\Theta_i^{n+1}}{\epsilon'} \right| \le 1 \Rightarrow -1 \le 1 - 4D \le 1,$$

and from this we deduce that

$$D \leq 1/2$$
.

If we now compare this constraint on the diffusion number derived from the asymptotic spreading of the disturbance with the initial requirement of preventing spurious oscillations we conclude that:

• The long-time stability requirement is equivalent to the requirement for zero overshoots for an isolated disturbance.

The von Neumann Stability Analysis

This approach was developed by John von Neumann in the early 1940s at Los Alamos, and it is the most popular method. It assumes that the domain is infinite in extent or periodic and employs the discrete analog of Fourier series expansions in the form

$$\Theta_j^n = \sum_{k=1}^{N-1} a_k^n \sin \pi k x_j$$

with $\Theta_0^n = \Theta_N^n = 0$ and $x_j = j \cdot \Delta x$; a_k^n are the Fourier-sine coefficients.

The main idea is to reduce the difference equation for Θ_j^n into an uncoupled ordinary difference equation for a_k^n , and subsequently require that

$$|a_k^{n+1}| \le |a_k^n|$$

in order to guarantee absolute stability.

To proceed we use the general identity

$$\Theta_{j-1} - 2\Theta_j + \Theta_{j+1} = \sum_{k=1}^{N-1} a_k [\sin \pi k \Delta x (j-1) - 2\sin \pi k \Delta x j + \sin \pi k \Delta x (j+1)]$$

$$= \sum_{k=1}^{N-1} a_k \sin \pi k \Delta x j (2\cos \pi k \Delta x - 2)$$

$$= -\sum_{k=1}^{N-1} \sigma_k a_k \sin \pi k \Delta x j,$$

where $\Delta x = 1/N$ and

$$\sigma_k \equiv 2(1 - \cos \pi k \Delta x)$$
 where $0 < \sigma_{\kappa} < 4$.

We analyze the time-space stencil corresponding to the θ -method again, and upon substitution in the difference equation (7.2) we obtain

$$\Theta_{j}^{n+1} - \Theta_{j}^{n} = \theta D(\Theta_{j-1}^{n+1} - 2\Theta_{j}^{n+1} + \Theta_{j+1}^{n}) + D(1-\theta)(\Theta_{j-1}^{n} - 2\Theta_{j}^{n} + \Theta_{j+1}^{n}),$$

and therefore

$$\sum_{k=1}^{N-1} (a_k^{n+1} - a_k^n) \sin \pi k x_j = D\theta \sum_{k=1}^{N-1} -\sigma_k a_k^{n+1} \sin \pi k x_j + D(1-\theta) \sum_{k=1}^{N-1} -\sigma_k a_k^n \sin k \pi x_j$$

or

$$a_k^{n+1} - a_k^n = -D\theta \sigma_k a_k^{n+1} - D(1-\theta)\sigma_k a_k^n$$

and finally

$$a_k^{n+1} = a_k^n \left(\frac{1 - D(1 - \theta)\sigma_k}{1 + D\theta\sigma_k} \right). \tag{7.7}$$

We can now obtain specific forms for the three main members of the family:

• $\theta = 0$ corresponds to Euler-forward scheme. We obtain

$$a_k^{n+1} = a_k^n (1 - D\sigma_k) \Rightarrow |1 - D\sigma_k| \le 1 \Rightarrow D \le 2/\sigma_k, \forall k$$

since $\max \sigma_k = 4$, $\forall k$, we have that $D \leq 1/2$, as before.

• $\theta = 1/2$ corresponds to Crank-Nicolson scheme. Here we obtain

$$a_k^{n+1} = \frac{1 - \frac{D\sigma_k}{2}}{1 + \frac{D\sigma_k}{2}} a_k^n.$$

This is an unconditionally stable scheme. It damps high-frequency components very weakly, although these components may (physically) decay very rapidly. However, for $\theta = 1/2 + \alpha \Delta t$, where α is a small positive constant, all components of the solution are damped and the method is still formally second-order accurate in time, i.e., $\mathcal{O}(\Delta t^2)$.

• $\theta = 1$ corresponds to Euler-backward scheme. We have

$$a_k^{n+1} = \frac{a_k^n}{1 + D\sigma_k} \,.$$

This is also an unconditionally stable scheme, and it damps high-frequency components very rapidly.

The structure of the instability can be studied by examining the most dangerous mode, which corresponds to

$$k = N - 1 \Rightarrow \sigma_{k, \max} = 4$$
,

and therefore

$$\epsilon \sim \sin \pi (N-1)x$$

with wavelength

$$\frac{2\pi}{\pi(N-1)} \sim \frac{2}{N} = 2\Delta x.$$

Therefore, the error has a "sawtooth" form with wavelength $2\Delta x$, as our previous perturbation analysis also showed.

Matrix Methods for Stability Analysis

This is the most general method and is suitable for studying the effect of boundary conditions on the stability of the difference equation. The main idea is to follow the *method of lines*, where we discretize the space derivatives first while we keep the time derivatives continuous. For example, for the heat equation (7.1) we obtain the vector equation

$$\frac{d\Theta_j}{dt} = \mathbf{A}\Theta_j,$$

where **A** is the discrete operator

$$\mathbf{A} = \frac{\kappa}{\Lambda x^2} (\mathbf{E} - 2\mathbf{I} + \mathbf{E}^{-1})$$

defined in terms of the displacement operator \mathbf{E} , see section 5.1. In this form, we can examine the eigenspectrum of the matrix obtained from the operator \mathbf{A} to determine the stability of the difference equation.

Before we treat the effect of boundary conditions, however, we revisit the heat equation with periodic boundary conditions, and in this context we introduce the matrix stability analysis. To this end, we obtain the fully-discrete form of equation (7.1)

$$\Theta_j^{n+1} = \mathbf{C}\Theta_j^n$$
, $\mathbf{C} = \mathbf{I} + \Delta t \cdot \mathbf{A} = \mathbf{I} + D(\mathbf{E} - 2\mathbf{I} + \mathbf{E}^{-2})$,

where D is the diffusion number. This form corresponds to the Euler-forward scheme ($\theta = 0$) as before. We can recover the von Neumann analysis here by introducing a *single* harmonic perturbation

$$\Theta_j^n = \hat{\Theta}^n e^{ij\phi}$$
 where $\phi_{\kappa} = \phi = k \frac{\pi}{N}$.

Upon substitution, we obtain

$$\hat{\Theta}^{n+1} = \underbrace{\mathbf{G}(\phi)}_{\text{Amplification Matrix}} \hat{\Theta}^n = [G(\phi)]^n \hat{\Theta}^1.$$

For stability $[\mathbf{G}(\phi)]^n$ has to be bounded, and thus the von Neumann necessary condition for stability is

$$\rho(\mathbf{G}) = \max_{k} |\lambda_k| \le 1,$$

where $\rho(\mathbf{G})$ is the spectral radius of the matrix \mathbf{G} . (Actually, the more precise condition is $\leq 1 + \mathcal{O}(\Delta t)$, see [77]). If \mathbf{G} is a normal matrix (i.e., it commutes with its Hermitian conjugate), then the condition $\rho(\mathbf{G}) \leq 1$ is also a sufficient condition.

Example 1 - The Euler-forward scheme: We can clarify this matrix further by considering the Euler-forward scheme, which we have already analyzed with the previous two methods. We first introduce a single harmonic in the equation

$$\Theta_j^{n+1} = \Theta_j^n + \frac{\kappa \Delta t}{\Delta r^2} (\Theta_{j+1}^n - 2\Theta_j^n + \Theta_{j-1}^n),$$

and we have also defined $D \equiv \frac{\kappa \Delta t}{\Delta x^2}$. Here we have a scalar equation, and the amplification factor is

$$e^{ij\phi}G(\phi) = e^{ij\phi} + \frac{\kappa \Delta t}{\Delta x^2} (e^{i(j+1)\phi} - 2e^{ij\phi} + e^{i(j-1)\phi})$$

$$\Rightarrow G(\phi) = 1 + D(e^{i\phi} - 2 + e^{-i\phi})$$

$$\Rightarrow G(\phi) = 1 - 4D\sin^2 \phi/2.$$

Thus, we require that

$$|G(\phi)| < 1 \Rightarrow 0 < D < 1/2,$$

and we recover the condition derived previously with the other two methods.

For the Euler-forward scheme we can also use the amplification factor $G(\phi)$ to analyze the nature of the error. The exact solution we obtained can be re-written as

$$\Theta^n = e^{-i\omega(n\Delta t)} \cdot e^{ikj\Delta x} \cdot \Theta_0^*(k),$$

where $\Theta_0^*(k)$ is the harmonic corresponding to initial data, so

$$\Theta^n = (e^{-i\omega\Delta t})^n e^{ikj\Delta x} \Theta_0^*(k) = G^n e^{ikj\Delta x} \Theta_0^*(k).$$

Thus, $G=e^{-i\omega\Delta t}$ is the exact amplification factor, where $\omega=\omega(k)$ defines the exact dispersion relation. Let

$$\omega = \omega_r + i\omega_i,$$

then

$$G = e^{\omega_i \Delta t} \cdot e^{-i\omega_r \Delta t} = |G|e^{-i\Phi}$$

where $\Phi = \omega_r \Delta t$ and $|G| = e^{\omega_i \Delta t}$. In order to compare the error in amplitude, we define

$$\epsilon_D = \frac{|G|}{e^{\omega_i \Delta t}},$$
 for numerical dissipation.

Similarly, to compare the error in *phase*, we define

$$\epsilon_{\phi} = \frac{\Phi}{\omega_r \Delta t}$$
, for numerical dispersion.

Specifically, for the Euler-forward time-discretization of equation (7.1) we have

$$\omega = -i\kappa k^2 = -iD\frac{\phi^2}{\Delta t}$$
, where $\phi = \frac{k\pi}{N} = k\Delta x$.

Therefore, the dissipation error is

$$\epsilon_D^{EF} = \frac{1 - 4D\sin^2\phi/2}{e^{-D\phi^2}} = \frac{1 - D\phi^2 + D\phi^4/12 + \dots}{1 - D\phi^2 + (D^2\phi^4/2) + \dots}$$

or

$$\epsilon_D^{EF} \approx 1 - \frac{D^2 \phi^4}{2} + \frac{D \phi^4}{12} + \dots \approx 1 - \frac{\kappa^2 k^4 \Delta t^2}{2} + \frac{\kappa k^4}{12} \Delta t \Delta x^2 - \dots$$

Clearly, for low frequencies ($\phi \approx 0$) there are only small errors, and specifically for D = 1/6 we obtain a high-order scheme, i.e., $\mathcal{O}(\Delta t^2, \Delta x^4)$!

Example 2 - The DuFort-Frankel scheme: This is an example of an *inconsistent scheme*, which sometimes converges and sometimes does not, and this is a very dangerous practice. It involves two previous time levels so the amplification factor is now a matrix and not a scalar as before. We first apply the *leap-frog* scheme to equation (7.1), i.e.,

$$\Theta_i^{n+1} - \Theta_i^{n-1} = 2D(\Theta_{i+1}^n - 2\Theta_i^n + \Theta_{i-1}^n).$$

Then, we average in time the term Θ_j^n on the right-hand-side to obtain $\frac{1}{2}[\Theta_j^{n+1} + \Theta_j^{n-1}]$, and upon substitution into the expression above, we have

$$\Theta_j^{n+1} - \Theta_j^{n-1} = 2D(\Theta_{j+1}^n - \Theta_j^{n+1} - \Theta_j^{n-1} + \Theta_{j-1}^n)$$

or

$$\Theta_j^{n+1}(1+2D) = \Theta_j^{n-1}(1-2D) + 2D(\Theta_{j+1}^n + \Theta_{j-1}^n).$$

Let us now introduce a new variable $z^n = \Theta^{n-1}$, and recast the above into the system:

$$\begin{array}{rcl} \Theta_{j}^{n+1} & = & \frac{1-2D}{1+2D}z^{n} + \frac{2D}{1+2D}(\Theta_{j+1}^{n} + \Theta_{j-1}^{n}) \\ z_{j}^{n+1} & = & \Theta_{j}^{n} \, . \end{array}$$

In matrix form, if we set $\mathbf{W} = \begin{bmatrix} \Theta_j^n \\ z_j^n \end{bmatrix}$, we have:

$$\mathbf{W}^{n+1} = \mathbf{C}\mathbf{W}^n \text{ where } \mathbf{C} = \begin{bmatrix} \frac{2D(E+E^{-1})}{1+2D} & \frac{1-2D}{1+2D} \\ 1 & 0 \end{bmatrix}.$$

The corresponding amplification matrix becomes

$$\mathbf{G} = \begin{bmatrix} \frac{2D(e^{i\phi} + e^{-i\phi})}{1+2D} & \frac{1-2D}{1+2D} \\ 1 & 0 \end{bmatrix} = \begin{bmatrix} \frac{4D\cos\phi}{1+2D} & \frac{1-2D}{1+2D} \\ 1 & 0 \end{bmatrix}$$

with eigenvalues

$$\lambda_{\pm} = \frac{2D\cos\phi \pm \sqrt{1 - 4D^2\sin^2\phi}}{1 + 2D},$$

and for

$$D > 0 \Rightarrow |\lambda \pm| < 1 \quad \forall \phi.$$

This result, which implies unconditional stability for an explicit scheme, is suspicious! It clearly violates the second stability barrier of Dahlquist, see section 6.2.1. To analyze this, we need to examine convergence, and thus we need to consider the consistency of this scheme. To this end, we obtain the *equivalent differential equation*

$$\Theta_{t} - \kappa \Theta_{xx} = -\frac{\Delta t^{2}}{6} \Theta_{tt} + \kappa \frac{\Delta x^{2}}{12} \left(\frac{\partial^{4} \Theta}{\partial x^{4}} \right) + \frac{\kappa \Delta x^{4}}{360} \left(\frac{\partial^{6} \Theta}{\partial x^{6}} \right) - \kappa \frac{\Delta t^{2}}{\Delta x^{2}} \Theta_{tt} - \frac{\kappa \Delta t^{4}}{12\Delta x^{2}} \left(\frac{\partial^{4} \Theta}{\partial t^{4}} \right) + \dots$$

In the limit as $\Delta t, \Delta x \to 0$ at the same rate the ratio $\Delta t/\Delta x$ is constant, and thus we solve a modified differential equation and not the original equation (7.1). Specifically, we solve

$$\frac{\partial \Theta}{\partial t} + \underbrace{\kappa \left(\frac{\Delta t}{\Delta x}\right)^2 \frac{\partial^2 \Theta}{\partial t^2}}_{\text{extra term}} = \kappa \frac{\partial^2 \Theta}{\partial x^2}.$$

We have a clear violation of the consistency condition! However, in practice one can take the diffusion number $D = \frac{\kappa \Delta t}{\Delta x^2}$ to be constant, and then the DuFort-Frankel scheme is almost consistent since $\Delta t \to 0$ faster than Δx .

7.1.4 Spectrum of the Diffusion Operator

Next, we examine the eigenspectrum of the matrix

$$\mathbf{A} = \frac{\kappa}{\Delta x^2} (\mathbf{E} - 2\mathbf{I} + \mathbf{E}^{-1})$$

corresponding to different types of boundary conditions. Specifically, we consider:

• Dirichlet Boundary Conditions: Let us assume that $\Theta_0(0) = a$; $\Theta_N = b$. Then the equations for the end-points are

$$\frac{d\Theta_1}{dt} = \frac{\kappa}{\Delta x^2} (a - 2\Theta_1 + \Theta_2),$$

and

$$\frac{d\Theta_{N-1}}{dt} = \frac{\kappa}{\Delta x^2} (b - 2\Theta_{N-1} + \Theta_{N-2}).$$

The corresponding linear system is

$$\frac{d\Theta}{dt} = \frac{\kappa}{\Delta x^2} \begin{bmatrix} -2 & 1 & & & \\ 1 & -2 & 1 & & \\ & 1 & -2 & 1 & \\ & & \ddots & \ddots & \\ & & & 1 & -2 \end{bmatrix} \begin{bmatrix} \Theta_1 \\ \vdots \\ \vdots \\ \Theta_{N-1} \end{bmatrix} + \begin{bmatrix} \frac{\kappa a}{\Delta x^2} \\ \vdots \\ 0 \\ \vdots \\ \frac{\kappa b}{\Delta x^2} \end{bmatrix}.$$

• Neumann Boundary Conditions: We assume that $\frac{\partial \Theta}{\partial x}|_{0}=a$ and $\Theta_{N}=b$. For the boundary point with the Neumann condition we have

$$\frac{1}{\Delta x}(\Theta_1 - \Theta_0) = a \Rightarrow \Theta_0 = \Theta_1 - a\Delta x,$$

and thus

$$\frac{d\Theta_1}{dt} = \frac{\kappa}{\Delta x^2} (\Theta_0 - 2\Theta_1 + \Theta_2) = \frac{\kappa}{\Delta x^2} (-\Theta_1 + \Theta_2 - a\Delta x).$$

The corresponding linear system is

$$\frac{d\Theta}{dt} = \frac{\kappa}{\Delta x^2} \begin{bmatrix} -1 & 1 & & & \\ 1 & -2 & 1 & & \\ & 1 & -2 & 1 & \\ & & & \ddots & \\ & & & 1 & -2 \end{bmatrix} \begin{bmatrix} \Theta_1 \\ \vdots \\ \vdots \\ \Theta_{N-1} \end{bmatrix} + \begin{bmatrix} -\frac{\kappa a}{\Delta x} \\ \vdots \\ 0 \\ \vdots \\ \frac{\kappa b}{\Delta x^2} \end{bmatrix}.$$

• Periodic Boundary Conditions: We assume that $\left\{ \begin{array}{l} \Theta_0 = \Theta_N \\ \Theta_{-1} = \Theta_{N-1} \end{array} \right.$

The equations for the boundary points are

$$\frac{d\Theta_0}{dt} = \frac{\kappa}{\Delta x^2} (\Theta_{N-1} - 2\Theta_0 + \Theta_1)$$

$$\frac{d\Theta_{N-1}}{dt} = \frac{\kappa}{\Delta x^2} (\Theta_0 - 2\Theta_{N-1} + \Theta_{N-2})$$

then:

$$\frac{d\Theta}{dt} = \frac{\kappa}{\Delta x^2} \begin{bmatrix} -2 & 1 & & & 1 \\ 1 & -2 & 1 & & \\ & 1 & -2 & 1 & \\ & & & \ddots & \\ 1 & & & 1 & -2 \end{bmatrix} \begin{bmatrix} \Theta_0 \\ \vdots \\ \vdots \\ \vdots \\ \Theta_{N-1} \end{bmatrix}$$

It can be shown (see section 7.3) that the eigenvalues of a general tridiagonal matrix $\mathbf{B}(a, b, c)$ of rank N determined from the condition det $|\mathbf{B}(a, b - \lambda, c)| = 0$ are given by

$$\lambda_j = b + 2\sqrt{(ac)}\cos\frac{j\pi}{N+1}, \ j = 1,\dots, N$$

while the corresponding eigenvectors are:

$$\phi_m^{(j)} = C_j \left(\frac{a}{c}\right)^{(m-1)/2} \sin m \left(\frac{j\pi}{N+1}\right), \ m = 1, \dots, N.$$

Notice that the eigenvectors do not depend on the diagonal element b and that C_j is arbitrary, and it is determined by normalization conditions.

Similarly, for the *periodic case* we have:

$$\lambda_j = b + ae^{-i2\pi j/N} + ce^{i2\pi j/N}$$

$$\phi_m^j = C_j e^{i(2\pi j/N)m}, \text{ where } m = 1, \dots, N$$

which can be obtained directly from the properties of a *circulant* matrix.

Specifically now for the Diffusion operator we have:

• Dirichlet boundary conditions:

$$\lambda_j = \frac{-4\kappa}{\Delta x^2} \sin^2\left(\frac{\pi j}{2(N+1)}\right), \ j = 1, \dots, N.$$

• Periodic boundary conditions:

$$\lambda_j = \frac{-4\kappa}{\Delta x^2} \sin^2\left(\frac{\pi j}{N}\right) .$$

• Neumann/Dirichlet boundary conditions:

$$\lambda_j = \frac{-4\kappa}{\Delta x^2} \sin^2 \frac{(2j-1)\pi}{(2N+1)2} \,.$$

The important conclusion from inspecting these eigenvalues is that they grow as N^2 since $\Delta x \sim 1/N$. We also note that for the Euler-forward scheme we recover again

$$\Delta t < 2/|\lambda| \Rightarrow D \le 1/2$$

as before. Finally, we observe that all type of boundary conditions lead to stable discretizations, unlike the discrete *convection* equation, see chapter 8. Stability here is satisfied if $Re(\lambda_j) \leq 0$, $\forall j$. In addition, if $\lambda_j = 0$ it has to be a simple eigenvalue.

Experiments with Euler-forward and Euler-backward

We will now consider both an Euler-forward and an Euler-backward time integration combined with the second-order finite difference spatial differentiation operator.



First, we present a C++ function which incorporates a central finite difference scheme with the Euler-forward time integration scheme. The function takes as input the number of points in the grid, the diffusion number DN, an array containing the current value of the solution uold, and an array unew into which to place the solution at the new time. Observe that we only update the interior points $1, \ldots, N-2$, leaving the boundary conditions to be handled separately.

Observe that this function is fully explicit, both in space and in time. Given the current value of the solution at a particular time, the solution at the next time step is obtained by finite differencing and explicit time marching. Compare this with the Euler-backward implementation, which uses implicit time-marching.

to be handled separately.

finite difference scheme with the Euler-backward time integration scheme. The function takes as input the number of points in the grid, the diffusion number DN, an array containing the current value of the solution uold, and an array unew into which to place the solution at the new time. To solve the Eulerbackward problem, we use our Thomas Algorithm tridiagonal solver. The tridiagonal system is formed by the second-order finite difference operator. Observe that we only update the interior points $1, \ldots, N-2$ (by passing only (N-2) points to the ThomasAlgorithm function), leaving the boundary conditions

We now present a C++ function which incorporates a central



Programming Note

• Notice the use of '&' in the code above. Recall what this means — the '&' operator gives us "the address of." So, in the expression above, we are passing to the *ThomasAlgorithm* routine the *address* of the second entry of the *uold* and the *unew* array. In figure 7.6, we show the memory addressing for the *unew* array and how the "&" operator can be used to obtain memory addresses for elements of the *unew* array.

As a model problem for the Euler-forward and Euler-backward implementations presented above, we will examine the problem of finding the solution $\Theta(x,t)$ of equation (7.1) on the interval [0,1] with the boundary conditions $\Theta(0,t) = \Theta(1,t) = 0.0$. We present a total of three numerical experiments, two cases using Euler-forward and one case using Eulerbackward.

Experiment 1 – Euler-forward with D = 0.7: For this experiment, we use the parameters as given in table 7.1. The results are presented in figure 7.7. This choice of the

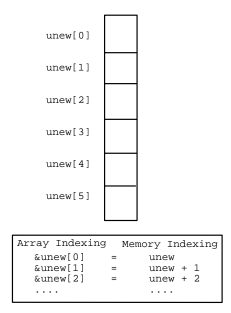


Figure 7.6: Memory addressing for the array unew.

parameter D is clearly above the $D \leq 0.5$ necessary for stability. Observe in figure 7.7 the $2\Delta x$ instability that renders the computation useless in very few iterations.

Parameter	Value
Method	Euler-forward Central Difference
Interval	[0,1]
N	21
Δt	0.00175
D	0.7
Initial Condition	$u(x,0) = \sin(\pi x)$
Boundary Conditions	u(0,t) = u(1,t) = 0

Table 7.1: Parameters used for the simulations presented in figure 7.7.

Experiment 2 – Euler-forward with D = 0.4: For this experiment, we use the parameters as given in table 7.2. The results are presented in figure 7.8. This choice of the parameter D is clearly below the $D \leq 0.5$ necessary for stability. Observe in figure 7.8 that the solution remains stable and agrees quite well with the exact solution for the three different times presented.

Experiment 3 – Euler-backward with D = 1.0: For this experiment, we use the parameters as given in table 7.3. The results are presented in figure 7.9. Because we now use an implicit method for the time integration, we are no longer bound by the $D \le 0.5$ limit imposed for Euler-forward. Observe in figure 7.9 that the solution remains stable and

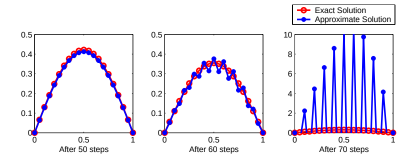


Figure 7.7: Euler-forward/central difference versus exact solution at different times for D = 0.7.

Parameter	Value
Method	Euler-forward Central Difference
Interval	[0,1]
N	21
Δt	0.001
D	0.4
Initial Condition	$u(x,0) = sin(\pi x)$
Boundary Conditions	u(0,t) = u(1,t) = 0

Table 7.2: Parameters used for the simulations presented in figure 7.8.

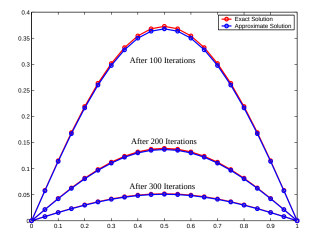


Figure 7.8: Euler-forward/central difference versus exact solution at different times for D = 0.4.

agrees quite well with the exact solution for the three time instances presented. Because of the use of a larger D, however, fewer iterations are necessary than in the Euler-forward case.

Parameter	Value
Method	Euler-Backward Central Difference
Interval	[0, 1]
N	21
Δt	0.0025
D	1.0
Initial Condition	$u(x,0) = \sin(\pi x)$
Boundary Conditions	u(0,t) = u(1,t) = 0

Table 7.3: Parameters used for the simulations presented in figure 7.9.

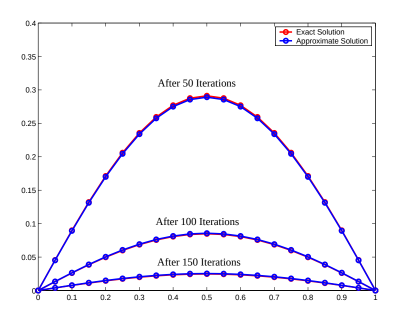


Figure 7.9: Euler-backward/central difference versus exact solution at varying times for D=1.0.

Multi-Dimensional Time-Space Stencils 7.1.5

We now turn to two-dimensional domains to demonstrate how we obtain discrete models of diffusion. Let us consider the unsteady diffusion equation in a domain Ω , see figure 7.10, with heat sources q(x,t), i.e.,

$$\frac{\partial \Theta}{\partial t} = \nabla^2 \Theta + q$$

$$\Theta(x, y, t) = 0 \text{ on } \partial \Omega$$
(7.8)

$$\Theta(x, y, t) = 0 \text{ on } \partial\Omega \tag{7.9}$$

$$\Theta(x, y, 0) = \Theta_0(x, y) \text{ in } \Omega. \tag{7.10}$$

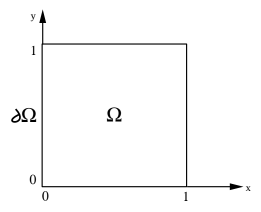


Figure 7.10: Two-dimensional domain.

We have assumed that we have homogeneous boundary conditions and that $\Theta_0(x, y)$ is a known function. We employ the θ -method to discretize this *in time* first, i.e.,

$$\frac{\Theta^{n+1} - \Theta^n}{\Delta t} = \theta(\nabla^2 \Theta^{n+1} + q^{n+1}) + (1 - \theta)(\nabla^2 \Theta^n + q^n).$$

We now use the Cartesian five-point stencil (see section 5.1) to discretize in space

$$\frac{\Theta_{ij}^{n+1} - \Theta_{ij}^{n}}{\Delta t} = \theta \left[\frac{\Theta_{i+1,j}^{n+1} - 2\Theta_{ij}^{n+1} + \Theta_{i-1,j}^{n+1}}{\Delta x^{2}} + \frac{\Theta_{i,j+1}^{n+1} - 2\Theta_{ij}^{n+1} + \Theta_{i,j-1}^{n+1}}{\Delta y^{2}} \right] + (1 - \theta) \left[\frac{\Theta_{i+1,j}^{n} - 2\Theta_{ij}^{n} + \Theta_{i-1,j}^{n}}{\Delta x^{2}} + \frac{\Theta_{i,j+1}^{n} - 2\Theta_{ij}^{n} + \Theta_{i,j-1}^{n}}{\Delta y^{2}} \right] + \theta q_{ij}^{n+1} + (1 - \theta) q_{ij}^{n}.$$

The accuracy of this discretization can be obtained by examining the truncation error, and as expected is

- For $\theta = 0$ or $\theta = 1$, the overall accuracy is $\mathcal{O}(\Delta t, \Delta x^2, \Delta y^2)$.
- For $\theta = 1/2$, the overall accuracy is $\mathcal{O}(\Delta t^2, \Delta x^2, \Delta y^2)$.

Directional Splitting Method

We consider again the diffusion equation (7.1) with $\kappa = 1$. One of the most economical approaches in solving this problem is to split directions and perform one-dimensional solves on corresponding and equivalent PDEs. The key to successful implementation of this idea is the construction of the new approximate equations. In the following, we demonstrate a few different ways of accomplishing this task.

1. Alternating-Direction-Implicit (ADI) Method

In two-dimensions, the Alternating-Direction-Implicit (ADI) method solves implicitly along the x-direction first, and subsequently it solves implicitly along the y-direction. This is done in two steps as follows

Pass I:

$$\frac{\Theta_{i,j}^* - \Theta_{i,j}^n}{\Delta t} = \Theta_{xx}^* + \Theta_{yy}^n$$

Pass II:

$$\frac{\Theta_{i,j}^{n+1} - \Theta_{i,j}^*}{\Delta t} = \Theta_{xx}^* + \Theta_{yy}^{n+1}.$$

From pass I we obtain Θ^* and from pass II we obtain the final solution Θ^{n+1} . To avoid bias towards one direction, we can alternate the order in the x-y directions every step. This method is $\mathcal{O}(\Delta t^2, \Delta x^2, \Delta y^2)$, and it is unconditionally stable. However, in the presence of time-dependent boundary conditions the two substeps introduce boundary errors of different orders, i.e., $\mathcal{O}(\Delta t)$, $\mathcal{O}(\Delta t^2)$ to the solution.

2. Approximate Factorization

We consider the time-discrete equation

$$\frac{\Theta_{ij}^{n+1} - \Theta_{ij}^{n}}{\Delta t} = \left(\frac{\partial^{2}}{\partial x^{2}} + \frac{\partial^{2}}{\partial y^{2}}\right) \Theta_{ij}^{n+1}$$

which is re-written in Helmholtz operator form

$$\left\{1 - \Delta t \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}\right)\right\} \Theta_{ij}^{n+1} = \Theta_{ij}^n.$$

We then approximate this by the product of two one-dimensional operators:

$$\left(1 - \Delta t \frac{\partial^2}{\partial x^2}\right) \left(1 - \Delta t \frac{\partial^2}{\partial y^2}\right) \Theta_{ij}^{n+1} \approx \Theta_{ij}^n,$$

which can be solved in two passes

Pass I:

$$\left(1 - \Delta t \frac{\partial^2}{\partial y^2}\right) \Theta_{i,j}^* = \Theta_{i,j}^n.$$

Pass II:

$$\left(1 - \Delta t \frac{\partial^2}{\partial x^2}\right) \Theta_{ij}^{n+1} = \Theta_{ij}^*.$$

Multiplying the two one-dimensional Helmholtz operators we obtain

$$\left\{1 - \Delta t \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}\right) + \underbrace{\Delta t^2 \left(\frac{\partial^2}{\partial x^2} \frac{\partial^2}{\partial y^2}\right)}_{\text{extra term}}\right\} \Theta_{ij}^{n+1} = \Theta_{ij}^n.$$

Therefore, the factorization has introduced an extra term which is $\mathcal{O}(\Delta t^2)$ and is equal to the local error of the Euler-backwards discretization.

Example: ADI Factorization versus Crank-Nicolson

We consider discretizations over a time interval of $2\Delta t$. We first construct a Crank-Nicolson time-discretization

$$\Theta^{n+2} - \Theta^n = \frac{2\Delta t}{2} \left\{ \left(\Theta_{xx}^{n+2} + \Theta_{yy}^{n+2} \right) + \left(\Theta_{xx}^n + \Theta_{yy}^n \right) \right\}$$

$$\Rightarrow \left\{ 1 - \Delta t \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} \right) \right\} \Theta^{n+2} = \left\{ 1 + \Delta t \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} \right) \right\} \Theta^n$$

and an approximate factorization:

$$\left(1 - \Delta t \frac{\partial^2}{\partial x^2}\right) \Theta^{n+1} = \left(1 + \Delta t \frac{\partial^2}{\partial y^2}\right) \Theta^n
\left(1 - \Delta t \frac{\partial^2}{\partial y^2}\right) \Theta^{n+2} = \left(1 + \Delta t \frac{\partial^2}{\partial x^2}\right) \Theta^{n+1}$$

where we have assumed that all these linear operators are commutative. Therefore, for the factorization we obtain

$$\left(1 - \Delta t \frac{\partial^2}{\partial x^2}\right) \left(1 - \Delta t \frac{\partial^2}{\partial y^2}\right) \Theta^{n+2} = \left(1 + \Delta t \frac{\partial^2}{\partial x^2}\right) \left(1 + \Delta t \frac{\partial^2}{\partial y^2}\right) \Theta^n$$

and expanding further we obtain

$$\left\{1 - \Delta t \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}\right) + \Delta t^2 \left(\frac{\partial^2}{\partial x^2} \frac{\partial^2}{\partial y^2}\right)\right\} \Theta^{n+2}$$

$$= \left\{1 + \Delta t \left(\frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2}\right) + \Delta t^2 \left(\frac{\partial^2}{\partial x^2} \frac{\partial^2}{\partial y^2}\right)\right\} \Theta^n.$$

Comparing this ADI form and the Crank-Nicolson form we see that there is an extra term in the factorization, namely

$$A = \Delta t^2 \left(\frac{\partial^2}{\partial x^2} \frac{\partial^2}{\partial y^2} \right) \left(\Theta^{n+2} - \Theta^n \right) .$$

We can approximate

$$\frac{\partial \Theta^n}{\partial t} = \frac{\Theta^{n+2} - \Theta^n}{2\Delta t} + \mathcal{O}(\Delta t) \Rightarrow A \approx 2\Delta t^3 \left(\frac{\partial^2}{\partial x^2} \frac{\partial^2}{\partial y^2}\right) \frac{\partial \Theta^n}{\partial t},$$

which is an $\mathcal{O}(\Delta t^3)$ term and thus consistent with the local error in the Crank-Nicolson approximation. Therefore, the accuracy between the two approaches is comparable, but in the approximate factorization only one-dimensional solves are involved.

3. ADI for Stationary Problems

ADI methods can also be used to solve the system resulting from a stationary elliptic equation, i.e.,

$$\Theta_{xx} + \Theta_{yy} = q.$$

Assuming that we discretize with the three-point central difference in each direction we obtain

$$[\mathbf{A}_x + \mathbf{A}_y]\mathbf{\Theta} = \mathbf{q} \tag{7.11}$$

where \mathbf{A}_x , \mathbf{A}_y are the discrete operators in x- and y-directions, respectively. In the following, we present algorithms for two-dimensional and three-dimensional discretizations.

Peaceman-Rachford Algorithms for Two-Dimensions

The idea here is to combine ADI with convergence acceleration tricks, i.e.,

$$(\mathbf{A}_x + \rho_n \mathbf{I})\Theta^* = (\rho_n \mathbf{I} - \mathbf{A}_y)\Theta^n + \mathbf{q}$$
$$(\mathbf{A}_y + \rho_n \mathbf{I})\Theta^{n+1} = (\rho_n \mathbf{I} - \mathbf{A}_x)\Theta^* + \mathbf{q},$$

where ρ_n , n = 1, ... is a sequence of positive acceleration parameters. Also, **I** is the identity operator.

For positive-definite systems, \mathbf{A}_x and \mathbf{A}_y have real eigenvalues and several theoretical results hold, see [89]. For example, if \mathbf{A}_x , \mathbf{A}_y are positive-definite and symmetric, any sequence $\rho_n = C > 0$ will produce a convergent iteration. The rate of convergence of ADI can be greatly enhanced by a sequence ρ_n in *cyclic* order.

Douglas-Rachford Algorithm for Three-Dimensions

Here we consider the three-dimensional analog of equation (7.11) and split in three directions as follows

$$(\mathbf{A}_x + \rho_n \mathbf{I})\Theta^* = -(\mathbf{A}_x + 2\mathbf{A}_y + 2\mathbf{A}_z - \rho_n \mathbf{I})\Theta^n + 2\mathbf{q}$$

$$(\mathbf{A}_y + \rho_n \mathbf{I})\Theta^{**} = -(\mathbf{A}_x + \mathbf{A}_y + 2\mathbf{A}_z - \rho_n \mathbf{I})\Theta^n - \mathbf{A}_x\Theta^* + 2\mathbf{q}$$

$$(\mathbf{A}_z + \rho_n \mathbf{I})\Theta^{***} = -(\mathbf{A}_x + \mathbf{A}_y + \mathbf{A}_z - \rho_n \mathbf{I})\Theta^n - \mathbf{A}_x\Theta^* - \mathbf{A}_y\Theta^{**} + 2\mathbf{q}$$

We note here that convergence is not always guaranteed for $\rho_n = C > 0$.

Stability Analysis

We can apply von Neumann analysis similarly to the one-dimensional case, i.e.,

$$\Theta_{ij}^n = \sum_k \sum_{\ell} a_{k\ell}^n \sin k\pi x_i \sin \ell \pi y_j.$$

We then substitute this expression into the *homogeneous* difference equation to obtain the uncoupled recursion for $a_{k\ell}^n$:

$$a_{k\ell}^{n+1} = a_{k\ell}^n \frac{1 - D(1 - \theta)\sigma_{k\ell}}{1 + D\theta\sigma_{k\ell}}$$

$$\sigma_{k\ell} = 2[(1 - \cos k\pi\Delta x) + \delta^2(1 - \cos \ell\pi\Delta y)] > 0$$

where $\delta = \frac{\Delta x}{\Delta y}$ and $D = \frac{\Delta t}{\Delta x^2}$ is the diffusion number. For the specific members of the θ -family we have:

- For $\theta = 1/2, 1$ unconditional stability.
- For $\theta = 0$ explicit time-discretization, thus

$$-1 \le 1 - D\sigma_{k\ell} \le 1 \Rightarrow D \le \frac{2}{\sigma_{k\ell}}$$

where

$$\max_{k,\ell} \sigma_{k\ell} = 4(1+\delta^2).$$

Therefore, $D \leq \frac{1}{2(1+\delta^2)}$ and thus

$$\Delta t \le \frac{1}{2} \frac{1}{2 \left(\frac{1}{\Delta x^2} + \frac{1}{\Delta y^2} \right)}.$$

If $\Delta x = \Delta y$ then

$$\Delta t \le \frac{\Delta x^2}{4} \, .$$

This condition is more restrictive than in the one-dimensional case where $\Delta t \leq \frac{\Delta x^2}{2}$!

Remark 1: In three-dimensions the diffusion number limit is 1/8 (obtained by similar analysis). Therefore, if we double the resolution in one-dimensional problems the cost is increased *eight times* whereas if we double the resolution in three-dimensional problems the cost is increased 32 times!

Remark 2: Implicit time-discretization requires the solution of a stationary elliptic equation of the Helmholtz type, since

$$\frac{\Theta^{n+1} - \Theta^n}{\Delta t} = \nabla^2 \Theta^{n+1} + q^{n+1}$$

and by re-arranging we obtain

$$\left(\nabla^2 - \frac{1}{\Delta t}\right)\Theta^{n+1} = -\frac{\Theta^n}{\Delta t} - q^{n+1},$$

which is an A-stable scheme.

We can also apply **matrix methods** as in the one-dimensional case to analyze the stability of multi-dimensional difference equations. For example, for two-dimensional problems with *periodic boundary* conditions we consider a single harmonic component $e^{i\phi_x}e^{i\phi_y}$, then discretizing with the θ -family we obtain

$$G - 1 = \theta DG[(e^{i\phi_x} + e^{-i\phi_x} - 2) + \delta^2(e^{i\phi_y} + e^{-i\phi_y} - 2)] + (1 - \theta)D[(e^{i\phi_x} + e^{-i\phi_x} - 2) + \delta^2(e^{i\phi_y} + e^{-i\phi_y} - 2)],$$

and therefore

$$G = \frac{1 - 4D(1 - \theta)[\sin^2\frac{\phi_x}{2} + \delta^2\sin^2\frac{\phi_y}{2}]}{1 + 4D\theta[\sin^2\frac{\phi_x}{2} + \delta^2\sin^2\frac{\phi_y}{2}]},$$

where $\delta \equiv \frac{\Delta x}{\Delta y}$. For the specific members of the family:

• For $\theta = 0$, we obtain

$$|1 - 4D(\sin^2\frac{\phi_x}{2} + \delta^2\sin^2\frac{\phi_y}{2})| \le 1,$$

which for $\Delta x = \Delta y$ leads to: $D \leq \frac{1}{4}$ or $\Delta t \leq \frac{\Delta x^2}{4\kappa}$, as before.

• For $\theta=1,1/2$ we have unconditional stability $\forall\, D>0.$

7.2 Iterative Solvers

Discretization of the diffusion equation leads us naturally to iterative solutions of elliptic problems of the form

$$\nabla^2 \Theta + q = 0 \text{ on } \Omega, \quad \Theta(x, y) = 0 \text{ on } \partial\Omega$$
 (7.12)

where Ω and $\partial\Omega$ are the domain and the boundary of the domain, respectively (see figure 7.11). Here we assume homogeneous boundary conditions for clarity in the presentation.

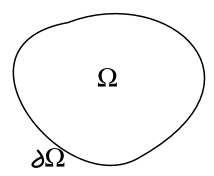


Figure 7.11: General two-dimensional domain Ω .

This connection becomes more clear by introducing a pseudo-time derivative to obtain

$$\frac{\partial \Theta}{\partial t} = \nabla^2 \Theta + q \text{ on } \Omega, \quad \Theta(x, y) = 0 \text{ on } \partial \Omega.$$
 (7.13)

We then compute the sequence

$$\Theta_{ij}^0, \, \Theta_{ij}^1, \, \Theta_{ij}^2, \, \dots, \, \Theta_{ij}^{n-1}, \, \Theta_{ij}^n \, \dots, \, \Theta_{ij}^{\infty}$$

from equation (7.13) in the limit of many iterations. Therefore, Θ_{ij}^{∞} is the steady-state solution, which is different than the exact solution due to *spatial* discretization error. Any other member of the sequence differs from the exact solution by the spatial discretization error as well as the solver convergence error; we will analyze the latter in the following.

7.2.1 Jacobi Algorithm

We now demonstrate how we can derive the algorithm for perhaps the oldest iterative solver, the Jacobi iterative method. We apply Euler-forward discretization to the diffusion equation (7.13) assuming that we have a two-dimensional domain Ω (such as in figure 7.12 (left)) and taking $\Delta x = \Delta y$. We also assume that we are given some function q defined on the domain (such as in figure 7.12 (right)). We then have

$$\frac{\Theta_{ij}^{n+1} - \Theta_{ij}^n}{\Delta t} = q_{ij} + \frac{1}{\Delta x^2} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n - 4\Theta_{ij}^n + \Theta_{i,j-1}^n + \Theta_{i-1,j}^n)$$

or:

$$\Theta_{ij}^{n+1} = \Delta t \cdot q_{ij} + D(\Theta_{i+1,j}^n + \Theta_{i,j+1}^n - 4\Theta_{ij}^n + \Theta_{i-1,j}^n + \Theta_{i,j-1}^n) + \Theta_{ij}^n,$$

where $D = \frac{\Delta t}{\Delta x^2}$ is the diffusion number defined in equation (7.5). In order to determine the rate of convergence, we write

$$\Theta_{ij}^n = \Theta_{ij}^\infty + \epsilon_{ij}^n$$

where ϵ_{ij}^n is the deviation from the fixed point, i.e., the steady-state. It is also clear that Θ_{ij}^{∞} satisfies the difference equation

$$\frac{1}{\Delta x^2} (\Theta_{i+1,j}^{\infty} + \Theta_{i,j+1}^{\infty} - 4\Theta_{ij}^{\infty} + \Theta_{i-1,j}^{\infty} + \Theta_{i,j-1}^{\infty}) + q_{ij} = 0.$$

By subtracting the two difference formulas above we obtain an equation for the convergence error

$$\epsilon_{ij}^{n+1} = \epsilon_{ij}^n + D(\epsilon_{i+1,j}^n + \epsilon_{i,j+1}^n - 4\epsilon_{ij}^n + \epsilon_{i-1,j}^n + \epsilon_{i,j-1}^n).$$

Convergence is obtained if $|\epsilon^n| \to 0$, which is implied by absolute stability. Therefore, we require that $D \leq 1/4$ and specifically we take D = 1/4 for the fastest possible convergence.

Given the above analysis we can now write the **Jacobi algorithm** for solving equation (7.12) in the form

$$\Theta_{ij}^{n+1} = \frac{1}{4} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n + \Theta_{i-1,j}^n + \Theta_{i,j-1}^n + \Delta x^2 q_{ij})$$

for all the *interior points*. Appropriate modifications with one-sided differences may be required to construct similar equations applied at the boundary.

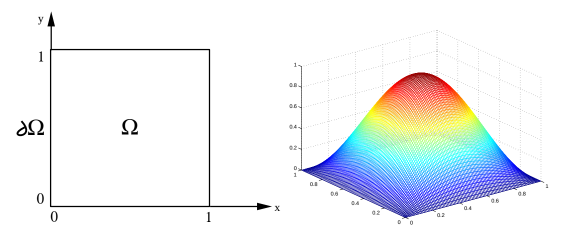
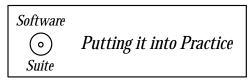


Figure 7.12: Example domain (left) and function q(x,y) (right) used to demonstrate the Jacobi algorithm.



We now present a C++ implementation of the algorithm previously discussed. This function takes as input the number of points N, the grid spacing dx, the δt , the matrix **A** as a double ** array, the right-hand-side vector **q**, and a stopping tolerance.

```
int Diffusion_Jacobi(int N, double dx, double dt,
      double **A, double **q, double abstol){
  int i,j,k;
  int maxit = 100000;
  double sum;
  double ** Aold = CreateMatrix(N,N);
  double D = dt/(dx*dx);
  for(i=1; i<N-1; i++)
    for(j=1;j<N-1;j++)
      Aold[i][j] = 1.0;
  /* Boundary Conditions -- all zeros */
  for(i=0;i<N;i++){
   A[0][i] = 0.0;
   A[N-1][i] = 0.0;
   A[i][0] = 0.0;
   A[i][N-1] = 0.0;
  }
  for(k=0; k < maxit; k++){
   for(i = 1; i < N-1; i++){
      for(j=1; j<N-1; j++){
         A[i][j] = dt*q[i][j] + Aold[i][j] +
           D*(Aold[i+1][j] + Aold[i][j+1] - 4.0*Aold[i][j] +
                   Aold[i-1][j] + Aold[i][j-1]);
      }
    }
    sum = 0.0;
    for(i=0;i<N;i++){
      for(j=0;j<N;j++){
        sum += (Aold[i][j]-A[i][j])*(Aold[i][j]-A[i][j]);
        Aold[i][j] = A[i][j];
      }
    }
    if(sqrt(sum)<abstol){</pre>
      DestroyMatrix(Aold,N,N);
      return k;
```

```
}

cerr << "Jacobi: Maximum Number of Interations Reached \n";
DestroyMatrix(Aold,N,N);

return maxit;
}</pre>
```

Remark: Remember to free dynamically allocated memory when exiting a routine. Often programmers forget to free things when they exit a function after a particular tolerance is found. Observe in the code above that if the tolerance is reached, we first free the dynamically allocated memory, and then return the value k.

Programming Note

• You must often decide which resource is more valuable - memory or computation. In this example, we expend extra memory to store boundary conditions, which are zero, so that the computational algorithm is simple. If we had to save memory, then we would need to explicitly write code, which takes into account the modification of the differencing stencil when we are computing along the edges and corners of the domain.

Convergence Rate

The convergence error ϵ^n satisfies the same equation as the $a_{k\ell}^n$ modes in the von Neumann equation (7.7) of the previous section, i.e.,

$$\epsilon_{k\ell}^{n+1} = \epsilon_{k\ell}^n \left\{ \frac{1 - \frac{1}{4}(1 - \theta)\sigma_{k\ell}}{1 + \frac{\theta}{4}\sigma_{k\ell}} \right\}$$

where we set $\theta = 0$ for Euler-forward discretization and $\delta = 1$ since $\Delta x = \Delta y$. Therefore

$$\epsilon_{k\ell}^{n+1} = \epsilon_{k\ell}^n (1 - \frac{1}{4}\sigma_{k\ell})$$

where

$$\sigma_{k\ell} = 2[(1 - \cos k\pi \Delta x) + (1 - \cos \pi \ell \Delta y)].$$

The minimum of $\sigma_{k\ell}$ corresponds to the most "resistant" error in decaying. This occurs for $(k = \ell = 1)$, and thus we obtain for an error estimate

$$\epsilon^{n+1} = \epsilon^n \left[1 - \frac{1}{2} \left[\left(1 - \cos \pi \Delta x \right) + \left(1 - \cos \pi \Delta x \right) \right] \right]$$

or

$$\epsilon^{n+1} = \epsilon^n [1 - (1 - \cos \pi \Delta x)],$$

which can be approximated as

$$\epsilon^{n+1} \approx \epsilon^n \left(1 - \frac{\pi^2 \Delta x^2}{2} \right).$$

We now define the convergence rate z_J of the Jacobi method

$$z_J \equiv \frac{\epsilon^{n+1}}{\epsilon^n} = \left(1 - \frac{\pi^2 \Delta x^2}{2}\right),$$

therefore

$$\epsilon^n = z_J^n \epsilon^0$$

where n on z_J^n is an exponent, whereas elsewhere denotes iteration level.

In order to reduce e^n to e^{-d} , that is to obtain accuracy defined by $z_J^n = e^{-d}$, we have

$$n \ln z_J = -d \Rightarrow -n \frac{\pi^2 \Delta x^2}{2} \sim -d \Rightarrow n \sim \frac{2}{\pi^2} \frac{d}{\Delta x^2}$$
 iterations.

Here $\Delta x = \frac{1}{N}$, and d is roughly equal to the number of correct digits in the solution. Therefore, the number of iterations is proportional to the total number of grid points, in a two-dimensional grid $N \times N$, i.e.,

$$n \sim \mathcal{O}(N^2)$$
.

The corresponding computational work is

$$z_J * \mathcal{O}(N^2) \approx \mathcal{O}(N^4)$$

since we need at least one operation for each grid point. This corresponds to more work than $\mathcal{O}(N^3)$ required for a backsolve in Gaussian elimination, although to factorize the matrix by LU requires $\mathcal{O}(N^4)$ (see section 9.1). These estimates assume that we solve the elliptic problem (7.12) on a grid $N \times N$ with center-differencing resulting in a matrix with order N^2 and bandwidth m = N. The number of backsolve operations in the Gaussian elimination is then $\mathcal{O}(mN^2) = \mathcal{O}(N^3)$.

In the above analysis, we considered specifically the *discrete diffusion equation* to derive the Jacobi iteration algorithm. However, this is not necessary, and the Jacobi iteration can be employed to solve general linear systems of the form

$$\mathbf{A}\mathbf{x} = \mathbf{b}$$
.

To this end, we decompose the matrix **A** as

$$A = \underbrace{L}_{\text{(strictly lower)}} + \underbrace{D}_{\text{(strictly upper)}} + \underbrace{U}_{\text{(strictly upper)}}$$

Denoting by n the iteration number, we formulate the following iteration procedure

$$\mathbf{D}\mathbf{x}^{n+1} = \mathbf{b} - (\mathbf{L} + \mathbf{U})\mathbf{x}^n.$$

This is the general Jacobi algorithm which does not requires matrix inversion as the diagonal matrix is trivially inverted, that is

$$\mathbf{D}^{-1} = \frac{1}{\mathbf{D}},$$

assuming non-zero diagonal entries.

As an example, we can recover the Jacobi algorithm specific to diffusion equation from:

$$\frac{-4}{\Delta x^2} \Theta_{ij}^{n+1} = -q_{ij} - \frac{1}{\Delta x^2} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n + \Theta_{i-1,j}^n + \Theta_{i,j-1}^n)$$

or

$$\Theta_{ij}^{n+1} = \frac{1}{4} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n + \Theta_{i-1,j}^n + \Theta_{i,j-1}^n + \Delta x^2 q_{ij})$$

as before.

The general Jacobi iteration converges if matrix **A** is strictly row-diagonally dominant. This means that the absolute value of each diagonal element of **A** is greater than the sum of absolute values of all the other entries in that row. However, this condition does not apply to the discrete diffusion equation considered here. Instead, a weaker condition is applied, by requiring that the matrix **A** is weakly row-diagonally dominant. This simply implies that some of the diagonal elements may be equal to the sum in each row, but greater on at least one row. This condition needed to be strengthened with the condition of *irreducibility*.

Th irreducibility condition is related to the *directed graph* of the matrix, which consists of a sequence of nodes with arrows connecting the non-zeros entries of a matrix. If each node in the graph is accessible by any other node, then this is a strongly-connected graph and the corresponding matrix is irreducible. Together the two conditions of weak row-diagonal dominance *and* of irreducibility guarantee that the Jacobi method will converge for systems resulting from central difference discretizations of the diffusion equation. Note that the cross-directional splitting of section 7.1.5 leads to a matrix with a corresponding graph which is not strongly connected.

Programming Note

• What tolerance should you use? This depends on the problem you are trying to solve. If the modeling error for your problem is $O(10^{-4})$, there may not be a need to converge your solution to $O(10^{-14})$. The bottom line know and understand the problem that you are trying to solve!



In the *software suite* we implement the Jacobi algorithm again but this time taken into account the pre-specified tolerance.

7.2.2 Parallel Jacobi Algorithm

In our implementation of parallel Jacobi, we will use an MPI command not previously discussed: $MPI_Allgather$. MPI_Gather and $MPI_Allgather$ provide us the ability to assemble (gather) data from a collection of processes and combine it either on one specific process (MPI_Gather) or on all processes ($MPI_Allgather$). MPI also provides an inverse operation called $MPI_Scatter$, which distributes data from one process to a collection of processes. We will now present for these three functions the function call syntax, argument list explanation, usage example and some remarks.

MPI_Gather:

Function Call Syntax

```
int MPI_Gather(
         void*
                         sendbuf
                                          in
         int
                         sendcount
                                         in
         MPI_Datatype
                         sendtype
                                          in
         void*
                         recvbuf
                                          out
         int
                         recvcount
                                         in
         MPI_Datatype
                         recytype
                                          in
                                          in
         int
                         root
         MPI_Comm
                                         in
                         comm
```

Understanding the Argument List

- sendbuf starting address of the send buffer.
- sendcount number of elements in the send buffer.
- sendtype data type of the elements in the send buffer.
- recvbuf starting address of the receive buffer.
- recvcount number of elements for any single receive.
- recvtype data type of the elements in the receive buffer.
- root rank of the root process obtaining the result.
- *comm* communicator.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be gathered
                    from each process
               // process to which the data is gathered
int root;
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
// Determine datasize and root
double * senddata = new double[datasize];
double * recvdata = NULL;
if(mynode == root)
  recvdata = new double[datasize*totalnodes];
// Fill in senddata array for each process
MPI_Gather(senddata,datasize,MPI_DOUBLE,recvdata,datasize,
            MPI_DOUBLE,root,MPI_COMM_WORLD);
// At this point, process root has received from each
// process the contents of senddata and stored them
// in recvdata according to the process ordering
```

Remarks

- MPI_Gather allows us to "gather" to one specified process information from all processes with the communicator. This collection function allows us to retrieve from all processes information that may need to be acted upon in an assembled form. In figure 7.13 we present a schematic demonstrating the results of a gather operation from all processes to process zero.
- The array *recvbuf* is only relevant on the process of rank root. All other processes may pass NULL in place of the argument for *recvbuf*.
- In most cases the *sendtype* and *recvtype* are identical, and the value of *sendcount* and the value of *recvcount* are identical. MPI requires that the amount of data sent (sendcount times the size in bytes of the datatype sendtype) equals the amount of data received (recvcount times the size in bytes of the datatype recvtype) per process/root pair.
- The allocated size of *recvbuf* should be equal to at least the value of *recvtype* times the number of processes (*totalnodes*).

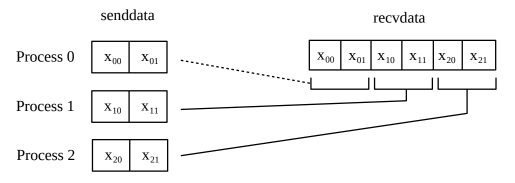


Figure 7.13: *MPI_Gather* schematic demonstrating a gather to process zero of two data objects from each process.

MPI_Allgather:

Function Call Syntax

```
int MPI_Allgather(
        void*
                         sendbuf
                                         in
        int
                         sendcount
                                         in
        MPI_Datatype
                         sendtype
                                         in
        void*
                         recvbuf
                                         out
        int
                         recvcount
                                         in
        MPI_Datatype
                         recvtype
                                         in
        MPI_Comm
                         comm
                                         in
```

Understanding the Argument List

- sendbuf starting address of the send buffer.
- sendcount number of elements in the send buffer.
- sendtype data type of the elements in the send buffer.
- recvbuf starting address of the receive buffer.
- recvcount number of elements for any single receive.
- recvtype data type of the elements in the receive buffer.
- *comm* communicator.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be
```

Remarks

- MPI_Allgather allows us to "gather" to all processes information from all processes with the communicator. The action of "all gather" is as if we were to gather to one process using MPI_Gather, and then send from that process to all other processes the assembled information. In figure 7.14 we present a schematic demonstrating the results of an "all gather" operation.
- Both the *sendbuf* and *recvbuf* arrays are relevant on all processes in the communicator.
- In most cases the *sendtype* and *recvtype* are identical and the value of *sendcount*, and the value of *recvcount* are identical. MPI requires that the amount of data sent (sendcount times the size in bytes of the datatype sendtype) equals the amount of data received (recvcount times the size in bytes of the datatype recvtype) per process/root pair.
- The allocated size of *recvbuf* should be at least equal to the value of *recvtype* times the number of processes (*totalnodes*).

MPI_Scatter:

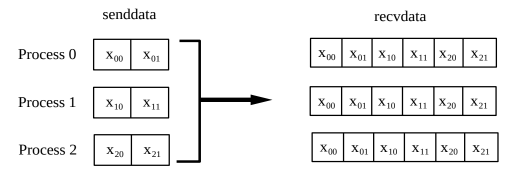


Figure 7.14: MPI_Allgather schematic demonstrating an allgather to all processes of two data objects from each process.

```
int MPI_Scatter(
         void*
                         sendbuf
                         sendcount
                                         in
         int
         MPI_Datatype
                         sendtype
                                         in
         void*
                         recvbuf
                                         out
         int
                         recvcount
                                         in
         MPI_Datatype
                         recvtype
                                         in
         int
                         root
                                         in
         MPI_Comm
                         comm
                                         in
```

Understanding the Argument List

- sendbuf starting address of the send buffer.
- sendcount number of elements in the send buffer.
- sendtype data type of the elements in the send buffer.
- recvbuf starting address of the receive buffer.
- recvcount number of elements for any single receive.
- recvtype data type of the elements in the receive buffer.
- root rank of the root process obtaining the result.
- comm communicator.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be scattered to each process
int root; // process from which the data is scattered

MPI_Init(&argc,&argv);
```

Remarks

- *MPI_Scatter* allows us to "scatter" from one specified process information to all processes with the communicator. This function allows us to distribute partitioned data to all processes. In figure 7.15 we present a schematic demonstrating the results of a scatter operation from process zero to all processes within the communicator.
- The array *sendbuf* is only relevant on the process of rank root. All other processes may pass NULL in place of the argument for *sendbuf*.
- In most cases the *sendtype* and *recvtype* are identical, and the value of *sendcount* and the value of *recvcount* are identical. MPI requires that the amount of data sent (sendcount times the size in bytes of the datatype sendtype) equals the amount of data received (recvcount times the size in bytes of the datatype recvtype) per process/root pair.
- The allocated size of *sendbuf* should be at least equal to the value of *sendtype* times the number of processes (*totalnodes*).
- Each recvdata array should contain at least datasize elements.

We now present a parallel implementation of the Jacobi method. This function assumes that the MPI initialization has already been accomplished by the calling function. It takes

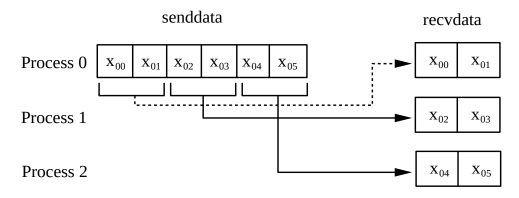
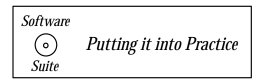


Figure 7.15: MPI_Scatter schematic demonstrating a scatter from process zero of two data objects to each process.



as input its process id number, the total number of processes being used, the size of the matrix, the matrix \mathbf{A} as a double** array, an array x into which the solution is to be stored, an array b containing the right-hand-side, and a stopping tolerance. On all processes other than the root process, the pointers A, x, and b should be NULL; process zero distributes the matrix and right-hand-side among the processes, and only process zero returns the solution vector \mathbf{x} . We first present the function definition and then present some remarks on the code.

```
int Jacobi_P(int mynode, int numnodes, int N, double **A,
             double *x, double *b, double abstol){
  int i,j,k,i_global;
  int maxit = 100000;
  int rows_local,local_offset,last_rows_local;
  int *count,*displacements;
  double sum1, sum2, *xold;
  double error_sum_local, error_sum_global;
  MPI_Status status;
  rows_local = (int) floor(N/numnodes);
  local_offset = mynode*rows_local;
  if(mynode == (numnodes-1))
    rows_local = N - rows_local*(numnodes-1);
  /*Distribute the Matrix and R.H.S. among the processors */
  if(mynode == 0){
    for(i=1;i<numnodes-1;i++){</pre>
      for(j=0;j<rows_local;j++)</pre>
```

```
MPI_Send(A[i*rows_local+j],N,MPI_DOUBLE,i,j,
               MPI_COMM_WORLD);
    MPI_Send(b+i*rows_local,rows_local,MPI_DOUBLE,i,
             rows_local, MPI_COMM_WORLD);
  }
  last_rows_local = N-rows_local*(numnodes-1);
  for(j=0;j<last_rows_local;j++)</pre>
    MPI_Send(A[(numnodes-1)*rows_local+j],N,MPI_DOUBLE,
             numnodes-1,j,MPI_COMM_WORLD);
  MPI_Send(b+(numnodes-1)*rows_local,last_rows_local,
           MPI_DOUBLE,numnodes-1,last_rows_local,
           MPI_COMM_WORLD);
}
else{
  A = CreateMatrix(rows_local, N);
  x = new double[rows_local];
  b = new double[rows_local];
  for(i=0;i<rows_local;i++)</pre>
    MPI_Recv(A[i],N,MPI_DOUBLE,O,i,MPI_COMM_WORLD,&status);
  MPI_Recv(b,rows_local,MPI_DOUBLE,0,rows_local,
           MPI_COMM_WORLD,&status);
}
xold = new double[N];
count = new int[numnodes];
displacements = new int[numnodes];
//set initial guess to all 1.0
for(i=0; i<N; i++){
  xold[i] = 1.0;
}
for(i=0;i<numnodes;i++){</pre>
  count[i] = (int) floor(N/numnodes);
  displacements[i] = i*count[i];
count[numnodes-1] = N - ((int)floor(N/numnodes))*
                     (numnodes-1);
for(k=0; k<\max it; k++){
  error_sum_local = 0.0;
  for(i = 0; i<rows_local; i++){</pre>
    i_global = local_offset+i;
    sum1 = 0.0; sum2 = 0.0;
```

```
for(j=0; j < i_global; j++)
      sum1 = sum1 + A[i][j]*xold[j];
    for(j=i_global+1; j < N; j++)
      sum2 = sum2 + A[i][j]*xold[j];
    x[i] = (-sum1 - sum2 + b[i])/A[i][i_global];
    error_sum_local += (x[i]-xold[i_global])*
                        (x[i]-xold[i_global]);
  }
  MPI_Allreduce(&error_sum_local,&error_sum_global,1,
                 MPI_DOUBLE,MPI_SUM,MPI_COMM_WORLD);
  MPI_Allgatherv(x,rows_local,MPI_DOUBLE,xold,count,
                  displacements,MPI_DOUBLE,MPI_COMM_WORLD);
  if(sqrt(error_sum_global) < abstol) {</pre>
    if(mynode == 0){
      for(i=0;i<N;i++)</pre>
        x[i] = xold[i];
    }
    else{
      DestroyMatrix(A,rows_local,N);
      delete[] x;
      delete[] b;
    }
    delete[] xold;
    delete[] count;
    delete[] displacements;
    return k;
 }
}
cerr << "Jacobi: Maximum Number of Interations Reached\n";</pre>
if(mynode == 0){
  for(i=0;i<N;i++)</pre>
    x[i] = xold[i];
}
else{
  DestroyMatrix(A,rows_local,N);
  delete[] x;
  delete[] b;
}
delete[] xold;
delete[] count;
delete[] displacements;
```

```
return maxit;
}
```

Remark 1: The rows are distributed across processes as shown in figure 7.16. Notice that in this routine we do not require that the number of processes equally divide the number of rows. This fact is taken into account using the *variable length* feature in MPI as discussed in the next remark.

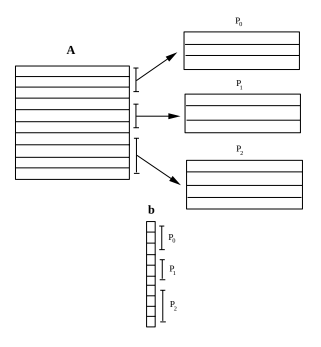


Figure 7.16: Distribution of matrix A across processes.

Remark 2: Observe in the code above that we use a slight modification to $MPI_Allgather$, $MPI_Allgatherv$. The function $MPI_Allgatherv$ allows us to specify per process how much information is expected. Recall that in the standard $MPI_Allgather$ it is assumed that each process is sending the same amount of data.

Remark 3: As you can see, other than the initial distribution of the matrix **A** and the right-hand-side vector **b**, the Jacobi method is highly parallelizable. Only two MPI calls are needed per iteration - MPI_Allreduce to compute the error and MPI_Allgatherv to distribute the updated solution vector at the conclusion of an iteration.

7.2.3 Gauss-Seidel Algorithm

This algorithm is usually twice as fast as the Jacobi algorithm but not as easily parallelizable. It is based on the observation that convergence usually is improved if we compute implicitly, hence in the Gauss-Seidel algorithm we use the most recently computed values. To illustrate

this algorithm, we use the matrix version of the Jacobi algorithm, where instead of the diagonal on the left-hand-side we employ the lower triangular matrix, i.e.,

$$(\mathbf{L} + \mathbf{D})\mathbf{x}^{n+1} = \mathbf{b} - \mathbf{U}\mathbf{x}^n.$$

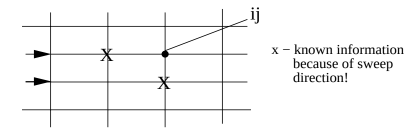


Figure 7.17: Horizontal sweeping of the grid from left to right.

We now apply this algorithm to the diffusion equation

$$\frac{1}{\Delta x^2} (\Theta_{i-1,j}^{n+1} + \Theta_{i,j-1}^{n+1} - 4\Theta_{ij}^{n+1}) = -q_{ij} - \frac{1}{\Delta x^2} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n)$$

or

$$\Theta_{ij}^{n+1} = \frac{1}{4} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n + \Theta_{i-1,j}^{n+1} + \Theta_{i,j-1}^{n+1} + \Delta x^2 q_{ij}).$$

We note that although on the right-hand-side we use values at the (n + 1) time level, these values are already known because of the "sweep" process we choose, from left to right in the grid of figure 7.17. Therefore, we use the newly computed information sooner than in the Jacobi method where we wait for a full cycle!

The *serial* Gauss-Seidel method is better than the *serial* Jacobi iteration as the convergence criteria are similar but the convergence rate is twice as large, i.e.,

$$z_{GS} = z_J^2$$

in most cases (always when we have consistent ordering, see [95]).

We can also use overwriting $\Theta_{ij}^{n+1} \leftrightarrow \Theta_{ij}^n$ to save in memory. However, the overall work is still $\mathcal{O}(N^4)$ on a grid with $N \times N$ points, so we have simply reduced the number of iterations by half but not the scaling with respect to the number of grid points. On a parallel computer, however, we need to use a special ordering of the unknowns on the grid in order to obtain an efficient algorithm as the standard Gauss-Seidel algorithm is inherently serial.



In the *software suite* we present implementations of the Gauss-Seidel algorithm for the diffusion equation and as a general iterative solver for the system $\mathbf{A}\mathbf{x} = \mathbf{b}$.

When examining the functions $Diffusion_GaussSeidel$ and GaussSeidel, we draw your attention to the following:

• Observe that the function definitions for $Diffusion_GaussSeidel$ and GaussSeidel are very similar to the definitions for $Diffusion_Jacobi$ and Jacobi, respectively. Recall from our previous discussion that the only modification necessary to transform the Jacobi method into the Gauss-Seidel method is to modify the updating pattern, as seen in these coding examples.

• Observe that although in Gauss-Seidel we could directly update the solution array with the new values, we still maintain an "old" and "new" solution array. We do this so that at the end of one sweep we can compare the two solutions to determine when the solution has converged.

In figure 7.18, we present the number of iterations necessary to converge to the solution for difference mesh sizes using our Jacobi and Gauss-Seidel implementations for the diffusion equation example previously given. Observe that when solving for the same mesh size, the Gauss-Seidel method requires half the number of iterations to converge as the Jacobi method, just as the theory discussed above predicted.

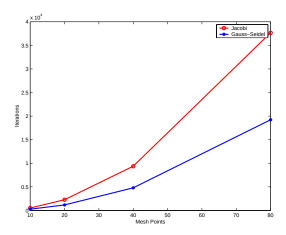


Figure 7.18: Comparison of the convergence rate for the Jacobi and Gauss-Seidel algorithms.

7.2.4 Parallel (Black-Red) Gauss-Seidel Algorithm

The computational complexity of the Gauss-Seidel iteration

$$\mathbf{x}^{n+1} = (\mathbf{L} + \mathbf{D})^{-1} [-\mathbf{U}\mathbf{x}^n + \mathbf{b}]$$

consists of

- Matrix-vector products for computing $\mathbf{U}\mathbf{x}^n$,
- Solution of the triangular system for $(\mathbf{L} + \mathbf{D})^{-1}$.

In particular, the solution of the triangular system can cause limitations with respect to parallelism, see chapter 9. However, we can apply a proper re-ordering of the difference equations in order to introduce more parallelism. This can be accomplished by a *multi-coloring scheme*, the simplest version of which is the *red-black* algorithm.

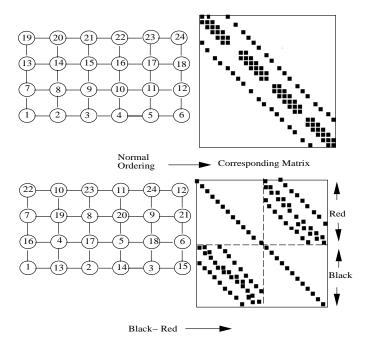


Figure 7.19: Ordering for standard Gauss-Seidel (top) and for black-red Gauss-Seidel (bottom).

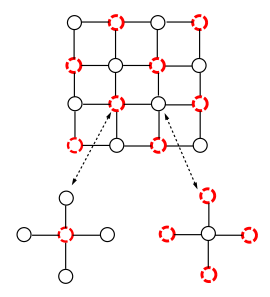


Figure 7.20: Typical black (solid line) and red (dashed line) stencils for the parallel Gauss-Seidel algorithm.

The key idea is to group the grid points into two groups, identified as black and red nodes, and observe that for Cartesian differencing the black nodes are surrounded by black nodes only, and the red nodes are surrounded by red nodes only. This is shown schematically in figures 7.19 and 7.20 along with the corresponding matrices **A**. Using this grouping and denoting with subscripts (b) and (r) the black and red nodes, respectively, we can decompose the system as follows

$$\left[egin{array}{ccc} \mathbf{D}_r & \mathbf{U} \ \mathbf{L} & \mathbf{D}_b \end{array}
ight] \left[egin{array}{c} \mathbf{x}_r \ \mathbf{x}_b \end{array}
ight] = \left[egin{array}{c} \mathbf{b}_r \ \mathbf{b}_b \end{array}
ight]$$

and using the Gauss-Seidel iteration, we obtain:

or

$$\begin{bmatrix} \mathbf{D}_r & 0 \\ \mathbf{L} & \mathbf{D}_b \end{bmatrix} \begin{pmatrix} \mathbf{x}_r^{n+1} \\ \mathbf{x}_b^{n+1} \end{pmatrix} = \begin{bmatrix} 0 & -\mathbf{U} \\ 0 & 0 \end{bmatrix} \begin{pmatrix} \mathbf{x}_r^n \\ \mathbf{x}_b^n \end{pmatrix} + \begin{pmatrix} \mathbf{b}_r \\ \mathbf{b}_b \end{pmatrix}$$
$$\begin{cases} \mathbf{x}_r^{n+1} = \mathbf{D}_r^{-1}[-\mathbf{U}\mathbf{x}_b^n + \mathbf{b}_r] \\ \mathbf{x}_b^{n+1} = \mathbf{D}_b^{-1}[-\mathbf{L}\mathbf{x}_r^{n+1} + \mathbf{b}_b] \end{cases}.$$

Thus, we do not have to invert a triangular system any more. Instead, we need to perform matrix-vector products and vector scaling operations.

An alternative way is to eliminate the red unknowns and solve for the black nodes first, i.e.,

$$(\mathbf{D}_b - \mathbf{U}\mathbf{D}_r^{-1}\mathbf{U})\mathbf{x}_b = \mathbf{b}_b + \mathbf{U}\mathbf{D}_r^{-1}\mathbf{b}_r,$$

with half as many unknowns. However, we need to solve a triangular system, which we can precondition with the diagonal for faster convergence, see chapter 9.

Remark 1: In general, the convergence rate depends on ordering unless it is a *consistent* ordering, see [95]. There have also been efforts to assign chaotic ordering (also known as *chaotic iteration*) but such schemes are not guaranteed that they will converge! A matrix $\mathbf{A} = \mathbf{L} + \mathbf{D} + \mathbf{U}$ is *consistently-ordered* if the eigenvalues of the matrix

$$\mathbf{D}^{-1}(\alpha^{-1}\mathbf{L} + \alpha\mathbf{U})$$

do not depend on α for any $\alpha \neq 0$. A different definition was introduced by Young based on the permutations of the matrix **A** [95]. Specifically, the matrix is termed 2-cyclic if there is a perturbation matrix **P** such that $\mathbf{P}\mathbf{A}\mathbf{P}^T$ can be written as a 2×2 matrix with the diagonal matrices being diagonal. For example, every block tridiagonal matrix with non-singular diagonal blocks is consistently-ordered and 2-cyclic.

Remark 2: The black-red Gauss-Seidel algorithm can be implemented in two parallel steps: one for updating the red points, and one for the black points. This is equivalent to two parallel Jacobi steps per iteration. Therefore, the total number of parallel steps between the Jacobi algorithm and the black-red Gauss-Seidel algorithm is the same given that the latter converges twice as fast as the former. However, the black-red algorithm operates on half the number of points. Overall, communications and latency are the factors that will ultimately determine if there is a substantial speed-up factor using the black-red algorithm. A better approach is to use the black-red coloring idea with an overelaxation method, as we discuss next.

7.2.5 Successive Acceleration Techniques - SOR

The convergence of the Gauss-Seidel method can be effectively accelerated if we introduce a relaxation procedure of the form

$$\Theta_{ij}^{n+1} = \Theta_{ij}^n + \omega d_{ij}^n, \quad d_{ij}^n \equiv \hat{\Theta}_{ij}^{n+1} - \Theta_{ij}^n$$

$$(7.14)$$

where $\hat{\Theta}_{ij}^{n+1}$ is predicted by the standard Gauss-Seidel method. This intermediate answer is then corrected to obtain Θ_{ij}^{n+1} ; such a method is called *predictor-corrector*. The displacement d_{ij}^n is given by

$$d_{ij}^{n} = \frac{1}{4} (\Theta_{i+1,j}^{n} + \Theta_{i,j+1}^{n} - 4\Theta_{ij}^{n} + \Theta_{i,j-1}^{n+1} + \Theta_{i-1,j}^{n+1} + \Delta x^{2} q_{ij}).$$

Upon substitution in equation (7.14), the corrected value is

$$\Theta_{ij}^{n+1} = \frac{\omega}{4} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n + \Theta_{i,j-1}^{n+1} + \Theta_{i-1,j}^{n+1} + \Delta x^2 q_{ij}) + (1 - \omega) \Theta_{ij}^n.$$

This method converges for $0 < \omega < 2$ assuming that the corresponding matrix **A** is positive-definite and symmetric.

- For $\omega = 1$, we recover the standard Gauss-Seidel;
- For $\omega > 1$, we have over-relaxation (SOR), and
- For $\omega < 1$ we have under-relaxation.

The latter is not used often in practice because of its slow convergence.

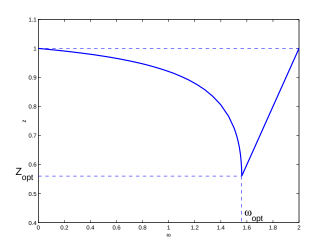


Figure 7.21: Convergence rate versus relaxation parameter for SOR.

For the diffusion equation (7.12) we can obtain the *optimum* relaxation parameter that maximizes the convergence rate. For central differencing we have:

$$\omega_{opt} = \frac{2}{(1 + \sqrt{1 - z_J^2})}$$
 and $z_{opt} = \frac{z_J^2}{(1 + \sqrt{1 - z_J^2})^2}$,

where z_J is the convergence rate of the Jacobi method for the same discretization. In particular, for $\Delta x = \Delta y$ the Jacobi convergence rate is $z_J = \cos \pi \Delta x$, and thus we obtain

$$\omega_{opt} = \frac{2}{1 + \sin \pi \Delta x} \approx 2(1 - \pi \Delta x),$$

and correspondingly

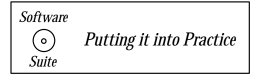
$$z_{opt} = \frac{\cos^2 \pi \Delta x}{(1 + \sin \pi \Delta x)^2} \approx 1 - 2\pi \Delta x$$
.

This approximation provides an estimate for the number of iterations to reach convergence with accuracy of about d correct digits, thus

Number of
$$SOR(\omega)$$
 iterations: $n \sim \frac{-d}{\ln z_{opt}} \sim N$

since $\Delta x = 1/N$. This implies, that for $SOR(\omega_{opt})$ we have that $n \sim \mathcal{O}(N)$ and thus the computational work is $\mathcal{O}(N^3)$ versus $\mathcal{O}(N^4)$ for the standard Gauss-Seidel iteration assuming an $N \times N$ grid. We will revisit the issue of convergence rate of over-relaxation methods in section 7.2.8.

Remark: For general matrices **A** an estimate of the form $\omega_{opt} = 2 - \mathcal{O}(\Delta x)$, where Δx is the spacing is a good initial guess. Subsequent numerical experiments can provide more refined values of the optimum relaxation parameter by comparing corresponding values of the convergence rate z and searching for a local minimum. Unfortunately, the minimum occurs over a quite narrow range in ω , as it is shown in figure 7.21.



We now present a function SOR which performs successive over-relaxation to obtain the solution \mathbf{x} to the system $\mathbf{A}\mathbf{x} = \mathbf{b}$. The function takes as input the relaxation parameter omega, the size of the system N, a double** array containing the matrix \mathbf{A} , an array x into which the solution is to be placed, the right-hand-side stored in the array b, and the convergence tolerance abstol.

```
//set initial guess
  for(i=0; i<N; i++){
    x[i] = 1.0e0;
    xold[i] = 1.0e0;
  }
  for(k=0; k < maxit; k++){
    for(i = 0; i < N; i++){
      sum1 = 0.0; sum2 = 0.0;
      for(j=0; j < i; j++)
        sum1 = sum1 + A[i][j]*x[j];
      for(j=i+1; j < N; j++)
        sum2 = sum2 + A[i][j]*xold[j];
      x[i] = (1.0-omega)*xold[i] + omega*
              (-sum1 - sum2 + b[i])/A[i][i];
    }
    if(sqrt(dot(N,x,xold))<abstol){</pre>
      delete[] xold;
      return;
    }
    for(i=0; i<N; i++)
      xold[i] = x[i];
  }
  cerr << "SOR: Maximum Number of Interations Reached\n";</pre>
  delete[] xold;
  return;
}
```

7.2.6 Symmetric Successive Acceleration Techniques - SSOR

The standard SOR method assumes a preferential direction as it sweeps the domain from left to right. This bias may cause error accumulation but in addition the amplification matrix G of the SOR may have complex eigenvalues. This may not be desirable as certain acceleration schemes, e.g. Chebyshev acceleration that we study next, require real eigenvalues. The idea of the $symmetric\ SOR\ (SSOR)$ is then to sweep the domain in two passes, one from the left to right and subsequently from right to left. To this end, we write the standard SOR presented above in matrix form

$$(\mathbf{D} + \omega \mathbf{L})\hat{\mathbf{x}}^{n+1} = [(1 - \omega)\mathbf{D} - \omega \mathbf{U}]\mathbf{x}^n + \omega \mathbf{b},$$

to obtain the intermediate solution $\hat{\mathbf{x}}^{n+1}$.

The SOR iteration in the reverse direction is

$$(\mathbf{D} + \omega \mathbf{U})\mathbf{x}^{n+1} = [(1 - \omega)\mathbf{D} - \omega \mathbf{L}]\hat{\mathbf{x}}^{n+1} + \omega \mathbf{b},$$

to obtain the final solution \mathbf{x}^{n+1} at the (n+1) iteration. The optimum relaxation parameter for SSOR is different than the ω_{opt} of SOR and it is not known exactly. An approximation proposed in [95] is

$$\omega_{opt} \approx \frac{2}{1 + \sqrt{2[1 - \rho(\mathbf{G})]}},$$

which leads to twice the convergence rate of the one-directional SOR, however the number of operations is also doubled. For the diffusion model that we consider we then have that

$$z_{SSOR} \approx 1 - \pi \Delta x \approx 1 - \frac{\pi}{N}$$
.

The corresponding amplification matrix is symmetric and has real eigenvalues with spectral radius

$$\rho_{SSOR}(\mathbf{G}) = 1 - \frac{\pi}{N}.$$

We will make use of this property of SSOR in the following section.

7.2.7 SSOR with Chebyshev Acceleration

What we have done so far is to convert the system $\mathbf{A}\mathbf{x} = \mathbf{b}$ to a fixed point iteration of the form

$$\mathbf{x}^{n+1} = \mathbf{G}\mathbf{x}^n + \mathbf{c}.$$

It is often the case, and we have seen that in the multi-step schemes for temporal discretization, that a linear combination of the iterates $\{\mathbf{x}^n\}$ converges faster than the last iterate. This is the idea of Chebyshev acceleration, and in the following we will see how Chebyshev polynomials come into play in this context; we will basically exploit their *minimax property* (see section 3.1.5). Let us define the linear combination

$$\mathbf{y}_n = \sum_{q=0}^n \gamma_n^q \mathbf{x}^q$$

and form the error with respect to the exact solution \mathbf{x}

$$\mathbf{y}_n - \mathbf{x} = \sum_{q=0}^n \gamma_n^q (\mathbf{x}^q - \mathbf{x}) = \sum_{q=0}^n \gamma_n^q \mathbf{G}^q (\mathbf{x}^0 - \mathbf{x}) = p_n(\mathbf{G})(\mathbf{x}_0 - \mathbf{x}).$$

Here we have defined the polynomial p_n of degree n with $p_n(1) = 1$ since the interpolation coefficients satisfy

$$\sum_{q=0}^{n} \gamma_n^q = 1.$$

In order to minimize the computational work we should choose the polynomial p_n carefully. We know (using the spectral mapping theorem) that the eigenvalues of $p_n(\mathbf{G})$ are

given by $p_n(\lambda(\mathbf{G}))$. We then need to minimize these eigenvalues, or even simpler we need to minimize the maximum eigenvalue (spectral radius ρ). The polynomials that best satisfy this requirement are the *Chebyshev polynomials* due to their *minimax property*. We have also assumed in the above that the eigenvalues of the amplification matrix \mathbf{G} are real.

Let us then assume that $x \in [-\rho, \rho]$ and define the polynomial

$$p_n(x) \equiv \frac{T_n(x/\rho)}{T_n(1/\rho)} = \frac{T_n(x/\rho)}{C_n},$$

where the last equation defines the normalization constant C_n . We will now use the threeterm recursive property of Chebyshev polynomials to obtain y_n recursively and construct the convergence acceleration algorithm. Note that C_n can also be obtained recursively in a similar way, i.e.,

$$C_n = \frac{2}{\rho} C_{n-1} - C_{n-2} \,.$$

For the error $(\mathbf{y}^n - \mathbf{x})$ we have

$$\mathbf{y}^{n} - \mathbf{x} = p_{n}(\mathbf{G})(\mathbf{x}^{0} - \mathbf{x})$$

$$= C_{n}^{-1}T_{n}(\mathbf{G}/\rho)(\mathbf{x}^{0} - \mathbf{x})$$

$$= C_{n}^{-1}[2 \cdot \mathbf{G}/\rho \cdot T_{n-1}(\mathbf{G}/\rho)(\mathbf{x}^{0} - \mathbf{x}) - T_{n-2}(\mathbf{G}/\rho)(\mathbf{x}^{0} - \mathbf{x})]$$

$$= C_{n}^{-1}[2 \cdot \mathbf{G}/\rho \cdot C_{n-1}p_{n-1}(\mathbf{G}/\rho)(\mathbf{x}^{0} - \mathbf{x}) - C_{n-2}p_{n-2}(\mathbf{G}/\rho)(\mathbf{x}^{0} - \mathbf{x})]$$

$$= C_{n}^{-1}[2 \cdot \mathbf{G}/\rho \cdot C_{n-1}(\mathbf{y}^{n-1} - \mathbf{x}) - C_{n-2}(\mathbf{y}^{n-2} - \mathbf{x})].$$

We can now write the recursive relation

$$\mathbf{y}^{n} = 2\frac{C_{n-1}}{C_{n}} \frac{\mathbf{G}}{\rho} \mathbf{y}^{n-1} - \frac{C_{n-2}}{C_{n}} \mathbf{y}^{n-2} + \mathbf{D}^{n}$$
(7.15)

where we have defined

$$\mathbf{D}^n = \frac{2}{\rho} \frac{C_{n-1}}{C_n} \mathbf{c}$$

and we recall the iteration $\mathbf{x}^{n+1} = \mathbf{G}\mathbf{x}^n + \mathbf{c}$.

We now have a recursion relation but we still need to initialize the first guess and the normalization constants. From the definition we have that $C_0 = 1$ and $C_1 = 1/\rho$, and we set $\mathbf{y}^1 = \mathbf{G}\mathbf{x}^0 + \mathbf{c}$.

We can also estimate the number of iterations by observing that in the above algorithm the error is multiplied in each iteration by the *inverse of*

$$C_n \equiv T_n(1/\rho).$$

For the SSOR algorithm we have that $\rho \approx 1 - \frac{\pi}{N}$ and thus $1/\rho \approx 1 + \frac{\pi}{N}$. Now, from the Chebyshev properties we have that

$$T_n(1+\pi/N) \ge \frac{1}{2}(1+n\sqrt{2\pi/N}).$$

In order to multiply the error by a factor less than one, from the above we need

$$\frac{2}{1 + n\sqrt{2\pi/N}} \le 1,$$

and thus the number of iteration n scales as $n \propto \mathcal{O}(\sqrt{N})$. The total computational work for the SSOR with Chebyshev acceleration is therefore:

$$\mathcal{O}(N^{5/2})$$
 versus $\mathcal{O}(N^3)$ for SOR.

We see that SSOR with Chebyshev acceleration beats even the direct solver, i.e. Gaussian elimination, which is is $\mathcal{O}(N^3)$ for the backsolve.

7.2.8 Convergence Analysis of Iterative Solvers

Basic Analysis

We have already discussed some of the conditions necessary for convergence of the basic iterative algorithms presented in this chapter. In general, if the coefficient matrix \mathbf{A} is symmetric and positive-definite the Gauss-Seidel iteration converges to a unique solution for any initial vector \mathbf{x}^0 although for the Jacobi iteration a stronger condition is needed. We have also seen the relation between the time-space stencils for diffusion and the iteration algorithms. In this section, we provide some more details on the convergence of the basic iterative solvers.

We start by re-stating the **analogy** between iterative procedures and equations of evolution. This analogy was formulated rigorously by Garabedian [41], who considered the following equation

$$K\frac{\partial\Theta}{\partial t} = \kappa \frac{\partial^2\Theta}{\partial x^2} - U\frac{\partial\Theta}{\partial x},\tag{7.16}$$

which is an advection-diffusion equation. By exploiting the properties of this PDE one can obtain information about the convergence of the discrete iterative algorithms. It can be shown, for example, that the smaller K is the more rapid the convergence is. Here, K depends on the specific iterative procedure considered. Convergence to steady solution is equivalent to existence of solutions of the PDE. A necessary condition for convergence is that equation (7.16) be parabolic for t > 0. This condition implies that

- For Jacobi: $K_J = 2\kappa \frac{\Delta t}{\Delta x^2} > 0$.
- For Gauss-Seidel: $K_{GS} = \frac{\kappa \Delta t}{\Delta x^2} \left(1 \frac{U \Delta x}{2\kappa} \right) > 0$, $(U \Delta x < 2\kappa)$.
- For SOR: $K_{SOR} = \frac{\kappa \Delta t}{\Delta x^2} \left(\frac{2-\omega}{\omega} \frac{U\Delta x}{2\kappa} \right) > 0.$

However, K > 0 is not sufficient for convergence as we need to have a stable scheme too.

The case of $U \neq 0$ will be studied in the next chapter. Here by setting U = 0 we obtain

$$K_J = 2K_{GS}$$
, and $\omega K_{SOR} = (2 - \omega)K_{GS}$.

For the SOR iteration, convergence is obtained if

$$0 < \frac{2-\omega}{\omega} < 1 \Rightarrow 2-\omega < \omega \Rightarrow \omega > 1 \text{ and } \omega < 2.$$

We now generalize the iteration algorithms by introducing the concept of *preconditioner*. Let us consider the system

$$\mathbf{A}\Theta = \mathbf{q}$$

where A is a stiffness matrix, e.g. the discrete Laplacian as before. Then, we define the convergence (vector) error:

$$\epsilon^n = \Theta^n - \Theta^\infty$$
.

which we will analyze in the following. We split the spatial discretization matrix

$$A = M + N$$

and set up the iteration

$$\mathbf{M}\Theta^{n+1} = \mathbf{q} - \mathbf{N}\Theta^n$$

or

$$\mathbf{M}\Delta\Theta^n = -\mathbf{r}^n, \quad \mathbf{r} \equiv \mathbf{A}\Theta - \mathbf{q}$$

where \mathbf{r} is the *residual*. The above equation is derived from

$$\Delta\Theta^n \equiv \Theta^{n+1} - \Theta^n \Rightarrow \mathbf{M}(\Theta^{n+1} - \Theta^n) = \underbrace{\mathbf{q}^n - (\mathbf{M} + \mathbf{N})\Theta^n}_{-r^n}.$$

We can recover all the schemes presented earlier by the proper choice of the preconditioner \mathbf{M} , i.e.,

- Jacobi: $\mathbf{M} = \mathbf{D}$.
- Gauss-Seidel: $\mathbf{M} = \mathbf{D} + \mathbf{L}$.
- SOR: $\mathbf{M} = \mathbf{D} + \omega \mathbf{L}$.
- SSOR: $\mathbf{M} = (\mathbf{D} + \omega \mathbf{L}) \mathbf{D}^{-1} (\mathbf{D} + \omega \mathbf{U})$.

Next, we obtain the error equation by subtracting the equations

$$\begin{split} \mathbf{M} \Theta^{n+1} &= \mathbf{q}^n - \mathbf{N} \Theta^n \\ (\mathbf{M} + \mathbf{N}) \Theta^{\infty} &= \mathbf{q} \end{split}$$

to get

$$\mathbf{M}\epsilon^{n+1} = -\mathbf{N}\epsilon^n.$$

This leads us to the definition of the *amplification matrix*, which we have already used earlier , i.e.

$$\epsilon^{n+1} = -\mathbf{M}^{-1}\mathbf{N}\epsilon^{n} = (-\mathbf{M}^{-1}\mathbf{N})^{n}\epsilon^{1}$$

$$= \underbrace{(\mathbf{I} - \mathbf{M}^{-1}\mathbf{A})^{n}}_{\mathbf{G}: \text{ amplification matrix}} \epsilon^{1}.$$

Therefore

$$\epsilon^{n+1} = \mathbf{G}^{n+1} \epsilon^0. \tag{7.17}$$

For stability we require that the spectral radius of the amplification matrix be less than one, i.e. ,

$$\rho(\mathbf{G}) < 1$$

since $\rho(\mathbf{A}) \leq ||\mathbf{A}||$ for any matrix norm.

This condition can be applied effectively as it may be sufficient to know the eigenvalues of the matrices that compose the original matrix \mathbf{A} . For example, for the Jacobi iteration we have

$$\mathbf{G}_J = \mathbf{I} - \mathbf{M}^{-1} \mathbf{A} \Rightarrow \lambda_J = 1 - (1/d_j) \lambda_j(\mathbf{A}),$$

where d_j are the diagonal elements of matrix **D**, and we have made use of the spectral mapping theorem.

We can now derive the **error-residual** relationship by subtracting the equations

$$\mathbf{r}^{n} = -\mathbf{q}^{n} + (\mathbf{M} + \mathbf{N})\Theta^{n}$$
$$0 = -\mathbf{q}^{n} + (\mathbf{M} + \mathbf{N}))\Theta^{\infty}$$

where the second equation is valid in steady-state. Upon subtraction we obtain

$$\mathbf{r}^n = \mathbf{A}\epsilon^n$$

which is a very useful equation relating the error and the residual.

Remark 1: When the residual \mathbf{r}^n is reduced to machine accuracy it does not mean that the solution Θ^n is within machine accuracy, because we still have to account for the discretization error.

Remark 2: To compute the matrix-vector products in the iterations we can use either of two ways suggested by the equations

$$\mathbf{w} = \mathbf{M}^{-1}\mathbf{A}\mathbf{v} = \mathbf{M}^{-1}(\mathbf{M} + \mathbf{N})\mathbf{v} = \mathbf{v} + \mathbf{M}^{-1}\mathbf{N}\mathbf{v} \,.$$

That is, we compute \mathbf{w} from

$$(I). \quad \left\{ \begin{array}{l} \mathbf{r} = \mathbf{A}\mathbf{v} \\ \mathbf{w} = \mathbf{M}^{-1}\mathbf{r} \end{array} \right.$$

or from

(II).
$$\begin{cases} \mathbf{r} = \mathbf{N}\mathbf{v} \\ \mathbf{w} = \mathbf{M}^{-1}\mathbf{r} \\ \mathbf{w} = \mathbf{v} + \mathbf{w} \end{cases}$$

It is more efficient to use approach (II) as it involves matrix-vector multiplies with matrices, which are substantially less dense than the original matrix \mathbf{A} .

Fourier Representation of Convergence Error

We can decompose the convergence error into normal modes by considering the eigenstructure of the matrix $\bf A$. For clearer illustration of the main points, we will assume that the amplification matrix $\bf G$ shares eigenvectors with the coefficient matrix $\bf A$. This is true, for example, for the Jacobi algorithm on a discrete Laplacian but not for the Gauss-Seidel iteration. With this assumption we can represent the convergence error as

$$\epsilon^n = \sum_j \lambda_j^n \epsilon_j^0 \Phi_j \,, \tag{7.18}$$

where ϵ_j^0 is the initial error distribution, and λ_j , Φ_j are associated with the matrices \mathbf{G} and \mathbf{A} , respectively. In particular, Φ_j are the eigenvectors of the discrete diffusion operator. This representation is possible because the error behaves like the homogeneous part of the solution. The eigenvectors Φ_j are also eigenvectors of \mathbf{G} only for special cases, e.g., periodic boundary conditions and Jacobi iteration, but not for Gauss-Seidel, as stated above.

As n increases, the low λ_j (corresponding to high frequencies) will decay faster. Asymptotically, after a few iterations, we are left with the highest λ_j which correspond to low frequencies. The convergence history of these methods would then have the form shown in figure 7.22: a very fast decay is obtained initially followed by an extremely slow decay.

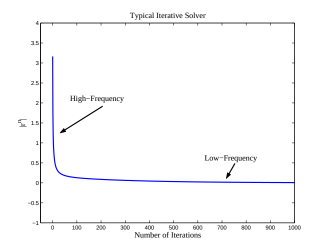


Figure 7.22: Typical convergence history of basic iterative solvers.

We can use equation (7.18) for the error to obtain a measure of convergence of the iterative solvers. To this end, we recall the *error-residual* relationship and substitute the

Fourier representation for the error

$$\mathbf{r}^{n} = \sum_{j} \lambda_{j}^{n}(\mathbf{G}) \epsilon_{j}^{0} \mathbf{A} \Phi_{j} = \sum_{j} \lambda_{j}^{n}(\mathbf{G}) \epsilon_{j}^{0} \underbrace{\lambda_{j}(\mathbf{A})}_{\text{eigenvalues of } A} \Phi_{j}.$$

If Φ_j are orthonormal eigenvectors, then we can obtain a simple expression for the L_2 -norm of the residual, i.e.,

$$\| \mathbf{r}^n \|_{L_2} = \left[\sum (\lambda_j^n(\mathbf{G}) \lambda_j(\mathbf{A}) \epsilon_j^0)^2 \right]^{1/2}.$$

This scalar residual is typically used as a measure of convergence of iterative solvers.

We can also use the eigenvalues of the amplification matrix to gain more insight into the convergence rate of individual modes (frequencies) for the iterative methods we presented. For example, in the Jacobi iteration

$$\lambda(\mathbf{G}_J) = 1 - (\sin^2 \phi_x / 2 + \sin^2 \phi_y / 2) = \frac{1}{2} (\cos \phi_x + \cos \phi_y), \qquad (7.19)$$

which is obtained from

$$G_J = I - M^{-1}A$$
, $J = k + (\ell - 1)N$, where $\ell, m = 1, ..., N - 1$

and $\phi_x = k\pi/N, \phi_y = \ell\pi/N$.

From equation (7.19) we see that at low frequencies, $\phi_x \simeq 0$, $\phi_y \simeq 0$, the damping rate (i.e., convergence rate) is very poor as $\lambda \sim 1$. At intermediate frequencies $(k, \ell \sim N/2)$ damping is very strong, and at high frequencies $(k, \ell \to N)$, damping is very weak again. Specifically, the most resistant error corresponds to $k = \ell = 1$ thus

$$\rho(\mathbf{G}_J) = \cos\frac{\pi}{N} \Rightarrow \rho(\mathbf{G}_J) \cong 1 - \frac{\pi^2}{2N^2}.$$

This analysis suggests that the number of iterations is proportional to $\mathcal{O}(N^2)$, as we have already seen earlier.

Following a similar analysis, we obtain that for the Gauss-Seidel iteration we have

$$\lambda(\mathbf{G}_{GS}) = \frac{1}{4}(\cos\phi_x + \cos\phi_y)^2 = \lambda^2(\mathbf{G}_J),\tag{7.20}$$

which shows that the Gauss-Seidel iteration converges twice as fast compared to the Jacobi iteration.

7.2.9 Relaxed Jacobi and Gauss-Seidel

We can also apply the above analysis to relaxation or weighted methods. Let us first introduce the $relaxed\ Jacobi$ algorithm:

$$\Theta_{ij}^{n+1} = \frac{\omega}{4} (\Theta_{i+1,j}^n + \Theta_{i,j+1}^n + \Theta_{i,j-1}^n + \Theta_{i-1,j}^n + q_{ij}^n) + (1 - \omega)\Theta_{ij}^n.$$

Thus, the new amplification matrix is

$$\mathbf{G}_{J(\omega)} = (1 - \omega)\mathbf{I} + \omega \mathbf{G}_J,$$

and its corresponding eigenvalues are

$$\lambda(\mathbf{G}_{J(\omega)}) = (1 - \omega) + \omega\lambda(\mathbf{G}_J). \tag{7.21}$$

For convergence we require that

$$\rho(\mathbf{G}_{J(\omega)}) \le |1 - \omega| + \omega \rho(\mathbf{G}_J) < 1$$

or

$$|1 - \omega| < 1 - \omega \rho(\mathbf{G}_J) \Rightarrow -[1 - \omega \rho(\mathbf{G}_J)] < 1 - \omega < 1 - \omega \rho(\mathbf{G}_J),$$

and thus

$$0 < \omega < \frac{2}{1 + \rho(G_J)}.$$

Therefore, in order to damp high frequencies we should select $\omega < 1$ and ω towards λ_{\min} . To damp low frequencies we should select $\omega > 1$ and ω towards λ_{\max} .

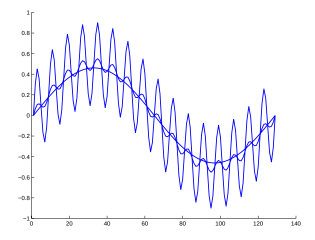


Figure 7.23: Jacobi algorithm with initial condition $(\Phi_2 + \Phi_{32})/2$. The curves correspond to the initial condition (large oscillations); results after 10 iterations (small oscillations); and results after 100 oscillations (smooth curve); $\omega = 2/3$.

In order to better understand this *selective damping* of the Fourier components of the error for the Jacobi method, we perform one-dimensional experiments. To this end, we revisit the Dirichlet problem

$$-\Theta_{j-1} + 2\Theta_j - \Theta_{j+1} = 0, \quad 1 \le j \le N - 1$$

$$\Theta_0 = \Theta_N = 0$$

so the corresponding matrix **A** is tridiagonal with elements (-1, 2, -1). The eigenvalues of this matrix are

$$\lambda_k(\mathbf{A}) = 4\sin^2(\frac{k\pi}{2N}), \quad 1 \le k \le N - 1, \tag{7.22}$$

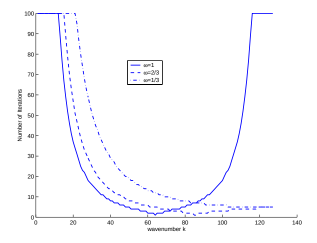


Figure 7.24: Number of Jacobi iterations to reduce the error 100 times for different values of ω . The initial condition is a single Laplace mode with wavenumber k.

and the corresponding eigenvectors for the grid points $0 \le j \le N$ are:

$$\Phi_j^k(\mathbf{A}) = \sin(\frac{jk\pi}{N}), \quad 1 \le k \le N - 1. \tag{7.23}$$

We have already seen these results in section 7.1.4 earlier in this chapter.

For the relaxed Jacobi algorithm we can derive from equation (7.21) the following

$$\lambda_k(\mathbf{G}_{J(\omega)}) = 1 - \frac{\omega}{2} \lambda_k(\mathbf{A}),$$

and thus

$$\lambda_k(\mathbf{G}_{J(\omega)}) = 1 - 2\omega \sin^2(\frac{k\pi}{2N}), \quad 1 \le k \le N - 1.$$
 (7.24)

We can now examine what is the best value for ω , i.e., the one that obtains

$$|\lambda_k(\mathbf{G}_{J(\omega)})| = minimum, \quad \forall \, 1 \le k \le N-1.$$

By inspection we can see that the *smooth components* of the error, i.e., the modes corresponding to $1 \le j \le N/2$, are affected very little by ω . In contrast, the damping of the rough components corresponding to $N/2 \le k \le N-1$ are affected significantly. We thus seek an optimum value of ω for the rough modes by limiting the above minimization to the upper half of the spectrum, i.e., $N/2 \le k \le N-1$. This minimum is achieved if

$$\lambda_N(\mathbf{G}_{J(\omega)}) = -\lambda_{(N/2)}(\mathbf{G}_{J(\omega)}),$$

which gives the optimum $\omega = 2/3$ for the one-dimensional problem, The corresponding reduction of the error is

$$\lambda_k(\omega=2/3)=1/3\,,$$

which means that the rough components will be reduced by a factor of three in each Jacobi iteration.

The effectiveness of this choice is shown in the next two figures. In figure 7.23 we plot the evolution of the solution from the relaxed Jacobi algorithm starting from the initial condition

$$\frac{1}{2}(\Phi_2 + \Phi_{32})$$

after 10 and also after 100 iterations with the relaxed Jacobi algorithm and $\omega = 2/3$. We see that the initially very rough solution relaxes to the smooth mode two, which remains nearly the same. The correct solution is $\Theta_j = 0$, j = 1, ..., N, since we solve the homogeneous one-dimensional Laplace's equation with zero Dirichlet conditions. A more comprehensive picture is shown in figure 7.24 where we record the number of relaxed Jacobi iterations to reduce the mean error by a factor of 100. Specifically, the initial condition here is a single Fourier mode, which coincides with the eigenmode of the Laplacian and the eigenmode of the relaxed Jacobi iteration matrix $\mathbf{G}_{J(\omega)}$. The choice of $\omega = 1$ is most effective for the intermediate range of modes while the choice $\omega = 2/3$ damps effectively the upper half of the spectrum. The actual number of iterations for the low modes is much higher than 100, in fact of the $\mathcal{O}(1000)$, but here we set a maximum 100 iterations for clarity of the plot.

Next we analyze the Gauss-Seidel over-relaxation method. We use the matrix form

$$(\mathbf{D} + \omega \mathbf{L}) \Delta \Theta^n = -\omega \mathbf{r}^n,$$

so the corresponding amplification matrix is

$$\mathbf{G}_{SOR}(\omega) = \mathbf{I} - \omega(\mathbf{D} + \omega \mathbf{L})^{-1}\mathbf{A}$$

or

$$\mathbf{G}_{SOR}(\omega) = \underbrace{(\mathbf{D} + \omega \mathbf{L})^{-1}}_{triangular} \underbrace{[(1 - \omega)\mathbf{D} - \omega \mathbf{U}]}_{triangular}.$$

Since $\mathbf{G}_{SOR}(\omega)$ is a product of triangular matrices, its determinant is the product of its diagonal element. Thus

$$\det \mathbf{G}_{SOR}(\omega) = \det(\mathbf{I} + \omega \mathbf{D}^{-1} \mathbf{L})^{-1} \det[(1 - \omega)\mathbf{I} - \omega \mathbf{D}^{-1} \mathbf{U}]$$
$$= 1 \cdot \det[(1 - \omega) - \omega \mathbf{D}^{-1} \mathbf{U}]$$
$$= (1 - \omega)^{N}$$

But

$$\det \mathbf{G}_{SOR}(\omega) = \prod_{i=1}^{N} \lambda_i \le \rho^N(\mathbf{G}_{SOR}(\omega))$$

so

$$|1 - \omega| \le \rho(\mathbf{G}_{SOR}(\omega)) < 1$$
,

and so for stability we recover the result $0 < \omega < 2$, as before.

Remark 1: For general symmetric **A** matrices we have that

$$\lambda(\omega) = 1 - \omega + \omega \lambda^{1/2}(\omega) \cdot \lambda(\mathbf{G}_J),$$

and for $\omega = 1 \Rightarrow \lambda(\omega = 1) = \lambda^2(\mathbf{G}_J)$, as before.

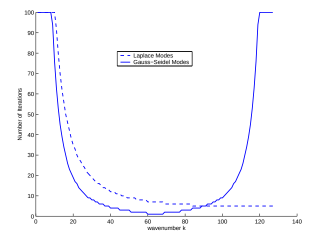


Figure 7.25: Number of Gauss-Seidel iterations to reduce the error 100 times for different initial conditions corresponding to single Laplace modes (dash line) and single Gauss-Seidel modes (solid line).

Remark 2: Unlike the Jacobi method, the Gauss-Seidel method does not share the same eigenvectors with the Laplacian matrix \mathbf{A} . Specifically, the eigenfunctions of the Gauss-Seidel iteration matrix \mathbf{G}_{GS} are

$$\Phi_j^k = \left[\cos(\frac{k\pi}{N})\right]^j \sin(\frac{jk\pi}{N}), \quad 1 \le k \le N - 1.$$

It is interesting to observe that the Gauss-Seidel method damps the Laplacian modes in a similar fashion to the relaxed Jacobi method, i.e. the rough components are damped faster. In contrast, the Gauss-Seidel eigenmodes are damped in a similar fashion as in the unrelaxed Jacobi method. This behavior is demonstrated in figure 7.25, where the number of iterations to reduce the mean error by a factor of 100 is recorded.

The selective damping achieved by the Jacobi and Gauss-Seidel methods is a typical property of all relaxation schemes. The Fourier error modes are determined by the grid spacing or equivalently the number of grid points. Therefore, a rough mode on a fine grid may appear as smooth mode on a coarse grid. This simple observation along with the selective damping of relaxation schemes is exploited in the multigrid method, one of the most effective solvers, that we present next.

7.2.10 The Multigrid Method

The multigrid method is not really a new algorithm, but rather a procedure to make the relaxation algorithms effective at *all* wavenumbers, including the low wavenumbers. The main idea of the multigrid procedure is based on the observation that the classical iterations rapidly kill high frequency (or wavenumber) errors but are slower for the low frequency, i.e., largest eigenvalues. Since this is all determined by the grid spacing, the idea of multigrid is to change the grid so that low frequencies on the fine grid look *higher* on the *coarser* grid. Conversely, errors that are invisible on the coarse grid are no problem on the fine grid. Figure

7.23 illustrates how the initial error consisting of both high (rough) and low (smooth) modes relaxes after a few iterations with a single grid discretization. The remaining low mode will appear as an intermediate or high mode in a discretization with a coarser grid. Relaxation on the new grid will effectively damp this mode as well.

The above discussion also suggests that the relaxation scheme will be more effective if we employ an initial condition that does not include smooth components. This can be accomplished by first solving the problem in a coarse grid and subsequently transfer the solution to the finer grid. Alternatively, we can start on a fine grid and subsequently transfer the solution into a coarser grid.

However, there is a further complication in these inter-grid transfers that we have to take into account. When we transfer from a fine grid to a coarse grid, the smooth modes become rough modes, but also the initially rough modes, although reduced in amplitude, are transformed (at least some of them) into smooth modes. This is the phenomenon of aliasing, which results in misrepresentation of the rough modes in the coarse grid after a fine-to-coarse grid transfer.

To make the above ideas more concrete let us introduce some notation. We will denote by h the fine grid and by H the coarse grid, and (in most cases) we will imply that we coarsen the grid by halving the number of points leading to grid spacing H = 2h. The question on how to *start* the inter-grid transfers, i.e., from a coarse or fine grid, that we posed above is also related to what exactly we should transfer from one grid to the other. Should that be

- The system $\mathbf{A}\mathbf{\Theta} = \mathbf{q}$ that we want to solve, or
- The residual $\mathbf{r} = \mathbf{A}\mathbf{\Theta} \mathbf{q}$ that we want to minimize?

Both ideas have been explored in multigrid research and have been used in practice for different problems.

The idea of transferring the original system is combined with the use of the coarse grid solution as an initial start, and it is called *nested iteration*. It consists of the following steps:

Nested Iteration Algorithm:

- 1. Relax $\mathbf{A}_H \mathbf{\Theta}_H = \mathbf{q}_H$ on a coarse grid to obtain an initial solution.
- 2. Transfer the solution Θ_H to the fine grid to obtain an initial condition Θ_h^0 .
- 3. Relax $\mathbf{A}_h \mathbf{\Theta}_h = \mathbf{q}_h$ using the initial condition $\mathbf{\Theta}_h^0$.
- 4. Test for convergence. If needed go to step 1 and continue.

In the above algorithm there can be several relaxation sweeps. Also, in each new execution of the algorithm the old fine grid is overwritten with the new coarse grid.

A second approach involves transferring the residual from one grid to the next starting with the fine grid. This is the *correction* method and it consists of the following steps:

Correction Algorithm:

- 1. On the fine grid, relax $\mathbf{A}_h \mathbf{\Theta}_h = \mathbf{q}_h$.
- 2. Transfer the residual \mathbf{r}_h to the coarser grid to obtain \mathbf{r}_H .
- 3. Obtain the correction $\delta \Theta_H$ by solving $\mathbf{A}_H(\delta \Theta_H) = \mathbf{r}_H$ at a reduced computational cost.
- 4. Transfer the correction to the fine grid in order to generate a new update $\Theta_h \leftarrow \Theta_h + \delta\Theta_H$.
- 5. Test for convergence. If needed go to step 1 and continue.

Similar to the nested iteration, in this algorithm also several coarser grids can be involved.

The above two algorithms make use of three important operations in multigrid:

- Relaxation of the solution or correction (Smoother).
- Transfer to coarse grid (*Restriction*).
- Transfer to fine grid (*Proplongation*).

In order to define a specific multigrid method we need to specify the corresponding three operators named in the above list (inside the parenthesis). We give details for each operators below.

The Smoother

To evaluate the smoothing properties of the relaxation method employed, e.g. relaxed Jacobi or Gauss-Seidel or any other variant, we need to examine the eigenspectrum of the amplification matrix \mathbf{G} . We have already seen in the previous section that we cannot really affect the low modes, and the best we can do is to come up with a good choice on how to most effectively damp the upper half of the eigenspectrum of \mathbf{G} . With this in mind, we define the *smoothing factor* as

$$\mu = \max |\lambda(\mathbf{G}(\phi))|, \quad \frac{\pi}{2} \le |\phi| \le \pi.$$
 (7.25)

Here $\phi = k\pi/N$; in multi-dimensions we have to extend this definition to account for ϕ_x, ϕ_y and ϕ_z which will depend on the resolution in x, y, and z, respectively.

Values of $|\phi|$ close to zero correspond to low wavenumbers while values of $|\phi|$ close to π correspond to high wavenumbers. In order to appreciate this correspondence between the angular variable ϕ and the spacing in the physical domain we sketch in figure 7.26 the one-dimensional discretization for a fine/coarse grid combination.

We have seen in the previous section that for the relaxed Jacobi method the eigenvalues are

$$\lambda_k(\mathbf{G}_{J(\omega)}) = 1 - 2\omega \sin^2(\phi/2) \,,$$

and thus the smoothing factor for the *one-dimensional* problem is

$$\mu = \lambda(\pi/2) = 1/3$$
 and $\omega = 2/3$.

h:
$$\Delta x = L/N \quad \phi \in [\pi/N, \pi]$$
H:
$$\Delta x = L/N \quad \phi \in [\pi/N, \pi/2]$$

$$\phi \quad \pi/N \quad \pi/2 \quad \pi$$
coarse grid

Figure 7.26: Fine/coarse grids in one-dimensional domain.

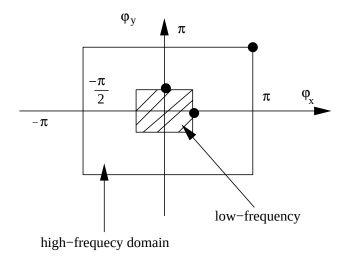


Figure 7.27: Frequency domain for estimating the Smoother.

In the two-dimensional case we also have to separate the domain into the high frequency (wavenumber) and low frequency (wavenumber) and formulate a minimax problem for the Smoother in the domain as shown diagrammatically in figure 7.27. We first note that the standard Jacobi algorithm has an eigenvalue of -1 at $\phi_x = \phi_y = \pi$ (in the high-frequency domain), and thus it is not a good smoother! However, the relaxed Jacobi can be used to damp high frequencies, as in the one-dimensional case.

In the high-frequency domain, the extreme eigenvalues of Jacobi are

$$\lambda(\mathbf{G}_J) \left\{ \begin{array}{ccc} -1 & \text{at} & (\pi, \pi) \\ +1/2 & \text{at} & (\pi/2, 0) \end{array} \right..$$

Then

$$\lambda(\mathbf{G}_{J(\omega)}) = [(1 - \omega) + \omega\lambda(G_J)]$$

and

$$\lambda(\mathbf{G}_{J(\omega)}) = \begin{cases} 1 - \omega + \omega/2 = 1 - \omega/2 \\ 1 - \omega - \omega = 1 - 2\omega \end{cases}.$$

This minimax problem can be solved graphically as shown in figure 7.28; we obtain for the optimum $\omega = 4/5 = 0.8$, with the corresponding smoothing factor

$$\mu_J = \text{minmax} \{ |1 - \omega/2|, |1 - 2\omega| \} = 0.6.$$

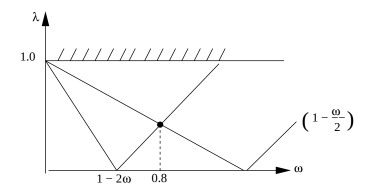


Figure 7.28: Estimation of the Smoother for the two-dimensional Jacobi method.

Remark 1: The total reduction of the error during the smoothing substep is proportional to μ^{n_1} where n_1 is the number of relaxation sweeps.

Remark 2: A similar but more elaborate analysis for Gauss-Seidel gives $\mu_{GS} = 0.5$ [92]. Therefore, the Gauss-Seidel Smoother achieves an order of magnitude reduction in the high frequency error after only three iterations.

Remark 3: In multi-dimensions we can relax by *points*, *lines*, or *planes* or by alternating directions and parallel versions such as the red-black coloring can be employed.

The Coarse-Grid Correction (CGC)

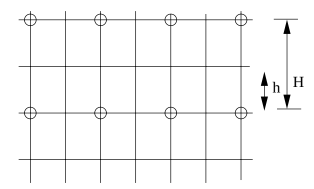


Figure 7.29: Two-grid transfer. The fine grid spacing is denoted by h and the coarse grid spacing is denoted by H.

The coarse-grid correction stage (CGC) is a very important operation and has to be done accurately and efficiently. To identify the main operations during this stage we define two grids, as shown in figure 7.29. Assuming that on the fine grid we have obtained

$$\mathbf{P}\left(\delta\mathbf{\Theta}_{h}\right)=\mathbf{r}_{h}$$

where **P** is an operator to be determined later, we perform the following steps in CGC:

• Transfer residual to coarse grid via the Restriction Operator, \mathbf{I}_h^H , i.e.,

$$\mathbf{r}_H = \mathbf{I}_h^H \mathbf{r}_h$$
.

- Solve: $\mathbf{A}_H(\delta \mathbf{\Theta}_H) = \mathbf{r}_H$ at low cost.
- Transfer correction $\delta \Theta_H$ to fine mesh via the *Prolongation Operator*, \mathbf{I}_H^h , i.e.,

$$\delta \mathbf{\Theta}_h = \mathbf{I}_H^h(\delta \mathbf{\Theta}_H) \,.$$

We have outlined the above multigrid substeps at a high level, and we now need to define the restriction and prolongation operators more precisely. We do that for simple finite difference representations following the two-grid model.

The **Restriction Operator** involves a projection from N points to N/2 points. The finite difference expressions for the residuals on the fine and coarse grids, respectively, are

$$r_{h,i} = \frac{1}{h^2} (\Theta_{i+1} - 2\Theta_i + \Theta_{i-1}) - q_i$$

$$r_{H,k} = \frac{1}{4h^2} (\Theta_{k-1} - 2\Theta_k + \Theta_{k+1}) - q_k$$

where i = 2k.

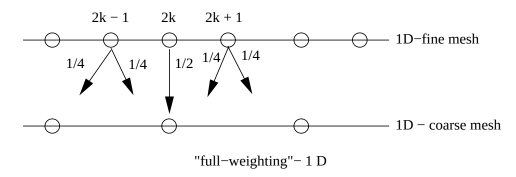


Figure 7.30: One-dimensional restriction operator (full-weighting).

Based on these residuals we can define a restriction operator that accomplishes the transfer; it is shown schematically in figure 7.30. This projection is referred to as *full-weighting* in contrast to a simpler projection by *injection*, which would imply that $\Theta_k^H = \Theta_i^h$. We can define more precisely the full-weighting restriction operator for the one-dimensional case in matrix form as follows:

which is an $(N/2) \times N$ matrix. The coarse grid residual is then obtained from

$$\mathbf{r}_H = \mathbf{I}_h^H \mathbf{r}_h$$
.

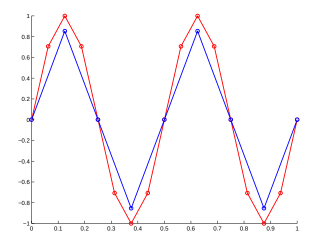


Figure 7.31: Action of a full weighting restriction operation on a smooth mode $(N=16;\,k=4)$.

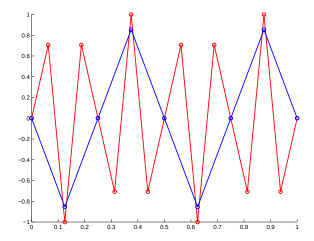


Figure 7.32: Action of a full weighting restriction operation on a rough mode $(N=16;\,k=12).$

Next, we explain how the restriction operator acts on the k modes of the Laplacian \mathbf{A}_h ; these modes are given by

 $\Phi_{k,j}^h = \sin(\frac{jk\pi}{N}), \quad 0 \le j \le N.$

In particular, the result is different depending on the mode, that is if it is a smooth or a rough mode. In the one-dimensional case for *smooth modes* we have that

$$\mathbf{I}_h^H \Phi_k^h = \cos^2(\frac{k\pi}{2N}) \, \Phi_k^H, \quad 1 \le k \le N/2.$$

This equation simply states that the projected k^{th} mode remains the k^{th} mode on the coarse grid. An example is shown in figure 7.31 where the k = 4 mode is projected onto a grid with 8 grid points from a grid of 16 points.

In contrast, for the high rough modes aliasing may transform them to smooth modes on the coarse grid according to

$$\mathbf{I}_{h}^{H} \Phi_{k}^{h} = -\sin^{2}(\frac{k\pi}{2N}) \Phi_{N-k}^{H}, \quad N/2 < k \le N.$$

This is shown schematically in figure 7.32 where the k = 12 mode is projected onto a grid with 8 grid points from a grid of 16 points. It is clear that the projected mode is a (16 - 12) = 4 mode, i.e., a smooth mode in the coarse grid.

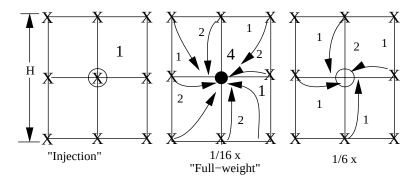


Figure 7.33: Three different schemes of restriction from fine to coarse grids. The full-weighting is the most popular choice.

In two- or three-dimensions, several choices for the restriction operator are possible representing different ways of weighting the contribution from the fine grid nodes to the coarse grid nodes. Examples are shown in figure 7.33. The full-weighting is the most popular choice as it leads to the following important property between the restriction and prolongation operators

$$\mathbf{I}_h^H = C\,\mathbf{I}_H^h\,,$$

which means that the restriction and interpolation operators are transpose of each other apart from a constant C.

We now examine some of the properties of the interpolation or **Prolongation Operator** that we need to define in order to transfer quantities, e.g. the correction, from the coarse to fine grid.

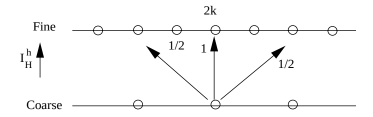


Figure 7.34: Schematic for the one-dimensional prolongation operation.

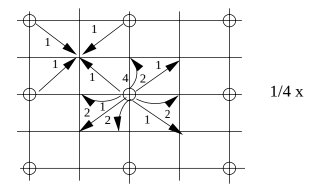


Figure 7.35: Schematic for the two-dimensional prolongation operation.

The one-dimensional prolongation operation is defined as shown in figure 7.34, while the two-dimensional prolongation operation is sketched in figure 7.35. They both involve *linear* interpolation, which is effective if the solution error is relatively smooth. A more accurate prolongation operator based on *cubic* interpolation has also been found to be very effective.

For the one-dimensional problem the prolongation matrix is defined as follows:

$$\mathbf{I}_{H}^{h} = \left| \begin{array}{ccccc} & k-1 & k & k+1 \\ & 0 & 1/2 & 1/2 & \dots \\ & 0 & 0 & 1 & 0 \\ & & 1/2 & 1/2 & 2k+1 \end{array} \right| \begin{array}{c} 2k-1 \\ 2k \\ 2k+1 \end{array}$$

In order to understand the action of the prolongation operator for the one-dimensional case, we examine how the modes of the Laplacian on the coarse grid are interpolated onto the fine grid. Specifically, we find that

$$\mathbf{I}_{H}^{h} \Phi_{k}^{H} = \left[\cos^{2}\left(\frac{k\pi}{2N}\right)\right] \Phi_{k}^{h} - \left[\sin^{2}\left(\frac{k\pi}{2N}\right)\right] \Phi_{N-k}^{h}.$$

This equation provides a lot of insight into the coarse-to-fine grid transfer. It states that the k mode is not preserved but is mixed up with the complimentary mode (N-k), so the smooth mode picks up some rough component as it is transferred onto the fine grid! This, in turn, suggests that after we transfer the correction $\delta \Theta_h$ to the fine grid we need to perform additional relaxation sweeps to get rid of this unwanted component. Fortunately, the rough component is of smaller magnitude compared to the smooth component as indicated from

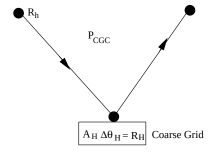


Figure 7.36: Schematic of the coarse-grid correction stage.

the approximation to the above equation, i.e.,

$$\mathbf{I}_{H}^{h} \Phi_{k}^{H} = [1 - \mathcal{O}(\frac{k^{2}}{N^{2}})] \Phi_{k}^{h} - [\mathcal{O}(\frac{k^{2}}{N^{2}})] \Phi_{N-k}^{h},$$

which is valid for $k \ll N/2$.

We now return to the coarse-grid correction operator \mathbf{P}_{CGC} , which we will define by the following "backwards" analysis. We begin with the correction at the fine grid and trace it back to the two-grid correction, see figure 7.36, as follows:

$$\delta \mathbf{\Theta}_{h} = \mathbf{I}_{H}^{h} (\delta \mathbf{\Theta}_{H})$$

$$= \mathbf{I}_{H}^{h} \mathbf{A}_{H}^{-1} \mathbf{r}_{H}$$

$$= \mathbf{I}_{H}^{h} \mathbf{A}_{H}^{-1} \mathbf{I}_{h}^{H} \mathbf{r}_{h}$$

$$\Rightarrow \underbrace{(\mathbf{I}_{H}^{h} \mathbf{A}_{H}^{-1} \mathbf{I}_{h}^{H})^{-1}}_{\mathbf{P}_{GGG}} \delta \mathbf{\Theta}_{h} = \mathbf{r}_{h}.$$

The corresponding amplification matrix or two-grid correction operator is then

$$\mathbf{G}_{CGC} = \mathbf{I} - \mathbf{P}_{CGC}^{-1} \mathbf{A}_h$$
$$= \mathbf{I} - \mathbf{I}_H^h \mathbf{A}_H^{-1} \mathbf{I}_h^H \mathbf{A}_h.$$

It is interesting again to examine what this operator does to the Laplacian modes, both smooth and rough. Combining the properties of the restriction and prolongation operators, we obtain

$$\mathbf{G}_{CGC}\Phi_{k} = [\sin^{2}(\frac{k\pi}{(2N)})]\Phi_{k} + [\sin^{2}(\frac{k\pi}{(2N)})]\Phi_{N-k}$$

$$\mathbf{G}_{CGC}\Phi_{N-k} = [\cos^{2}(\frac{k\pi}{(2N)})]\Phi_{k} + [\cos^{2}(\frac{k\pi}{(2N)})]\Phi_{N-k}.$$

The above equations simply state that the coarse-grid correction operator mixes up the modes irrespective of the mode (smooth or rough) to which it is applied, unlike, for example, the restriction operator. However, the amplitudes are different depending on whether the original mode is smooth or rough, and this is really what saves the day for multigrid! In

particular, when \mathbf{G}_{CGC} operates on smooth modes the amplitudes of the resulted smooth and rough modes are proportional to $\mathcal{O}(k^2/N^2)$ (for $k \ll N/2$), in contrast with the high modes whose resulted modes have amplitudes of $\mathcal{O}(1)$.

The above analysis is useful as it also suggests the potential benefits of relaxation. To this end, we recall that most of the relaxation solvers do not mix up the modes. Assuming, for example, that we employ the relaxed Jacobi method as smoother, and we perform n_1 relaxation sweeps then the amplification matrix is

$$\mathbf{G}_{CGC}\cdot\mathbf{G}_{J(\omega)}^{n_1}$$
 .

The corresponding action of this operator to the Laplacian modes is then given by

$$\mathbf{G}_{CGC} \cdot \mathbf{G}_{J(\omega)}^{n_1} \cdot \Phi_k = [\sin^2(\frac{k\pi}{(2N)})] \lambda_k^{n_1} \Phi_k + [\sin^2(\frac{k\pi}{(2N)})] \lambda_k^{n_1} \Phi_{N-k}$$

$$\mathbf{G}_{CGC} \cdot \Phi_{N-k} = [\cos^2(\frac{k\pi}{(2N)})] \lambda_{N-k}^{n_1} \Phi_k + [\cos^2(\frac{k\pi}{(2N)})] \lambda_{N-k}^{n_1} \Phi_{N-k} ,$$

where λ_k are the eigenvalues of the Jacobi amplification matrix $\mathbf{G}_{J(\omega)}$, which are, of course, all less than one. Therefore, relaxation reduces significantly the already small amplitudes of the smooth modes, but the more significant effect is on the rough modes, since $\lambda_{N-k} < \lambda_k$ for small wavenumbers.

Multigrid Algorithm

We now have all the ingredients required in order to write the complete multigrid algorithm. Two popular choices of multigrid algorithms are shown schematically in figure 7.37, corresponding to a *V-cycle* and a *W-cycle*. They both start with a fine grid and transfer the residual down to the coarsest grid and then they interpolate back to a fine grid. The path of return to fine grid is different for the V-cycle and the W-cycle, as illustrated in the figure.

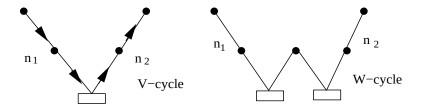


Figure 7.37: Schematic of typical V-cycle and W-cycle multigrid algorithms.

For the *V-cycle*, in particular, we provide all the steps of the algorithm:

- Perform n_1 relation sweeps with smoother S_1 on solution Θ_h^n $(n_1 \sim 2-4)$.
- Perform coarse-grid correction, $\Theta_h^{n+1} = \Theta_h^n + \delta \Theta_h$.
- Perform n_2 relaxation sweeps with smoother S_2 on solution $\Theta_h^{n+1}(n_2 \sim 2-4)$.

The total amplification factor is then:

$$\mathbf{G}_{h,H} = \mathbf{G}_{S_2}^{n_2} \cdot \mathbf{G}_{CGC} \cdot \mathbf{G}_{S_1}^{n_1}$$
.

Remark 4: In the above we have implemented the V-cycle for the correction scheme; it transfers the residual between grids. A similar algorithm can be obtained using nested iteration; it transfers the linear system from one grid to the other, starting from the coarse grid solution. This is the so called *Full Multigrid cycle*; for more details, the reader is referred to [11, 10, 92].

Convergence and Computational Cost

The question of computational complexity of the multigrid method is related to the convergence rate of the method. The storage requirements are relatively simple to estimate. They are less than *twice* the storage required on the fine grid in either one-dimensional or multi-dimensional problems.

The convergence rate of multigrid is defined by the spectral radius of the *total amplifica*tion matrix which is less than one, i.e.,

$$\rho(\mathbf{G}_{h,H}) < 1.$$

Asymptotically, the multigrid algorithm is particularly effective for large-scale problems, since ρ is (in most cases) independent of the spacing h or equivalently the number of grid points N. The major cost is due to smoother as the intergrid transfers cost about 15% of the entire cycle.

We can provide an estimate for the work required for a V-cycle assuming that we only perform one relaxation sweep per grid that costs δt_r . Assuming also that we employ m coarse grids, and since we visit twice each grid, then the total work is

$$W_V = 2[1 + 2^{-d} + 2^{-2d} + \dots + 2^{-md}]\delta t_r = \frac{2}{1 - 2^{-d}}\delta t_r$$

where d denotes space dimension. For example, in three-dimensions (d=3) we have that the work for the entire V-cycle, W_V , is slightly more than twice the work (time) required for one relaxation sweep, δt_r .

Next, we can estimate the number of required sweeps per level n_1 and n_2 . For simplicity, we assume that $n_1 = n_2 = n$ and that the spectral radius $\rho(\mathbf{G}_{h,H})$ is indeed independent of h. The multigrid V-cycle should reduce the algebraic error from $\mathcal{O}(1)$ to (at least) the level of the discretization error $\mathcal{O}(h^p) = \mathcal{O}(N^{-p})$, where p is the order of the finite difference discretization accuracy, e.g. p = 2. This condition implies that

$$\rho(\mathbf{G}_{h,H})^n = \mathcal{O}(N^{-p}) \Rightarrow n \approx \mathcal{O}(\log N).$$

The total cost of the V-cycle is then

$$W_V \approx n \times \mathcal{O}(N^d) \approx \mathcal{O}(N^d \log N)$$

where $\mathcal{O}(N^d)$ is the minimum number of operations for the N^d grid points involved in d space dimensions. In most problems the value of $n \sim \mathcal{O}(1)$, for example n = 4 and thus,

for two-dimensions the optimum cost is $\mathcal{O}(N^2)$ versus $\mathcal{O}(N^3)$ for direct solvers and $\mathcal{O}(N^4)$ for the Jacobi or the Gauss-Seidel solvers. These estimates assume an $N \times N$ grid. For preconditioned conjugate gradients (see section 9.4) the work is $\mathcal{O}(N^{5/2})$. Therefore, the multigrid is indeed a very efficient method!

Remark 5: We have presented here only the main elements of multigrid for linear systems. For more in-depth study of multigrid including applications to nonlinear systems the reader should consult the references [11, 92, 10]. A very good online resource is the site **www.MGNet.org**.

Remark 6: The computational efficiency of the multigrid method depends critically on the parallel implementation of the smoother and the solution of the system on the coarse grid. Modern implementations of multigrid exploit the memory hierarchies of the processors and in particular the L2 and L1 cache, see [34]. In terms of available parallel multigrid software, the free C/MPI package ParMGridGen from the University of Minnesota generates an optimum sequence of successive coarse grids which are suitable for the multigrid algorithm.

7.3 Homework Problems

1. Use von Neumann analysis to show that leap-frog time differencing combined with central differencing to discretize

$$\frac{\partial u}{\partial t} = \frac{\partial^2 u}{\partial x^2}$$

leads to an unstable scheme. Verify your results with heuristic arguments using the corresponding equivalent differential equation.

2. Write a C++ code to implement the DuFort-Frankel scheme for solving

$$\frac{\partial u}{\partial t} = \frac{\partial^2 u}{\partial x^2}, \quad x \in [-1, 1]$$

with

$$u(-1,t) = 0;$$
 $u(1,t) = 1$ and $u(x,0) = -(x+1)/2.$

Consider two grids with number of points N=20 and N=40, and advance the solution to time t=1:

- (a) First, using a constant diffusion number D, and
- (b) Second, using a timestep Δt that scales linearly with Δx .

What do you observe?

3. Show that the eigenvalues of the one-dimensional diffusion matrix obtained from secondorder central differencing with Dirichlet boundary conditions are:

$$\lambda_j = -\frac{4}{\Delta x^2} \sin^2 \left(\frac{j\pi}{2(N+1)} \right) ,$$

and the corresponding eigenvectors are

$$\phi_j = \left[\sin(\frac{j\pi}{2(N+1)}), \sin(\frac{2j\pi}{2(N+1)}), \dots, \sin(\frac{Nj\pi}{2(N+1)})\right]^T.$$

4. Show that the eigenvalues of the partial differential equation

$$\frac{\partial^2 u}{\partial x^2} + \delta^2 \frac{\partial^2 u}{\partial y^2} = \lambda u$$

with homogeneous Dirichlet boundary conditions are given by

$$-4\sin^2(\frac{i\pi}{2(N+1)}) - 4\delta^2\sin^2(\frac{j\pi}{2(N+1)})$$
 with $i, j = 1, \dots, N$.

We also assume that the discretization is second-order central differencing and that $\Delta x = \Delta y = 1$.

5. Consider the diffusion equation with the "wrong sign" (negative diffusion)

$$\frac{\partial u}{\partial t} = -\frac{\partial^2 u}{\partial x^2}$$

with

$$u(-1,t) = u(1,t) = 0$$
 and $u(0,0) = 1$.

- (a) What is the solution u(x,t) after time t=0.1;1 and also $t\to\infty$.
- (b) Use explicit time-stepping to advance the solution and observe a numerical instability developing. Is the physical solution stable?
- 6. Solve the nonlinear parabolic equation

$$\frac{\partial u}{\partial t} + u + u^3 = \frac{\partial^2 u}{\partial x^2} + \cos(\pi x)$$

with

$$u(0,t) = u(t,1) = 0$$
 and $u(x,0) = \sin(\pi x)$.

- (a) First, using an explicit method to obtain empirically an equivalent diffusion number for stability.
- (b) Second, implicitly using also Newton's method for the nonlinear equation.
- 7. Solve the one-dimensional heat equation

$$\frac{d^2T}{ds^2} = \sin 2\pi s, \quad s \in [0, 1]$$

with periodic boundary conditions using

- (a) The Thomas algorithm.
- (b) The serial Jacobi method.
- (c) The parallel Jacobi method.

Use more than one grid and compare convergence rates as well as computational times.

8. Consider the Helmholtz equation

$$\frac{\partial^2 u}{\partial^2 x} + \frac{\partial^2 u}{\partial y^2} - \lambda^2 u = 0$$

where $(x, y) \in [-1, 1] \times [-1, 1]$ and

$$u(-1, y) = 0$$
; $u(1, y) = 0$; $u(x, -1) = 0$; $u(x, 1) = 1$.

Discretize this equation using central differencing and solve the corresponding algebraic system using the Jacobi and SOR methods. Experiment with $N_x = N_y = 10$, 50, 100 grid points.

Is convergence of these iterative solvers guaranteed for all values of λ^2 ?

What is the optimum value of the relaxation parameter ω ?

- 9. Consider the 2×2 matrix with a constant value a in the main diagonal and b and -b in the cross-diagonal. Determine the relationship between a and b so that the Jacobi, Gauss-Seidel, and SOR methods are guaranteed to converge.
- 10. Discretize the equation $\nabla^2 u = f$ with the exact solution

$$u(x,y) = \cos(\pi x)\sin(\pi y), \quad (x,y) \in [-1,1] \times [-1,1]$$

using the fourth-order compact (Padè) scheme. Use the ADI and Peaceman-Rachford methods to obtain the numerical solutions and determine the convergence rate of the solver as well as the spatial accuracy of the discretization.

What do you observe?

- 11. Show that in the relaxed Jacobi algorithm, when ω is chosen to damp the smooth modes, then the rough modes are amplified.
- 12. Write a C++ code for the one-dimensional relaxed Jacobi and SOR. Then experiment with different initial modes and compare against the Laplacian modes.
- 13. Show that the prolongation operator (with linear interpolation) and the restriction operator (with full-weighting) satisfy the transpose property, i.e.,

$$\mathbf{I}_H^h = C(\mathbf{I}_h^H)^T$$

where C is a scaling constant.

14. Implement the multigrid V-cycle in two dimensions for the unsteady diffusion equation using second-order central differencing. Consider first a square domain and subsequently a rectangle with aspect ratio 25. Use a relaxed Jacobi for smoother and full-weighting for the restriction operator.

What is the convergence rate for the square domain and what is it for the large aspect ratio domain?

15. Implement an MPI version of the SOR algorithm. Solve the one-dimensional heat equation

$$\frac{d^2T}{ds^2} = \sin 2\pi s, \ \ s \in [0, 1]$$

with periodic boundary conditions using:

- (a) The relaxation parameter chosen so that the method becomes a Gauss-Seidel solver.
- (b) Using an optimal relaxation parameter.

Use more than one grid and compare convergence rates as well as computational times. Compare computational timing results (using MPI_Wtime) for these two methods against parallel Jacobi for P = 2, 4, and 8 processes.

- 16. Consider $\nabla^2 u = A \sin(2\pi x) \sin(2\pi y)$ on the unit square. Solve for A = 1.0, 2.0, 3.0, 5.0 and A = 10.0 using:
 - (a) The serial Jacobi method.
 - (b) The serial SOR method.
 - (c) The parallel Jacobi method.

Use more than one grid and compare convergence rates as well as computational times. If you fix the number of grid points per direction to 20 and examine the number of iterations that each methods takes to converge, does this number of iterations vary as you change the value of A?

Chapter 8

Propagation: Numerical Diffusion and Dispersion

In this chapter we continue with mixed discretizations for initial value problems (IVP) and boundary value problems (BVP), but our emphasis is on advection equations and wave propagation with and without dissipation. Specifically, we introduce two important non-dimensional numbers that define the accuracy and the stability of discretizations we present: The *Courant number* or CFL condition first proposed by Courant in 1928, and the (grid) *Peclet number* for advection-diffusion systems. We also provide C++ functions and corresponding results that illustrate the effects of numerical diffusion and numerical dispersion in pure advection and in advection-diffusion systems.

On the parallel computing side, we introduce the concept of non-blocking communications, specifically the use of MPI_Isend and MPI_Irecv . Non-blocking communications may lead to a reduction in the computational time by allowing the programmer to appropriately intertwine computation and communication.

8.1 Advection Equation

We consider as a prototype problem the linearized wave or advection equation

$$\frac{\partial\Theta}{\partial t} + U\frac{\partial\Theta}{\partial x} = 0 \quad \left(\text{or } \frac{D\Theta}{Dt} = 0\right),\tag{8.1}$$

which expresses the (passive) advection of heat or a species in one-dimension by the flow with transport velocity U(x,t). For a constant U this is a deceptively "simple-looking" equation, but it is perhaps one of the most difficult equations to resolve in scientific computing. The reason is that, as we will see below, it exhibits no physical diffusion or dispersion. However, there is no practical numerical method for the solution of equation (8.1) that does not introduce at least some amount of either numerical diffusion or dispersion! This equation is therefore a good yardstick by which we can measure these two fundamental properties of a numerical method.

$$x = 0$$
 $x = 1$

Figure 8.1: Domain for the one-dimensional linear advection equation.

Let us assume that we have periodic boundary conditions, i.e., $\Theta(0,t) = \Theta(1,t)$, and that the initial conditions are $\Theta(x,0) = \Theta_0(x) = \sin 2\pi x$, then the exact solution is $\Theta = \Theta_0(x-Ut) = \sin 2\pi (x-Ut)$, which is a traveling wave of constant amplitude that propagates to the right in figure 8.1, with constant velocity U. The exact solution can be verified by direct substitution in equation (8.1). Any initial waveform (see figure 8.2) is simply advected to the right without diffusion or dispersion.

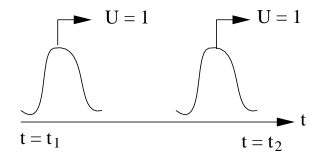


Figure 8.2: A waveform advected with constant velocity to the right is described by equation (8.1).

8.1.1 Dispersion and Diffusion

We explain next the concepts of dispersion and diffusion that a partial differential equation may exhibit as we need to distinguish these from the numerically induced ones. We can use complex arithmetic to write the solution of equation (8.1) as

$$\Theta = \mathcal{R}e\left\{ \left(e^{i2\pi kx} \right) \left(e^{-i2\pi\omega t} \right) \right\}$$
$$= \mathcal{R}e\left\{ e^{i2\pi k(x - \frac{\omega}{k} \cdot t)} \right\}$$

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of constant amplitude; here, ω is the eigenvalue and it is related to the frequency of the wave. The definition of *phase speed* is suggested by the above expression

$$C_{\phi} = \omega/k$$
,

and for the linear advection equation it is obtained upon substitution of the assumed solution into equation (8.1), i.e.,

$$\frac{\partial \Theta}{\partial t} = -i2\pi\omega\Theta$$

$$-U\frac{\partial \Theta}{\partial x} = -2\pi ki\Theta U$$

$$\Rightarrow \omega = kU \Rightarrow \frac{\omega}{k} = C_{\phi} = U.$$

Thus, the waveforms are not dispersed as time progresses, as the phase speed is equal to the constant transport velocity U; they are also non-dissipative, and thus we obtain solutions as sketched in figure 8.2.

In contrast, in the diffusion equation

$$\frac{\partial\Theta}{\partial t} = \frac{\partial^2\Theta}{\partial x^2},\tag{8.2}$$

and assuming a solution of the form

$$\Theta = \mathcal{R}e\{e^{\sigma t}e^{i\pi 2kx}\},\,$$

where σ is the eigenvalue, we obtain upon substitution in equation (8.2)

$$\frac{\partial \Theta}{\partial t} = \sigma \Theta = -(2\pi k)^2 \Theta = \frac{\partial \Theta}{\partial x^2}.$$

Therefore, we have dissipation of the waveform, and σ is the attenuation factor, i.e., the solution $\Theta(x,t)$ is dissipative.

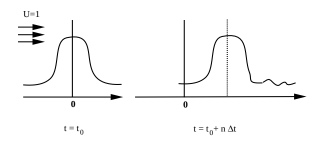


Figure 8.3: Dispersion leads to short waves leading the wave form.

Similarly, we consider a third-order equation

$$\frac{\partial\Theta}{\partial t} = \frac{\partial^3\Theta}{\partial x^3} \tag{8.3}$$

and assuming a solution of the form

$$\Theta = \mathcal{R}e\{e^{i2\pi kx}e^{-i2\pi\omega t}\}.$$

Upon substitution in equation (8.3), we obtain

$$\frac{\partial \Theta}{\partial t} = -i2\pi\omega\Theta = -i(2\pi k)^3\Theta = \frac{\partial^3 \Theta}{\partial x^3}$$

and therefore

$$\frac{\omega}{k} = C_{\phi} = (2\pi k)^2.$$

This is the dispersion equation for the third-order partial differential equation with periodic boundary conditions. It shows that the phase speed depends on the wave number, with short waves traveling quadratically faster. Each wave corresponding to wave number k travels at a speed proportional to the square of its wavenumber, and we obtain solutions which graphically look like the sketch in figure 8.3.

In summary, we have seen that the order of spatial derivative determines if a PDE exhibits dispersion or diffusion. Specifically,

- (i) $\Theta_t = -\Theta_x$ leads to non-dissipative, non-dispersive solutions.
- (ii) $\Theta_t = \Theta_{xx}$ leads to dissipative solutions.
- (iii) $\Theta_t = \Theta_{xxx}$ leads to non-dissipative but dispersive solutions.
- (iv) $\Theta_t = -\Theta_{xxxx}$ leads to dissipative solutions (similar justification as above).

Therefore, the *odd derivatives* (higher than first) are associated with dispersion, whereas the *even derivatives* are associated with dissipation. We will use this fact in the following in order to determine the numerical properties of a specific discretization by examining its corresponding *equivalent differential equation*.

8.1.2 Other Advection Equations

Burgers Equation

The nonlinear advection equation has the form

$$\frac{\partial\Theta}{\partial t} + \frac{1}{2}\Theta\frac{\partial\Theta}{\partial x} = 0. \tag{8.4}$$

It has been introduced by the fluid mechanist J.M. Burgers [12] to model quadratic nonlinearities that appear in many physical phenomena including turbulence and shock waves. In particular, the initial conditions are very important in determining the structure of solutions to this *inviscid* Burgers equation. The basic discretizations employed for the solution of equation (8.1) can also be used to solve equation (8.4) with appropriate treatment of the nonlinear terms.

Second-Order Wave Equation

The advection equation (8.1) is first-order and admits one-directional wave solutions. Its second-order analogue has the form

$$\frac{\partial^2 \Theta}{\partial t^2} = c^2 \frac{\partial^2 \Theta}{\partial x^2} \,, \tag{8.5}$$

and admits left- and right-traveling waves with velocity c. This equation can be reduced to a system of first-order linear advection equations, similar to equation (8.1), as follows

$$\frac{\partial v}{\partial t} = c \frac{\partial \Theta}{\partial x}; \quad \frac{\partial \Theta}{\partial t} = c \frac{\partial v}{\partial x}.$$

Stochastic Advection Equation: We assume that the transport velocity is a stochastic variable that can be decomposed into its mean and a fluctuation component as follows $U = \bar{U} + u$. If the fluctuation component follows a Gaussian distribution with variance σ^2 , its probability distribution function has the form

$$f(u) = \frac{1}{\sqrt{2\pi\sigma^2}} exp[-\frac{u^2}{2\sigma^2}].$$

An exact expression can be obtained for the *mean solution*, i.e.,

$$\bar{\Theta} = \Theta_0(x - \bar{U}t)exp[-\pi^2 \alpha t].$$

Here, $\Theta_0(x)$ is the initial (deterministic) condition, and α is an effective diffusivity due to random fluctuations given by

$$\alpha = \frac{\sigma^2 \tau}{2} \,,$$

where τ is a time scale that determines the sampling rate. For example, if the initial distribution u is not updated during the time integration, then $\tau = t$, but if the fluctuation is updated every, say, 10 time steps, then $\tau = 10\Delta t$. Note that numerical discretizations of equation (8.1) can be employed for different distributions of u in order to obtain a statistically converged solution, which will be an approximation to the above solution. This is the so-called Monte Carlo method, the efficiency of which depends on the efficiency of numerical solutions for the deterministic equation (8.1).

In the following, we consider first- and second-order discretizations of the linear advection equations. For the other type of advection equations the same scheme apply.

8.1.3 First-Order Discrete Schemes

Most explicit schemes for the advection equation are conditionally stable and depend on a non-dimensional number, i.e., the Courant or CFL number. In the following, we first introduce some first-order schemes, which are useful for short-term integration only, and subsequently we present second-order schemes that are suitable for both short-term and long-term time integration.

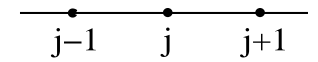


Figure 8.4: The EF/CD schemes employs a three-point stencil.

Euler-Forward/Center-Difference Scheme (EF/CD)

This is a straight-forward discretization for equation (8.1) on the stencil of figure 8.4, it is not as useful in practice, but it brings out the subtleties associated with the numerical solution of hyperbolic equations. For simplicity, let us assume that U=1 and we also define the first-order central difference

$$\Delta_x \Theta_j \equiv \frac{\Theta_{j+1} - \Theta_{j-1}}{2\Delta x}.$$

The discrete EF/CD scheme is

$$\frac{\Theta_{j}^{n+1} - \Theta_{j}^{n}}{\Delta t} + \frac{\Theta_{j+1}^{n} - \Theta_{j-1}^{n}}{2\Delta x} = 0, \ j = 1, \dots N$$

with periodic boundary conditions

$$\Theta_0^n = \Theta_N^n$$

and initial conditions given by

$$\Theta_j^0 = \sin 2\pi x_j \,.$$

The total truncation error is obtained by Taylor-expanding all terms to obtain

$$T_j = \frac{\Delta t}{2} \frac{\partial^2 \Theta^n}{\partial t^2} + \mathcal{O}(\Delta t^2, \Delta x^2) = \frac{\Delta t}{2} \cdot \frac{\partial}{\partial x} \left(\frac{\partial \Theta}{\partial x} \right) + \mathcal{O}(\Delta t^2, \Delta x^2).$$

Clearly, this is a consistent scheme of first-order accuracy in time and second-order accuracy in space as expected. The *equivalent differential equation* is obtained by the differential equation (8.1) enhanced with the truncation term, i.e.,

$$\frac{\partial \hat{\Theta}}{\partial t} + \frac{\partial \hat{\Theta}}{\partial x} + T_j = 0 \Rightarrow \frac{\partial \hat{\Theta}}{\partial t} + \frac{\partial \hat{\Theta}}{\partial x} = -\frac{\Delta t}{2} \frac{\partial^2 \hat{\Theta}}{\partial x^2} + \mathcal{O}(\Delta t^2, \Delta x^2).$$

Therefore, we see that the equivalent differential equation introduces a second-order derivative, which according to the previous discussion, indicates numerical dissipation. Moreover, the coefficient is negative, i.e., the numerical diffusivity is negative, which in turn implies exponential growth. Based on this heuristic argument we expect that this scheme is unstable at least for long-term integration.

A more rigorous analysis can be obtained based on the **von Neumann stability** method. Let us assume that

$$\Theta_j^n = \sum_{k=-\infty}^{\infty} a_k^n e^{2\pi i k x_j}$$

and form the difference

$$\Theta_{j+1} - \Theta_{j-1} = \sum a_k e^{2\pi i k x_j} (e^{2\pi i k \Delta x} - e^{-2\pi i k \Delta x})$$
$$= \sum a_k e^{2\pi i k x_j} \cdot 2i \sin 2\pi k \Delta x.$$

Upon substitution in the EF/CD scheme we obtain

$$\sum_{k} (a_{k}^{n+1} - a_{k}^{n}) e^{2\pi i k x_{j}} = -\frac{C}{2} \sum_{k} (2i \sin 2\pi k \Delta x) \cdot a_{k}^{n} e^{2\pi i k x_{j}}$$

where

$$C \equiv \frac{U\Delta t}{\Delta x}.\tag{8.6}$$

This non-dimensional number above is the **Courant number** or **CFL number** or condition, and it was first introduced in the work of Courant, Friedrich & Lewy (1928) [18].

We can now obtain the amplification factor of EF/CD from the magnitudes of the amplitudes

$$a_k^{n+1} = a_k^n (1 - iC\sin 2\pi k\Delta x)$$

or

$$|a_k^{n+1}| = |a_k^n| |1 + C^2 \sin^2 2\pi k \Delta x|^{1/2}.$$

This equation shows that the amplitudes grow in time, and thus the EF/CD scheme is absolutely unstable, irrespective of the value of the Courant number C, i.e., irrespective of the time step!

It is interesting to note, however, that the EF/CD scheme is conditionally stable in the general stability sense. Let us examine the magnitude of the amplitude

$$|a_k^{n+1}| \le |a_k^n| (1+C^2)^{1/2} \le |a_k^n| e^{C^2/2}$$

 $\le |a_k^0| e^{nC^2/2}$

Thus,

$$|a_k^n| \le |a_k^0| e^{\left(\frac{\Delta t}{2\Delta x^2}\right)(n\Delta t)}$$

and this inequality indicates general stability. In practice, this means that for $(n \cdot \Delta t)$ fixed we can get a stable result for:

$$\begin{pmatrix} \frac{\Delta t}{\Delta x^2} < \text{constant} \\ \Delta t, \Delta x \to 0 \end{pmatrix}.$$

This condition is in fact as restrictive as the condition for diffusion analyzed in the previous chapter. For $U \neq 1$ the above condition is equivalent to

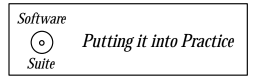
$$D = \frac{U\Delta t}{\Delta x^2} < \mathcal{O}(1),$$

which is a condition on the diffusion number D! Clearly, this is not a practical scheme as the solution blows up after a few time periods as shown in figure 8.5. However, it can be used as a *starter* in multi-step stable high-order explicit schemes.

It is also instructive to examine the stability of this scheme by investigating how the eigenspectrum of the advection operator fits into the stability region of the Euler-forward scheme, which is a unit circle on the left complex plane (see section 5.2). The eigenvalues are obtained from

$$\frac{a_k^{n+1} - a_k^n}{\Delta t} = \underbrace{\left(\frac{-i\sin 2\pi k \Delta x}{\Delta x}\right)}_{\lambda} a_k^n,$$

and are purely imaginary. This means that there is only one point of contact between the stability region of EF and the eigenvalues, and thus the EF/CD is an unstable scheme. A successful scheme, however, would be one that employs leap-frog time integration, (see section 5.2), as its stability region is the imaginary axis. Thus, for $C \leq 1$ the leap-frog/CD scheme is absolutely stable.



Below we present a C++ implementation of the advection equation using central differencing in space and Euler-Forward in time. This function takes as input the number of grid points, the CFL number, and the values at the current time level stored in the array *uold*. This function updates the array *unew* with the new values at the next time level.

Parameter	Value
Method	Euler-Forward Central Difference
Interval	[0, 1]
N	20
Δt	0.005
C	0.1
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Periodic

Table 8.1: Parameters used for the solution shown in figure 8.5.

Key Concept

• Often boundary conditions are handled separately from interior points for looping simplicity.

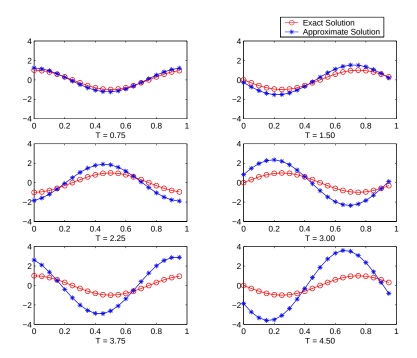


Figure 8.5: Euler-forward/central difference versus exact solution at various times T for the parameters listed in table 8.1.

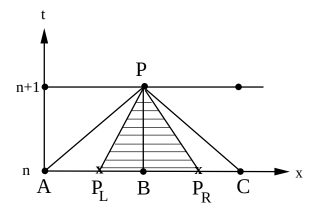


Figure 8.6: Time-space discretization in an explicit scheme. The lines PP_L and PP_R are the characteristics emanating from the point P.

Propagation and the CFL Condition

The Courant number or CFL condition C has a physical meaning and a geometric interpretation that we present next. From an order of magnitude analysis of equation (8.1) we have that

$$\frac{\Delta\Theta}{\Delta t} \propto U \frac{\Delta\Theta}{\Delta x}$$
 or $C = \frac{U\Delta t}{\Delta x} \propto \mathcal{O}(1)$.

For the geometric interpretation we refer to figure 8.6 and make use of the properties of characteristics, i.e. lines along which information is propagated. In our case the characteristic lines are defined by

$$\frac{dX}{dt} = U,$$

and they are straight lines for constant velocity U. In the figure we plot characteristics for both U and -U for the point P. The points P_L and P_R are the locations of point P at time level $(n\Delta t)$ if the wave was advected with velocity U and -U, respectively. By definition the lengths (P_LB) and (P_RB) are equal to the magnitude $U\Delta t$, and the region (PP_LP_R) is the domain of influence of the differential equation. We have already seen that $C \leq \mathcal{O}(1)$ and this simply means that the domain of dependence of the differential equation should be contained within the domain of dependence of the difference equation, the latter defined in figure 8.6 by the region (PAC).

Euler-Forward/Upwind-Differencing Scheme (EF/UD)

The EF/CD scheme is inconsistent with the physics of the problem, which suggests that information is propagating from left to right. However, in the previous scheme we used information from the node (j + 1), which is downstream of node (j). Here, we present a scheme consistent with the physics of the problem and employ an upwind derivative for spatial discretization. The discrete EF/UD scheme is

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} + \frac{\Theta_j^n - \Theta_{j-1}^n}{\Delta x} = 0, \ j = 1, \dots, N$$

which can be re-written as

$$\Theta_{j}^{n+1} = \Theta_{j}^{n} - \frac{C}{2} \left(\Theta_{j+1}^{n} - \Theta_{j-1}^{n} \right) + \frac{C}{2} \left(\Theta_{j+1}^{n} - 2\Theta_{j}^{n} + \Theta_{j-1}^{n} \right).$$

In this form, we can see that the EF/UD scheme is similar to the EF/CD with the addition of an extra term on the right-hand-side. This term is the discrete central difference second-order derivative, and thus it represents *numerical diffusion* added to the difference equation. Correspondingly, the equivalent differential equation is

$$\frac{\partial \hat{\Theta}}{\partial t} + \frac{\partial \hat{\Theta}}{\partial x} = \frac{1}{2} \underbrace{(\Delta x - U \Delta t) \frac{\partial^2 \hat{\Theta}_j}{\partial x^2} + \mathcal{O}(\Delta x^2, \Delta t^2)}_{T_j}.$$

We note that if $\Delta x = \Delta t$ then the Courant number is C= 1 and the above scheme is exact, that is $\Theta_i^{n+1} = \Theta_{i-1}^n$! This is because all high-order terms have the factor $(\Delta x - U\Delta t)$.

With regards to stability, if:

- $C < 1 : U\Delta t \le \Delta x$, and the equivalent differential equation has a positive numerical diffusivity, which in turn implies stability. However, the scheme is of first-order, i.e., $\mathcal{O}(\Delta t, \Delta x)$.
- C > 1: then an instability occurs due to negative diffusivity and associated exponential growth as before.

These results are confirmed from **von Neumann stability** analysis. Specifically, we obtain for the amplitude

$$a_k^n = a_k^n \left[1 - C(1 - e^{-2\pi i k \Delta x}) \right]$$

and taking its magnitude

$$|a_k^{n+1}| = |a_k^n| |1 - C(1 - e^{-2\pi i k \Delta x})|$$

= $|a_k^n| \left[1 + 2(C^2 - C)(1 - \cos 2\pi k \Delta x) \right]^{1/2}$.

For stability, we need to have attenuation of the amplitude, and thus we require

$$2C^2 - 2C \le 0 \Rightarrow C \le 1,$$

which is the same condition for stability as given by the equivalent differential equation.

We can also examine the region of stability in the *complex plane* by identifying the eigenspectrum for the UD scheme, i.e.,

$$\frac{a_k^{n+1} - a_k^n}{\Delta t} = \underbrace{\frac{1 - e^{-2\pi i k \Delta x}}{\Delta x}}_{\lambda \text{ complex}} a_k^n,$$

and thus we obtain

$$\lambda \Delta t = -C(1 - e^{-2\pi i k \Delta x})$$

= $-C(1 - \cos 2\pi k \Delta x) - iC \sin 2\pi k \Delta x.$

In the complex $\lambda \Delta t$ plane, the curves described above are circles enclosed within the unit circle for $C \leq 1$, and thus stability is ensured (see figure 8.7). In contrast to the EF/CD scheme, here the damping due to upwinding brings the $\lambda \Delta t$ into the stability region of the Euler scheme (the C=1 curve).

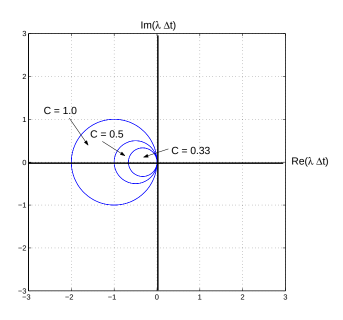


Figure 8.7: Stability region of the EF/UD scheme.

Therefore, with regards to stability the EF/UD is the opposite of the EF/CD, in fact it is too stable! This is suggested by re-examining the equivalent differential equation and write it in the form

$$\frac{\partial \hat{\Theta}}{\partial t} + U \frac{\partial \hat{\Theta}}{\partial t} = \nu_{eff} \frac{\partial^2 \hat{\Theta}}{\partial x^2} + \dots$$

where

$$\nu_{eff} = \frac{U\Delta x}{2}(1-C) \sim U\Delta x$$
.

The numerical solution will be so damped in time that the initial waveform will disappear after some time. This is shown in figure 8.8, where the parameters as of table 8.2 are used, similar to the parameters for the EF/CD method listed in table 8.1.

Below we present a C++ implementation of the advection equation using first-order upwinding in space and Euler-forward in time. This function takes as input the number of grid points, the CFL number, and the values at the current time level stored in the array *uold*. This function updates the array *unew* with the new values at the next time level.

Parameter	Value
Method	Euler-Forward Upwind Difference
Interval	[0, 1]
N	20
Δt	0.005
C	0.1
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Periodic

Table 8.2: Parameters used for the solution shown in figure 8.8.

Programming Note

• A sign mistake can make all the difference. Take for instance this upwind scheme. The sign of the last term (CFL*(uold[i]-uold[i-1])) determines which way the wave is moving. If the sign is correct for upwinding, then the scheme is stable. If not, your code will blow up. Bottom line check your signs carefully.

Remark: Numerical discretization of propagation problems should have the so-called $trans-portive\ property$ according to Roache [78]. This means that the effect of a perturbation is advected only in the direction of the velocity. Therefore, the upwind scheme is transportive but the centered difference scheme is non-transportive. For a variable transport velocity U the following algorithm has been suggested by Roache:

1. First-order:

$$\frac{\partial \Theta}{\partial t} + U \frac{\partial \Theta}{\partial x} \Rightarrow \frac{\Delta \Theta_j}{\Delta t} = -\frac{U_j \Theta_j - U_{j-1} \Theta_{j-1}}{\Delta x}.$$

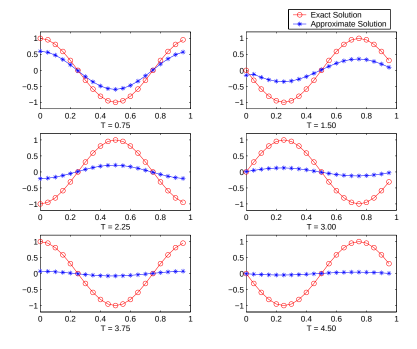
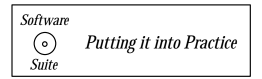


Figure 8.8: Euler-forward/upwind difference versus exact solution at various times T for the parameters listed in table 8.2.



This scheme is transportive but not conservative when U changes sign.

2. Second-order: If the spatial derivative is discretized as

$$-\frac{U_R\Theta_R - U_L\Theta_L}{\Delta x}$$

where

$$\begin{array}{rcl} U_{R} & = & \frac{1}{2}(U_{j+1}+U_{j}); & U_{L} = \frac{1}{2}(U_{j-1}+U_{j}) \\ \Theta_{R} & = & \Theta_{j}, U_{R} > 0 \\ \Theta_{L} & = & \Theta_{j-1}, U_{L} > 0 \end{array}$$

then this scheme is transportive, conservative, and (almost) second-order.

Lax-Friedrichs Scheme (LF) and Tadmor's Correction

The idea of the LF scheme is to stabilize the EF/CD scheme by replacing the contribution from node j, i.e. the value Θ_i^n by its average as shown in figure 8.9. Thus, the discrete

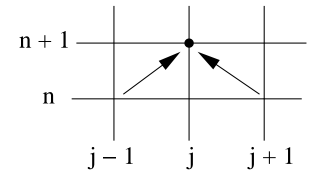


Figure 8.9: Time-space stencil for the Lax-Friedrichs scheme.

scheme is

$$\Theta_j^{n+1} = \underbrace{\frac{1}{2}(\Theta_{j+1}^n + \Theta_{j-1}^n)}_{\text{average for }\Theta_j^n} - \underbrace{\frac{C}{2}(\Theta_{j+1}^n - \Theta_{j-1}^n)}_{}. \tag{8.7}$$

This can be considered as a correction to the *unstable* EF/CD scheme as can be seen by re-writing the above as

$$\Theta_j^{n+1} - \Theta_j^n = \underbrace{\frac{1}{2}(\Theta_{j+1}^n - 2\Theta_j^n + \Theta_{j-1}^n)}_{\text{fixed dissipation}} - \underbrace{\frac{C}{2}(\Theta_{j+1}^n - \Theta_{j-1}^n)}_{\text{optimized}}.$$

The von Neumann analysis shows that

$$a_k^{n+1} = a_k^n [\cos 2\pi k \Delta x - iC \sin 2\pi k \Delta x],$$

and thus stability is obtained for $C \leq 1$. For C = 1 we obtain $|a^{n+1}| = |a^n|$, which implies that there are no dissipation errors. In general, however, this scheme is *overly dissipative*, in fact more dissipative than the first-order upwind scheme. This is shown from the analysis of the *equivalent differential equation*

$$\frac{\partial \hat{\Theta}}{\partial t} + U \frac{\partial \hat{\Theta}}{\partial x} = \frac{\Delta x^2}{2\Delta t} (1 - C^2) \hat{\Theta}_{xx} + \frac{U \Delta x^2}{3} (1 - C^2) \hat{\Theta}_{xxx} + \dots$$

The numerical viscosity is

$$\nu_{eff} = \frac{U}{2C} \Delta x (1 - C^2)$$
$$= \frac{U \Delta x}{2} (1 - C) \frac{(1 + C)}{C}.$$

Therefore, the diffusion of this scheme is higher than the numerical viscosity of UD/CD by the factor $\left(\frac{1+C}{C}\right)$.

The phase errors are measured by

$$\epsilon_{\phi} \equiv \frac{\omega/k}{U}, \quad \phi = 2\pi k \Delta x,$$

the exact value of which is $\epsilon_{\phi} = 1$ for equation (8.1). For C = 1 there are no dispersion errors but in general

$$\epsilon_{\phi} = \frac{\omega/k}{U} \approx 1 + \frac{\phi^2}{3} (1 - C^2) + \mathcal{O}(\Delta x^4),$$

and since $\epsilon_{\phi} > 1$ we have *leading* phase errors.

The Lax-Friedrichs scheme is of first-order, but a *second-order version* has been proposed more recently by Tadmor, see [71]. It employs piecewise linear interpolation instead of constant values within the cell for the standard Lax-Friedrichs scheme. For the nonlinear case the scheme can be written as a predictor-corrector procedure (see homework problems), but for the linear advection equation we consider here, it can be performed in a single step as follows:

$$\Theta_j^{n+1} = \frac{1}{2} [\Theta_{j+1}^n + \Theta_{j-1}^n] + \frac{1 - C^2}{4} [\Theta_{j-1}' - \Theta_{j+1}'] - \frac{C}{2} [\Theta_{j+1}^n - \Theta_{j-1}^n]. \tag{8.8}$$

The Tadmor correction therefore introduces the extra (middle) term in the above equation. Here, the first derivatives are computed properly so that they are of second-order accuracy, but they also satisfy non-oscillatory reconstruction procedures. To this end, we have

$$\Theta'_{j} = MinMod[\Delta\Theta^{n}_{j+1/2}, \Delta\Theta^{n}_{j-1/2}],$$

where we define

$$MinMod[x,y] = \frac{1}{2}[sgn(x) + sgn(y)]Min[|x|, |y|]$$

and also

$$\Delta\Theta_{j+1/2}^n \equiv \Theta_{j+1}^n - \Theta_j^n.$$

The above is just one of several possible ways of computing Θ'_{j} . An alternative way that may lead to crisper discontinuities has also been suggested in [71]; it has the form

$$\Theta'_{j} = MinMod[\alpha \Delta \Theta^{n}_{j+1/2}, \frac{1}{2}(\Theta^{n}_{j+1} - \Theta^{n}_{j-1}), \alpha \Delta \Theta^{n}_{j-1/2}]. \tag{8.9}$$

The free parameter α is chosen so that a steeper slope near discontinuities is obtained. It is in the range $\alpha \in [1, 4)$ in order to satisfy the CFL condition and monotonicity constraints.

Example:



To demonstrate the difference between the Lax-Friedrichs scheme and Tadmor's correction, we have implemented both equation (8.7) and equation (8.8) with Θ'_j defined as in equation (8.9). The implementations can be found in the *software suite*. For this experiment, we will compare the exact solution versus the Lax-Friedrichs (LF) scheme, LF with Tadmor's correction using $\alpha = 1$, and LF with Tadmor's correction using $\alpha = 3$. The parameters used for these experiments are presented in table 8.3 with a parabola as initial condition.

Parameter	Value
Method	Lax-Friedrichs/Tadmor's Correction
Interval	[0, 1]
N	1000
Δt	0.0005
C	0.5
Steps N	10000
Final Time T	5.0
Initial Condition	$u(x,0) = \begin{cases} 100(x-0.4)(0.6-x) & \text{if } 0.4 \le x \le 0.6\\ 0 & \text{otherwise} \end{cases}$
Boundary Conditions	Periodic

Table 8.3: Parameters for the solutions shown in figure 8.10.

In figure 8.10, we present the results of our numerical experiments. The exact solution to this problem is for the initial condition to be advected with speed *one* through the periodic domain. Observe that although the Lax-Friedrichs solution has reasonably maintained the correct phase, the numerical solution is highly dissipated compared to the exact solution. Using Tadmor's correction with $\alpha=1$, we see a significant improvement in the dissipation error. By setting $\alpha=3$, we see slight improvement over the $\alpha=1$ case; however, this improvement is at the cost of the solution being slightly more dispersive. Observe that for $\alpha=3$ we see slight trailing wiggles in the solution indicating the dominance of the dispersion error over the dissipation error.

In the following, we present other high-order schemes suitable for propagation problems.

8.1.4 High-Order Discrete Schemes

Second-Order Upwind Scheme (EF/UD2)

This scheme was introduced by Warming & Beam [91] in order to correct the first-order upwind scheme EF/UD. It has the form

$$\Theta_j^{n+1} = \Theta_j^n - \underbrace{C(\Theta_j^n - \Theta_{j-1}^n)}_{\text{1st-order upwind}} + \frac{1}{2}C(C-1)\underbrace{(\Theta_j^n - 2\Theta_{j-1}^n + \Theta_{j-2}^n)}_{\text{dissipation at }(j-1)}.$$

Therefore, it adds to the upwind node (j-1) positive or negative dissipation depending on the value of C. The corresponding equivalent differential equation

$$\hat{\Theta}_t + U\hat{\Theta}_x = \frac{U}{6}\Delta x^2 (1 - C)(2 - C)\hat{\Theta}_{xxx}$$
$$- \frac{C}{8}\Delta x^3 (1 - C)^2 (2 - C) \left(\frac{\partial^4 \hat{\Theta}}{\partial x^4}\right) + \dots$$

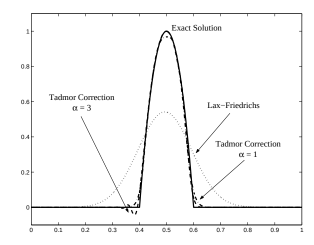


Figure 8.10: Comparison of exact solution (solid), Lax-Friedrichs scheme (dotted), Lax-Friedrichs with Tadmor correction, $\alpha=1$ (dot-dashed), and Lax-Friedrichs with Tadmor correction, $\alpha=3$ (dashed). Details of the numerical experiment are given in the text; the parameters are listed in table 8.3.

shows that there is no second-order derivative and therefore dissipation is reduced compared to first-order upwind.

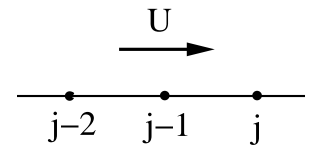


Figure 8.11: Stencil for second-order upwind scheme

The precise stability condition is determined from **von Neumann analysis** as follows:

$$a_k^{n+1} = a_k^n \left\{ 1 - 2C[1 - (1-C)\cos\phi]\sin^2\phi/2 - iC\sin\phi[1 + 2(1-C)\sin^2\phi/2] \right\}.$$

and we can obtain

$$\left| \frac{a_k^{n+1}}{a_k^n} \right|^2 = 1 - 4C(1 - C^2)(2 - C)\sin^4 \phi/2.$$

For stability, we require that $C \leq 2$.

Analysis of **phase errors** shows that

$$\epsilon_{\phi} = \frac{\omega/k}{U} \approx 1 + \frac{1}{6}(1 - C)(2 - C)\phi^2 + \mathcal{O}(\phi^4),$$

which suggests that the phase errors are leading errors for C < 1, and lagging error for C > 1. This is demonstrated in figure 8.12; the parameters for the solution are in table 8.4.

Software

O Putting it into Practice
Suite

Below we present a C++ implementation of the advection equation using second-order upwinding in space and Euler-forward in time. This function takes as input the number of grid points, the CFL number and the values at the current time level stored in the array *uned*. This function updates the array *unew* with the new values at the next time level.

Remark: Unlike in previous examples, second-order upwinding, due to its stencil, requires us to handle the first two values (unew[0] and unew[1]) separately instead of the two endpoints of the domain (unew[0] and unew[N-1]).

Lax-Wendroff Scheme (EF/LW)

This is a second-order scheme as well as the previous EF/UD2. In fact, these two schemes are the only single-step schemes that achieve second-order accuracy on a three-point stencil. The key idea is somewhat different: instead of substituting the Taylor expansions of all derivatives in the equation (8.1), we first expand in time the numerical solution and subsequently replace the time derivatives with the first- and second-order wave equations. To wit, we present this

Parameter	Value
Method	Euler-forward Second-Order Upwind Difference
Interval	[0,1]
N	30
Δt	0.01667 and 0.05
C	0.5 and 1.5
T	10.0
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Periodic

Table 8.4: Parameters used for the solutions of figure 8.12.

derivation below. We start with

$$\Theta_j^{n+1} = \Theta_j^n + \Delta t(\Theta_t)_j + \frac{\Delta t^2}{2} (\Theta_{tt})_j^n + \mathcal{O}(\Delta t^3)$$

and replace the first-order time derivative from equation (8.1) and also the second-order time derivative from

$$\Theta_{tt} = U^2 \Theta_{xx}$$
 (second-order wave equation)

thus,

$$\Theta_j^{n+1} = \Theta_j^n - U\Delta t(\Theta_x)_j + \frac{U^2\Delta t^2}{2}(\Theta_{xx})_j + \mathcal{O}(\Delta t^3)$$

or

$$\Theta_j^{n+1} = \Theta_j^n - \frac{C}{2}(\Theta_{j+1}^n - \Theta_{j-1}^n) + \frac{C^2}{2} \underbrace{(\Theta_{j+1}^n - 2\Theta_j^n + \Theta_{j-1}^n)}_{\text{dissipation}}.$$

The last term is a high-order numerical dissipation contribution as it scales with C^2 , i.e., with $(\Delta t)^2$.

The corresponding equivalent differential equation is

$$\hat{\Theta}_t + U\hat{\Theta}_x = -\frac{U}{6}\Delta x^2 (1 - C^2)\hat{\Theta}_{xxx} - \frac{U\Delta x^3}{8} \cdot C(1 - C^2)\hat{\Theta}_{xxxx} + \mathcal{O}(\Delta x^4),$$

which shows predominantly *phase errors* since the dominant term in the truncation error is a third-order derivative.

The von Neumann stability analysis shows that

$$a_k^{n+1} = a_k^n [1 - iC\sin 2\pi k\Delta x - C^2(1 - \cos 2\pi k\Delta x)],$$

and thus for stability we require that C < 1.

The phase errors are *lagging* in this scheme since

$$\epsilon_{\phi} \approx 1 - \frac{1}{6}(1 - C^2)\phi^2 + \mathcal{O}(\phi^4)$$

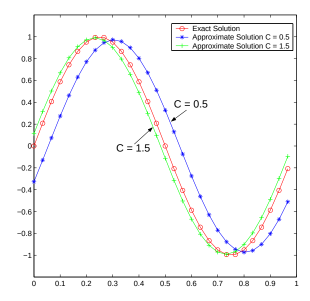
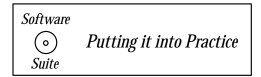


Figure 8.12: Euler-forward/second-order upwind difference versus exact solution at T = 10.0 for C=0.5 and C = 1.5; the parameters are listed in table 8.4.

with the largest phase errors present at high frequencies. This behavior of the Lax-Wendroff scheme is demonstrated in figure 8.13 which is obtained based on the parameters of table 8.5.



Below we present a C++ implementation of the advection equation using the Lax-Wendroff scheme. This function takes as input the number of grid points, the CFL number, and the values at the current time level stored in the array *uold*. This function updates the array *unew* with the new values at the next time level. The following code was used to produce the results shown in figure 8.13.

Parameter	Value
Method	Lax-Wendroff
Interval	[0, 1]
N	30
Δt	0.01667 and 0.0333
C	0.5 and 1.0
T	10.0
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Periodic

Table 8.5: Parameters used for the solutions of figure 8.13.

Remark 1: Grouping terms to reduce operations can increase computational efficiency. Take the example here. Observe that $\operatorname{uold}[i+1]$ is used twice on the right-hand-side expression. The scheme below could be re-written as $\operatorname{unew}[i] = \operatorname{c1*uold}[i] + \operatorname{c2*uold}[i+1] + \operatorname{c3*uold}[i-1]$ * where c1 = 1.0 - CFL * CFL, c2 = CFL(-1.0 + CFL)/2.0 and c3 = CFL(1.0 + CFL)/2.0. Since c1, c2, c3 do not change with respect to i, we can compute them ahead of time and merely use them within the loop. A good compiler will attempt to optimize this for you, but the programmer should always be mindful of such things. The bottom line if you can save the computer time by thinking a little, then you should try thinking a little.

Remark 2: Given that the EF/UD2 and EF/LW schemes have leading and lagging errors, respectively, in the range $0 < C \le 1$, we can construct *hybrid schemes* by combining the two second-order schemes to obtain fourth-order accuracy in the phase properties. This is important especially for long-time integration.

Adams-Bashforth/Center Differencing Scheme (AB/CD)

So far we have considered single-step explicit schemes but higher order schemes can be easily achieved using either multi-stage schemes such as Runge-Kutta or multi-steps schemes such as Adams-Bashforth. Here we present results for the latter; specifically first we present a second-order Adams-Bashforth scheme and subsequently a third-order Adams-Bashforth scheme. The previous stable single-step schemes can be employed in the initial steps.

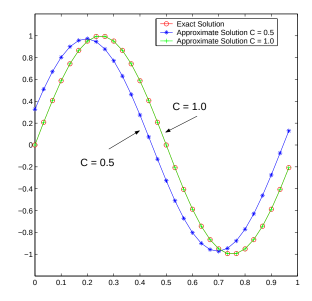


Figure 8.13: Lax-Wendroff versus exact solution at T = 10.0 for C=0.5 and C = 1.0; parameters are listed in table 8.5.

The **second-order** scheme is

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} + \frac{3}{2} \Delta_x \Theta_j^n - \frac{1}{2} \Delta_x \Theta_j^{n-1} = 0,$$

with corresponding equivalent differential equation

$$\frac{\partial \hat{\Theta}}{\partial t} + \frac{\partial \hat{\Theta}}{\partial x} = \left(\frac{5\Delta t^2}{12} - \frac{\Delta x^2}{6}\right) \frac{\partial^3 \hat{\Theta}}{\partial x^3} + \dots$$

The term on the right-hand-side indicates that the dominant errors are phase errors. Note that unlike the previous cases, here the terms Θ_{xxx} , Θ_{vi} , ... are *not* all zero, and thus we have amplitude errors as well but they are subdominant.

The von Neumann stability analysis leads to

$$a_k^{n+1} + a_k^n \left(-1 + \frac{3}{2}iC\sin 2\pi k\Delta x \right) + a_k^{n-1} \left(-\frac{iC}{2}\sin 2\pi k\Delta x \right) = 0.$$

Let us define

$$\mu \equiv \frac{a_k^{n+1}}{a_k^n} \Rightarrow \mu^2 + \mu \left(-1 + \frac{3}{2} i \sin 2\pi k \Delta x \right) + \left(\frac{-iC}{2} \sin 2\pi k \Delta x \right) = 0,$$

where μ is complex. For $C\ll 1$ we can obtain approximate expressions for the two roots: The larger one

$$\mu_{+} = 1 - iC\sin 2\pi k\Delta x - \frac{C^{2}}{2}\sin^{2} 2\pi k\Delta x - \frac{1}{4}iC^{3}\sin^{3} 2\pi k\Delta x - \frac{1}{8}C^{4}\sin^{4} 2\pi k\Delta x + \dots$$

Parameter	Value
Method	2nd order Adams-Bashforth/Central Difference
Interval	[0,1]
N	30
Δt	0.003333
C	0.1
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Periodic

Table 8.6: Parameters used for the solutions of figure 8.14.

corresponds to the physical mode because it is approximately 1, while the smaller root

$$\mu_{-} = -\frac{1}{2}iC\sin 2\pi k\Delta x + \frac{1}{2}C^{2}\sin^{2} 2\pi k\Delta x + \dots$$

is a spurious (numerical) mode.

Note that both roots μ_+, μ_- depend on the initial conditions, and that the numerical mode is damped relatively to the physical mode. For C small we have that $\rightarrow |\mu_-| \ll 1$.

The root corresponding to the physical mode can be bounded from above as shown below by performing leading order analysis:

$$|\mu_{+}| = \left(1 + \frac{1}{2}C^{4}\right)^{1/2} + \mathcal{O}(C^{n})$$

$$|a_{k}^{n}| \approx |a_{k}^{0}|(1 + \frac{1}{2}C^{4})^{n/2}$$

$$\leq |a_{k}^{0}|e^{\frac{C^{4}n}{4}} = |a_{k}^{0}|e^{\frac{\Delta t^{3}(n \cdot \Delta t)}{4\Delta x^{4}}}.$$

The last inequality shows that the AB2/CD is a generally stable scheme. More specifically, the scheme is similar to EF/CD, but here the growth is proportional to C^4 not C^2 and requires only $\Delta t^3 < \Delta x^4$ and not $\Delta t < \Delta x^2$ as in the EF/CD scheme.

Therefore, we see that this is a stable scheme in practice, so we have the case of an unstable scheme, which however is usable for C small. The phase errors for this scheme are shown in figure 8.14, which are large for long time integration.

A better scheme is the third-order Adams-Bashforth/CD scheme

$$\frac{\Theta^{n+1} - \Theta_j^n}{\Delta t} + \frac{23}{12} \Delta_x \Theta_j^n - \frac{16}{12} \Delta_x \Theta_j^{n-1} + \frac{5}{12} \Delta_x \Theta_j^{n-2} = 0.$$

Its corresponding equivalent differential equation is

$$\frac{\partial \hat{\Theta}^2}{\partial t} + \frac{\partial \hat{\Theta}}{\partial x} = \underbrace{\frac{-\Delta x^2}{6} \frac{\partial^3 \hat{\Theta}}{\partial x^3}}_{\text{dispersive}} + \underbrace{\mathcal{O}(\Delta t^3) \frac{\partial^4 \hat{\Theta}}{\partial x^4}}_{\text{diffusive}} + \dots \mathcal{O}(\Delta t \Delta x^2).$$

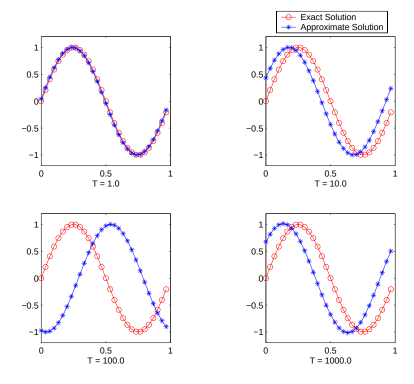


Figure 8.14: Second-order Adams-Bashforth/central difference versus exact solution at various times T; parameters are listed in table 8.6.

Therefore, the phase errors are second-order, and if high-order spatial discretization is used this scheme will exhibit very small dispersion. It is conditionally stable, and the Courant number limit is given by considering the imaginary eigenvalues

$$\lambda \Delta t = -iC \sin 2\pi k \Delta x < 0.723i.$$

For stability, we require that C < 0.723.

In the homework problems we investigate the phase errors for short-time as well as long-time integration.

Crank-Nicolson/Center-Differencing Scheme (CN/CD)

We now study an *implicit* scheme to discretize equation (8.1), the Crank-Nicolson scheme as follows

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} + \frac{1}{2} \Delta_x \Theta_j^{n+1} + \frac{1}{2} \Delta_x \Theta_j^n = 0.$$

The von Neumann stability analysis leads to

$$a_k^{n+1} = \left(\frac{1 - \frac{iC}{2}\sin 2\pi k\Delta x}{1 + \frac{iC}{2}\sin 2\pi k\Delta x}\right) a_k^n \Rightarrow |a_k^{n+1}| = |a_k^n| \quad \forall C, k$$

so we have neutral stability, more specifically absolute but weak stability. The eigenvalues

$$\lambda \Delta t = \frac{-\frac{C^2}{4} \sin^2 2\pi k \Delta x - \frac{iC}{2} \sin 2\pi k \Delta x}{1 - \frac{C^2}{4} \sin^2 2\pi k \Delta x}$$

Parameter	Value
Method	Crank-Nicolson/Central Difference
Interval	[0, 1]
N	30
Δt	0.0333
C	1.0
T	10.0
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Periodic

Table 8.7: Parameters used for the solutions of figure 8.15.

are on the left half-plane for any positive value of C.

Although there are no amplitude errors there are *phase errors*, which are determined from the *equivalent differential equation*

$$\frac{\partial \hat{\Theta}}{\partial t} + \frac{\partial \hat{\Theta}}{\partial x} = \left(-\frac{1}{12}\Delta t^2 - \frac{1}{6}\Delta x^2\right) \frac{\partial^3 \hat{\Theta}}{\partial x^3}$$

where all Θ_{xx} , Θ_{xxxx} , and so on vanish. Let us now assume that

$$\hat{\Theta} = \mathcal{R}e \left\{ e^{2\pi ikx} e^{-2\pi i\omega t} \right\}$$

then we can obtain the approximate dispersion relationship

$$-2\pi i\omega = -2\pi ik - \left(\frac{1}{12}\Delta t^2 + \frac{1}{6}\Delta x^2\right)(-i)(2\pi k)^3,$$

and therefore

$$\epsilon_{\phi} = 1 - \underbrace{\left(\frac{1}{12}\Delta t^2 + \frac{1}{6}\Delta x^2\right)(2\pi k)^2}_{\text{phase error}}.$$

Thus, the *phase errors* are $\Delta \phi = -\left(\frac{1}{12}\Delta t^2 + \frac{1}{6}\Delta x^2\right)(2\pi k)^2$, which lead to errors in period and peak as shown in figure 8.16; the parameters used in the solution are listed in table 8.7.

In general, the largest errors correspond to high wave numbers but for k=N/2 in particular, we have that

$$\Theta \sim Re\left\{e^{2\pi i k \Delta x \cdot j}\right\} = Re\left\{e^{2\pi i \frac{N}{2} \frac{j}{N}}\right\} = Re\left\{e^{\pi i j}\right\}.$$

and thus

$$\frac{\Theta_{j+1} - \Theta_{j-1}}{2\Delta x} = 0.$$

Therefore, this mode never moves although it should be traveling with speed $C_{\phi} = 1!$ Instead, it develops a standing wave with wave length $2\Delta x$ as shown in figure 8.16.

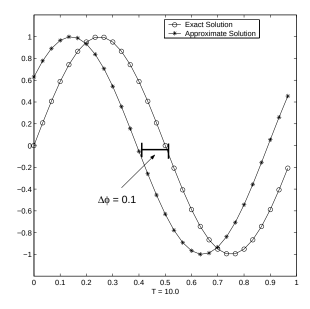


Figure 8.15: Crank-Nicolson/central difference versus the exact solution at T=10.0; parameters are listed in table 8.7.

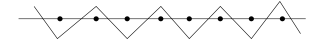


Figure 8.16: Stationary mode develops a 2 Δx wave pattern for the CN/CD scheme.

Below we present a C++ implementation of the advection equation using central differencing in space and Crank-Nicolson in time. This function takes as input the number of grid points, the CFL number and the values at the current time level stored in the array *uold*. This function updates the array *unew* with the new values at the next time level.

Software

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Putting it into Practice

}

Key Concept

• Re-use of code is important. Notice how easily Crank-Nicolson can be implemented once we know that we have our Thomas Algorithm tridiagonal solver?

Remark: The computational complexity of this algorithm is determined from the work required in solving the linear system, i.e. the implicit part, which includes the terms

$$\frac{1}{\Delta t}\Theta_{j}^{n+1} + \frac{1}{4\Delta x}(\Theta_{j+1}^{n+1} - \Theta_{j-1}^{n+1}).$$

This is a triadigonal system and it requires linear work if no pivoting is required. This is guaranteed if we have diagonal dominance, which here means that

$$\frac{1}{4\Delta x} + \frac{1}{4\Delta x} < \frac{1}{\Delta t},$$

and this condition is satisfied if

$$C < 2$$
.

• In other words for stability any value of C leads to a stable scheme for CN/CD, however for linear work we should have C < 2 otherwise pivoting is required which may lead to quadratic or even cubic work!

8.1.5 Effects of Boundary Conditions

So far we have considered periodic boundary conditions, essentially avoiding complications in the discretization due to boundaries. In general, hyperbolic equations like equation (8.1) are very sensitive to boundary conditions, compared, for example, to parabolic equations that we studied in the previous chapter.

To illustrate this effect, let us revisit the linear advection problem but enforce Dirichlet boundary conditions on the left boundary, i.e.,

$$\frac{\partial\Theta}{\partial t} + U \frac{\partial\Theta}{\partial x} = 0; \Theta(x, t = 0) = \sin 2\pi x \tag{8.10}$$

$$0 < x < 1; \quad \Theta(x = 0, t) = -\sin 2\pi t. \tag{8.11}$$

Physically, we have that the left boundary located at x = 0 generates the information that is then propagated through the domain. Note that no boundary conditions are needed at the right boundary (outflow) located at x = 1.

Figure 8.17: An extra node is added at the outflow in order to compute the derivative at the node N.

To proceed, we first consider the **Crank-Nicolson** scheme, which for the periodic case was absolutely stable. The new discretization should include the boundary conditions as follows

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} + \frac{1}{2} \left(\frac{\Theta_{j+1}^{n+1} - \Theta_{j-1}^{n+1}}{2\Delta x} + \frac{\Theta_{j+1}^n - \Theta_{j-1}^n}{2\Delta x} \right) = 0, \ j = 1, \dots, N - 1$$

$$\Theta_0^{n+1} = -\sin 2\pi t^{n+1}; \quad \Theta_j^0 = \sin 2\pi x_j.$$

In the above we have excluded node j = N, as we cannot write the central difference for the first-order spatial derivative. To this end, we add a fictitious node at j = N + 1, see figure 8.17, and obtain Θ_{N+1} by linear extrapolation as follows

$$\Theta_{N+1} = \Theta_N + \Delta x \frac{\Theta_N - \Theta_{N-1}}{\Delta x} = 2\Theta_N - \Theta_{N-1},$$

which can be re-written as

$$\frac{\Theta_{N+1} - \Theta_N}{\Delta x} = \frac{\Theta_N - \Theta_{N-1}}{\Delta x} \,,$$

so at the outflow node we employ a first-order, one-sided derivative. This will affect the accuracy locally, but overall we still obtain $\mathcal{O}(\Delta x^2)$ as errors are advected to the left and are propagated outside the domain. Therefore, at node j = N we have

$$\frac{\Theta_N^{n+1} - \Theta_N^n}{\Delta t} + \frac{1}{2} \left(\frac{\Theta_N^{n+1} - \Theta_{N-1}^{n+1}}{\Delta x} + \frac{\Theta_N^n - \Theta_{N-1}^n}{\Delta x} \right) = 0.$$

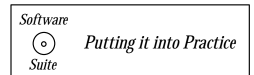
The addition of the upwind derivative does not influence adversely the stability of the CN scheme as it produces eigenvalues in the left half-plane.

In contrast, if we consider **explicit schemes** the outflow condition influences very strongly their stability. For example, the EF/CD scheme which was unstable in the periodic case now becomes stable due to the dissipation of the upwind derivative at this outflow that makes the corresponding eigenvalues land in the left half-plane instead of the imaginary axis as in

the periodic case. Let us examine what happens with the second-order Adams-Bashforth scheme we presented earlier. The discrete equation at the outflow node j = N is

$$\frac{\Theta_N^{n+1} - \Theta_N^n}{\Delta t} + \frac{3}{2} \underbrace{\left(\frac{\Theta_N^n - \Theta_{N-1}^n}{\Delta x}\right) - \frac{1}{2} \left(\frac{\Theta_N^{n-1} - \Theta_{N-1}^{n-1}}{\Delta x}\right)}_{\text{dissipative}} = 0.$$

In the following we first implement the above equation and subsequently we perform numerical experiments to study its behavior.



Below we present a C++ implementation of the advection equation using central differencing in space and 2nd-order Adams-Bashforth in time. This function takes as input the number of grid points, the CFL number, and the values at the current time level stored in the array *uold*. This function updates the array *unew* with the new values at the next time level.

Remark: The function above assumes that the left-hand boundary condition (unew[0]) is handled by the calling function. For the numerical results presented below, the exact solution was used at the inflow. We have obtained numerical results for C = 0.1, which are shown in figure 8.18. There is some small error at the outflow, however we obtain a stable result.

We now turn to the **Leap-frog** scheme which was a stable scheme for the periodic case, but it now becomes unstable due to the effects of dissipation at the outflow node. The scheme

Parameter	Value
Method	2nd order Adams-Bashforth/Central Difference
Interval	[0,1]
N	40
Δt	0.005
C	0.1
T	40.0
Initial Condition	$u(x,0) = \sin(2\pi x)$
Boundary Conditions	Non-periodic

Table 8.8: Parameters used for the solutions of figure 8.18.

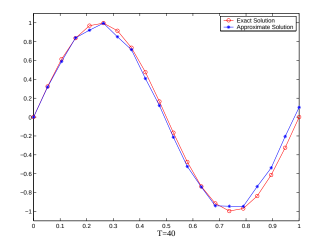


Figure 8.18: Second-order Adams-Bashforth/central difference at T=40.0 for C=0.1, N=20; parameters are listed in table 8.8.

is given by

$$\frac{\Theta_j^{n+1} - \Theta_j^{n-1}}{2\Delta t} + \frac{\Theta_{j+1}^n - \Theta_{j-1}^n}{2\Delta x} = 0, \ j = 1, \dots, N-1$$
$$\frac{\Theta_N^{n+1} - \Theta_N^{n-1}}{2\Delta t} + \frac{\Theta_N^n - \Theta_{N-1}^n}{\Delta x} = 0.$$

This is an unstable scheme as we ask you to verify in the homework problems.

8.2 Advection-Diffusion Equation

We now introduce physical dissipation by including a second-order derivative (in space) to the linearized advection equation (8.1). The initial waveform is both advected and diffused at the same time as shown in figure 8.19.

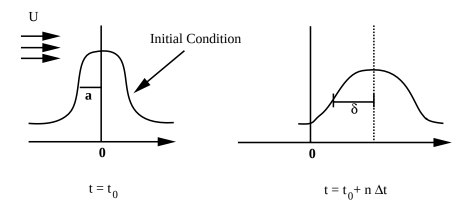


Figure 8.19: Advection and diffusion of an initial waveform by a uniform airstream.

This simple model describes, for example, advection and diffusion of dye by a uniform stream, and has the form

$$\frac{\partial \Theta}{\partial t} + U \frac{\partial \Theta}{\partial x} = \alpha \frac{\partial^2 \Theta}{\partial x^2}, \quad -\infty < x < \infty \tag{8.12}$$

where α is the diffusion coefficient and depends on the material.

The ratio

$$\frac{U\Theta_x}{\alpha\Theta_{xx}} \sim \frac{U\frac{\Delta\Theta}{a}}{\alpha \cdot \frac{\Delta\Theta}{a^2}} \sim \frac{U \cdot a}{\alpha} = Pe$$

is defined as the (physical) Peclet number of the process, and expresses the relative importance of advection compared to diffusion. Here a is the characteristic length scale of the problem.

In practice, we need to truncate the domain to finite limits and thus the interval of interest is

$$-L_I$$
 (inflow) $\leq x \leq L_O$ (outflow).

Later, we will discuss how to choose the size of the truncated domain defined by L_I and L_O , and what are the appropriate boundary conditions to impose there.

8.2.1 Discrete Schemes

We present next representative discretizations of fully implicit, fully explicit, semi-implicit as well as semi-Lagrangian schemes. By this we refer to the time-stepping algorithms involved in handling the advection and diffusion terms, which may or may not be identical for each contribution.

Fully Implicit Discretization

A typical second-order fully implicit scheme is one based on Crank-Nicolson time stepping with central differencing in space. Defining the central discrete first and second derivative operators Δ_x and Δ_{xx} respectively, as before, we have

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} = \frac{1}{2} (\alpha \Delta_{xx} \Theta_j^{n+1} - U \Delta_x \Theta_j^{n+1}) + \frac{1}{2} (\alpha \Delta_{xx} \Theta_j^n - U \Delta_x \Theta_j^n),$$

which can recast in a form that leads more easily to constructing the algebraic system

$$\left(\Delta_{xx} - \frac{U}{\alpha}\Delta_x - \frac{2}{\alpha\Delta t}\right)\Theta_j^{n+1} = -\frac{2}{\alpha\Delta t}\Theta_j^n - \left(\Delta_{xx} - \frac{U}{\alpha}\Delta_x\right)\Theta_j^n.$$

From von Neumann analysis we obtain the amplitude relation

$$a_k^{n+1} = a_k^n \left(\frac{1 - \frac{\lambda \Delta t}{2}}{1 + \frac{\lambda \Delta t}{2}} \right),$$

where the eigenvalues are complex

$$\lambda = -\frac{i \sin 2\pi k \Delta x \cdot U}{\Delta x} - \alpha \underbrace{\frac{\sigma_k}{(1 - \cos 2\pi k \Delta x)}}_{\Delta x^2}.$$

Combining the above two equations, we can conclude that the scheme is stable with no amplitude errors but it suffers from dispersion (phase) errors as we have seen before in the previous section.

Fully Explicit Discretization

Here we use an Euler-forward time stepping algorithm combined with central difference; a higher order scheme for time discretization could be employed in practice, e.g. third-order Adams-Bashforth. However, the issue of stability associated with explicit discretization for an advection-diffusion system are more clearly demonstrated with the first-order scheme

$$\frac{\Theta_j^{n+1} - \Theta_j^n}{\Delta t} = \alpha \Delta_{xx} \Theta_j^n - U \Delta_x \Theta_j^n.$$

This scheme has the following equivalent differential equation

$$\frac{\partial \hat{\Theta}}{\partial t} + U \frac{\partial \hat{\Theta}}{\partial x} = \left(\alpha - \frac{1}{2}\Delta t \cdot U^2\right) \frac{\partial^2 \hat{\Theta}}{\partial x^2} + \ldots + \mathcal{O}(\Delta t^2, \Delta x^2) \cdot \frac{\partial^3 \hat{\Theta}}{\partial x^3}.$$

For stability we require that the modified diffusion coefficient be positive, and thus

$$\frac{1}{2}\Delta t \cdot U^2 \le \alpha \Rightarrow \Delta t < \frac{2\alpha}{U^2}.$$

Remark: For $\alpha = \frac{1}{2}\Delta t \cdot U^2$, we recover the Lax-Wendroff scheme presented in the previous section for the linear advection equation (8.1).

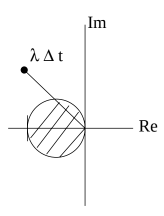


Figure 8.20: Stability diagram and eigenvalues of the EF/CD scheme for the advection-diffusion equation.

A more rigorous von Neuman stability analysis leads to

$$\frac{a_k^{n+1} - a_k^n}{\Delta t} = \lambda a_k^n$$

with complex eigenvalues

$$\lambda = \frac{-i\sin 2\pi k \Delta x U}{\Delta x} - \frac{\sigma_k \alpha}{\Delta x^2},$$

where $\sigma_k = 2(1 - \cos 2\pi k \Delta x)$ as before. It is clear that as $\Delta x \to 0$, the second term (due to diffusion) dominates, and thus we require that

$$\Delta t \le \frac{\Delta x^2}{2\alpha}$$

or

$$\Delta t \lambda_{diff} > -2 \Rightarrow -\frac{\sigma_k \alpha}{\Delta x^2} \cdot \Delta t > -2 \dots,$$

so we recover the limit for the diffusion number, i.e., $D \leq \frac{1}{2}$ as before. We note here that the highest derivative "governs" the stability of the advection-diffusion equation, which is true for other systems as well.

In summary, we have obtained two independent upper limits for the maximum allowable time step in order to guarantee stability of the EF/CD scheme, i.e.,

$$\begin{pmatrix} \Delta t & \leq & \frac{\Delta x^2}{2\alpha} \\ & \text{and} & \\ \Delta t & \leq & \frac{2\alpha}{U^2} \end{pmatrix}.$$

The question then becomes which of the two limits we should use. If we assume that

$$\frac{\Delta x^2}{2\alpha} < \frac{2\alpha}{U^2}$$

then we obtain that

$$Pe_g \equiv \frac{U\Delta x}{\alpha} < 2.$$

The above equation defines the *grid Peclet number*, which should be less than 2 in order to obtain a stable explicit scheme if we only satisfy the diffusion limit.

This statement is *not* equivalent to the previous two conditions, i.e., it is not a necessary condition, but it is sufficiently restrictive. In fact, it imposes a restriction on the mesh size (for stability) independent of the time step Δt , which is somewhat strange! The condition $Pe_g \leq 2$ has been used for a long time in simulations although it does not present the true stability limit for advection-diffusion as we will see next. However, because the numerical solution develops spatial oscillations for $Pe_g > 2$, it was assumed that it was the right stability criterion.

We now re-examine more carefully the stability of the EF/CD scheme for equation (8.12) by following the analysis presented in [56].

If we re-write the amplitude equation from the von Neumann analysis in terms of the Courant (C) and Diffusion (D) numbers, we have

$$a_k^{n+1} = a_k^n [1 + 2D(\cos\phi - 1) - iC\sin\phi].$$

We can now construct the polar plot in the complex plane shown in figure 8.21.

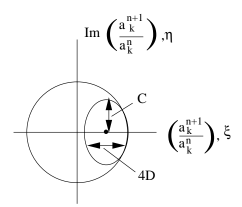


Figure 8.21: Polar plot in the complex plane of the EF/CD scheme.

By inspection of the plot, from first glance we see that stability is obtained if the ellipse is within the unit circle of stability, i.e.,

$$C < 2D < 1$$
.

However, this is not quite correct, and in fact we recover the same condition as before, i.e., the overly restrictive limit on the grid Peclet number! The correct analysis computes exactly the possible intersection points of the ellipse with the circle [56]. We first write the equations for each curve, i.e.,

Circle:
$$\xi^2 + \eta^2 = 1$$

Ellipse:
$$\frac{[\xi - (1 - 2D)]^2}{4D^2} + \frac{\eta^2}{C^2} = 1$$

In order to ensure that we have stability, i.e., $\left|\frac{a_k^{n+1}}{a_k^n}\right| \leq 1$ we have to ensure that there is no intersection next to the point (1,0). To this end, we compute the (hypothetical) second intersection point from

$$\begin{cases} \xi = \frac{(1-4D)C^2 + 4D^2}{C^2 - 4D^2} \\ \eta^2 = 1 - \xi^2 \end{cases}$$

In order to avoid intersection inside the circle and next to the point (1,0), we require that $\xi \geq 1$. This leads to the condition

$$C^2 \le 2D$$
,

which is different than the previous condition, i.e., $C \leq 2D$. Note that we still need to satisfy the diffusion limit, i.e., $2D \leq 1$, and thus finally we have

$$C^2 \le 2D \le 1 \Rightarrow C \le \frac{Pe}{2} \le \frac{1}{C} \text{ or } C \le \frac{2}{Pe} \le \frac{1}{C}.$$

This is the *necessary* and *sufficient* condition for stability. It is equivalent to what we derived heuristically initially using the equivalent differential equation.

Remark: Note that as $\alpha \to 0$ we recover the unconditionally unstable scheme for convection as before. Thus, diffusion (physical here) stabilizes the difference EF/CD equation leading to conditional stability.

Don't Suppress the Wiggles!

We now demonstrate that for $Pe_g > 2$, the numerical solution develops spatial oscillations although we may have a stable scheme, and thus convergence, if the aforementioned two stability criteria are satisfied. To proceed, we need to introduce a more general type of analysis that examines amplification both in time and space. In particular, we can use the time-space normal mode analysis introduced by Godunov & Ryabenkii to analyze the advection-diffusion equation [44]. Similar to having a time amplification factor z, we now have a space amplification factor κ . We write

$$\Theta_i^n = \tilde{\Theta} z^n \kappa^j$$

where the tilde variable denotes amplitude. When introduced into a numerical scheme, including the boundary conditions, we obtain a quadratic equation for κ with two solutions κ_1, κ_2 , i.e.,

$$\Theta_i^n = \tilde{\Theta}z^n (A\kappa_1^j + B\kappa_2^j).$$

The constants A and B are determined from the boundary conditions, and thus we can also investigate the effects of boundary conditions on the stability.

To examine the effect of $Pe_g > 2$ in the advection-diffusion equation, we consider the stationary problem $\left(\frac{\partial}{\partial t} = 0\right)$

$$U\frac{\partial\Theta}{\partial x} = \alpha \frac{\partial^2\Theta}{\partial x^2}, \quad 0 \le x \le L$$

$$\Theta(0) = \Theta_0, \quad x = 0$$

$$\Theta(L) = \Theta_L, x = L.$$

The corresponding difference equation is

$$U\frac{\Theta_{j+1} - \Theta_{j-1}}{2\Delta x} = \alpha \frac{\Theta_{j+1} - 2\Theta_j + \Theta_{j-1}}{\Delta x^2}$$

or

$$(2 - Pe)\Theta_{i+1} - 4\Theta_i + (2 + Pe)\Theta_{i-1} = 0.$$

We now introduce the mode $\Theta_j = \kappa^j$ and obtain the quadratic algebraic equation

$$(2 - Pe)\kappa^2 - 4\kappa + (2 + Pe) = 0$$

with roots

$$\kappa_1 = 1 \text{ and } \kappa_2 = \frac{2 + Pe}{2 - Pe},$$

with the general solution

$$\Theta_j = A\kappa_1^j + B\kappa_2^j$$

or

$$\Theta_j = A + B \left(\frac{2 + Pe}{2 - Pe} \right)^j.$$

From the boundary conditions we determine that

$$A = \frac{\Theta_0 \kappa_2^n - \Theta_L}{\kappa_2^N - 1}$$
 and $B = \frac{\Theta_L - \Theta_0}{\kappa_2^N - 1}$,

where N is the total number of grid points, and thus the solution is

$$\Theta_j = \Theta_0 + (\Theta_L - \Theta_0) \frac{\kappa_2^j - 1}{\kappa_2^N - 1}.$$

We note that for U > 0 then $\kappa_2 < 0$ for Pe > 2. Therefore, the numerical solution will alternate in sign from one grid point to the next producing an oscillatory behavior. If it is stable, the amplitude of the oscillation will be decreasing as $\Delta x \to 0$.

In terms of accuracy, the exact solution is given by

$$\Theta_{ex}(x) = \Theta_0 + (\Theta_L - \Theta_0) \frac{e^{Pe\frac{x}{L}} - 1}{e^{Pe} - 1}$$

and so the numerical solution is a second-order approximation to it.

Semi-Implicit Discretization

Semi-implicit discretizations allow us to treat the advection contribution differently than the diffusion terms. Specifically, since the diffusion limits the time step more severely than the advection, we will treat the former implicitly while we will treat the latter explicitly. To show this, let us integrate equation (8.12) in time with $t \in [t_n, t_{n+1}]$, i.e.,

$$\int_{t_n}^{t_{n+1}} d\Theta + \int_{t_n}^{t_{n+1}} U \frac{\partial \Theta}{\partial x} dt = \int_{t_n}^{t_{n+1}} \alpha \frac{\partial^2 \Theta}{\partial x^2} dt.$$

It is clear that we can approximate the above integrals differently. Here we choose to use Adams-Bashforth of second-order to compute the second integral on the left while we use Euler-backward to compute the integral on the right-hand-side. It is more convenient to do this in two steps by *splitting* this equation, i.e.,

$$(Step I:) \quad \frac{\bar{\Theta}_j^{n+1} - \Theta_j^n}{\Delta t} = -\frac{3}{2}U\Delta_x\Theta_j^n + \frac{1}{2}\Delta_x\Theta_j^{n-1}$$
(8.13)

$$(Step II:) \quad \frac{\Theta_j^{n+1} - \bar{\Theta_j}^{n+1}}{\Delta t} = \alpha \Delta_{xx} \Theta_j^{n+1}$$
(8.14)

The above discretization is $\mathcal{O}(\Delta t^2)$ in the advection but only $\mathcal{O}(\Delta t)$ in the diffusion term, but a Crank-Nicolson scheme can be employed in the second substep, if uniform second-order convergence is desired. However, it is useful to have a high-order (e.g. third-order Adams-Bashforth or fourth-order Runge-Kutta) in the convection scheme to reduce phase errors which are crucial in propagation phenomena, whereas second-order accuracy for amplitude errors may suffice. With regards to stability, a semi-implicit scheme has typically the same stability characteristics as the explicit scheme employed. This is true if the implicit step for diffusion is up to second-order, and thus unconditionally stable. For higher order treatment of the diffusion, however, the time step limit of the semi-implicit scheme is determined as the minimum of the two limits for advection and diffusion. A useful criterion in practice in this case is

$$\Delta t \le \frac{1}{\frac{1}{\Delta t_{conv}} + \frac{1}{\Delta t_{diff}}},$$

which is an empirical rule.

Semi-Lagrangian Discretization

This scheme, which has been used very effectively in metereology simulations, allows the use of a *very large time step* as it bypasses, in effect, the CFL limit by introducing a *variable stencil* defined by the characteristics of wave propagation. It is a semi-implicit scheme with the diffusion treated implicitly but with the advection handled in semi-Lagrangian manner. The Lagrangian form of equation (8.12) in multi-dimensions is

$$\frac{D\Theta}{Dt} = \alpha \nabla^2 \Theta$$

$$\frac{D\boldsymbol{x}}{Dt} = \mathbf{U}(\boldsymbol{x}, t).$$

The idea of a true Lagrangian scheme is to solve the first equation above along the characteristics defined by the second equation. This leads to an effective decoupling of the advection and diffusion terms and an *unconditionally stable* scheme. However, a moving grid is involved, which, if distorted substantially, may lead to great inaccuracies. To this end, re-gridding is typically performed but that may me a substantial overhead to the computation.

The idea of a semi-Lagrangian scheme is to use a fixed grid with the arrival points (along the characteristics) coinciding with the grid points. The departure points are obtained by backwards extrapolation, and they typically do not coincide with the grid points; instead, they are obtained by interpolation. The second-order Crank-Nicolson semi-Lagrangian form for the multi-dimensional advection-diffusion equation is

$$\frac{\Theta^{n+1} - \Theta_d^n}{\Delta t} = \alpha \nabla^2 \left(\frac{\Theta^{n+1} + \Theta^n}{2}\right)$$
$$\frac{D\boldsymbol{x}}{Dt} = \mathbf{U}(\boldsymbol{x}, t), \quad \boldsymbol{x}^{n+1} = \boldsymbol{x}_a.$$

Here Θ_d^n denotes the numerical solution at the departure point \mathbf{x}_d while \mathbf{x}_a is the location of the arrival point, which is a grid point. The characteristic equation is solved backward to obtain \mathbf{x}_d^n given the initial condition $\mathbf{x}^{n+1} = \mathbf{x}_a$.

The *backward integration* could be either explicit or implicit as we show next for secondorder accuracy. First, we use a predictor-corrector method to obtain

$$\hat{\boldsymbol{x}} = \boldsymbol{x}_a - \frac{\Delta t}{2} \mathbf{U}(\boldsymbol{x}_a, t^n)$$

 $\boldsymbol{x}_d = \boldsymbol{x}_a - \Delta t \mathbf{U}(\hat{\boldsymbol{x}}, t^n + \frac{\Delta t}{2}).$

By defining

$$\beta \equiv \boldsymbol{x}_a - \boldsymbol{x}_d$$

we can re-write the above explicit scheme as

$$\beta = \Delta t \mathbf{U}(\boldsymbol{x}_a - \frac{\Delta t}{2} \mathbf{U}(\boldsymbol{x}_a, t^n), t^n + \frac{\Delta t}{2}).$$

If instead we employ implicit integration we obtain

$$\hat{\boldsymbol{x}} = \boldsymbol{x}_a - \frac{\Delta t}{2} \mathbf{U}(\hat{\boldsymbol{x}}, t^n + \frac{\Delta t}{2}),$$

and finally for the difference β we have

$$\beta = \Delta t \mathbf{U}(\boldsymbol{x}_a - \frac{\beta}{2}, t^n + \frac{\Delta t}{2}).$$

Although both the explicit and implicit integration algorithms shown above are of secondorder, better results have been achieved with the implicit scheme, which can be solved using the Newton-Raphson method. This semi-Lagrangian scheme requires extra searching and interpolation algorithms to obtain the departure points and that may be expensive in multi-dimensional computations. However, it is stable for any time step, and it has accuracy

$$\mathcal{O}(\Delta^k + \frac{\Delta x^{p+1}}{\Delta t})$$

where k is the time-integration order (e.g., here k=2) and p is the spatial discretization (e.g., for standard central differencing p=2). For small time steps the second contribution to the overall error typically dominates, but for larger time steps the first term dominates. These two competing contributions create a plateau in the error versus time step, which can be exploited in practice to perform computations with large time step while maintaining high accuracy.

8.2.2 Effects of Boundary Conditions

In advection-diffusion systems the solution is advected while at the same time its amplitude is decreasing. Therefore, the concept of periodic boundary conditions cannot be employed here. Instead, boundary conditions that allow the solution to leave the domain without upstream effects are required. In general, we have *proper* boundary conditions, i.e., mathematically required physical boundary conditions at a physical boundary. However, in a simulation we may also need *artificial* boundary conditions, which are introduced by discretization and the domain truncation but not required by the continuous mathematics or physics. Such boundary conditions are application-specific, but some generalizations can be made. Let us revisit the advection-diffusion problem of advecting a Gaussian waveform to the right in the sketch shown in figure 8.22.

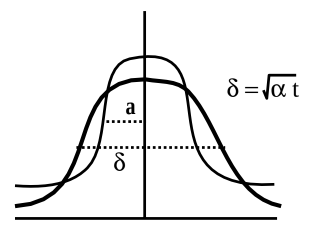


Figure 8.22: Advection-diffusion of a Gaussian waveform. Here a is the initial width and δ is the width after time t.

From an energy balance budget assuming an initial amplitude of unity and width of a, we can obtain the amplitude at time $t = t_f$:

$$\begin{array}{c} \bullet & \text{Initial } \frac{1 \cdot a}{\Theta \cdot \delta} \\ \bullet & \text{Later } \overline{\Theta} \cdot \delta \\ \delta \sim \sqrt{\alpha t} \end{array} \right\} \Rightarrow \overline{\Theta} = \frac{1}{\sqrt{\alpha t}} \, .$$

The waveform is advected in time t by d=Ut. For the mean temperature to decay to $\overline{\Theta}=0.1$ (say) it takes

$$t_f \sim \frac{\delta^2}{\alpha} = \frac{(10a)^2}{\alpha} = 100 \frac{a^2}{\alpha}$$
$$d_f = 100 \frac{Ua^2}{\alpha}$$

and thus

$$\frac{d_f}{a} \sim 100 Pe.$$

From this simple analysis we see that for high Peclet number, which implies faster advection, the required domain should be larger. In general, we first need to obtain such estimates based on conservation laws in order to decide on the extent of the domain.

In addition to selecting the size of the domain, we need to provide boundary conditions at the truncated boundaries consistent with the physics of the problem. However, imposing Dirichlet boundary conditions leads to artificial normal boundary layers that may render the solution erroneous. Instead, weaker boundary conditions that do not affect the solution substantially are preferable, e.g. Neumann boundary conditions or radiation-type boundary conditions, as we explain next.

We revisit the steady-state advection-diffusion problem that we encountered earlier and obtained the following exact solution for *Dirichlet* boundary conditions

$$\Theta_1(x) = \Theta_0 + (\Theta_L - \Theta_0) \frac{e^{Pe\frac{x}{L}} - 1}{e^{Pe} - 1}.$$

Next, we assume that on the right boundary we impose a Neumann boundary condition, i.e.,

$$\frac{\partial \Theta}{\partial x}(x=L) = \Theta_L'$$

with corresponding exact solution

$$\Theta_2(x) = \Theta_0 + e^{Pe} \frac{\Theta_L' L}{Pe} \left[e^{Pe\frac{x}{L}} - 1 \right].$$

By examining the two solutions above, we see that the left (Dirichlet) boundary condition affects the solution independent of Pe. However, for small Peclet number, i.e., Pe << 1, the right boundary condition affects the entire solution, whereas for Pe >> 1, it affects the solution in a region proportional to $\mathcal{O}(L/Pe)$ for the Dirichlet case and only $\mathcal{O}(1/Pe)$ for the Neumann case. In other words, the normal boundary layer developed at the downstream boundary due to truncation of the domain can be minimized at high Peclet number if a Neumann boundary condition is employed there.

For propagation problems, specifically, a radiation type boundary condition is similar to the linearized advection equation, i.e.,

$$\frac{\partial\Theta}{\partial t} + c\frac{\partial\Theta}{\partial x} = 0$$

which leads to $\Theta(x, y, t) = \Theta(x - ct, y)$. In two-dimensional problems it works effectively; it is important, however, to choose the advection velocity c properly. This can be done following, for example, a mass-conservation principle, locally. However, even this boundary condition has been shown to reflect waves especially oblique waves in the x - y plane. To this end, it has been found in [35] that when solving

$$\Theta_{tt} = c^2 [\Theta_{xx} + \Theta_{yy}], \quad t, x \ge 0,$$

where at the x = 0 boundary the boundary condition

$$c\frac{\partial^2\Theta}{\partial x\partial t} - \frac{\partial^2\Theta}{\partial t^2} + \frac{1}{2}c^2\frac{\partial^2\Theta}{\partial y^2} = 0$$

is employed, reflections are substantially reduced. For example, it was reported that the amplitude of the reflections were reduced from 17% to only 3% of a 45 degrees incidence angle wave.

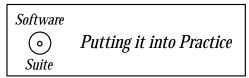
Example: We present the results from the CN/CD discretization presented earlier with Dirichlet and Neumann boundary conditions employed downstream for two different truncations of the domain:

$$\Theta_t + \Theta_x = \frac{1}{100} \Theta_{xx}$$

$$I.C. \quad \Theta(x, t = 0) = e^{-2x^2}$$

$$\Theta(-3, t) = 0; \quad \frac{\partial \Theta}{\partial x}(L_0, t) = 0$$
outflow errors

The discrete parameters are described in tables 8.9 and 8.10, and the corresponding results are shown in figures 8.23 and 8.24 for a short domain and a longer domain. We verify that in the latter case more accurate results are produced as the outflow errors do not influence the results as much.



Below we present a C++ implementation of the advection-diffusion equation using central differencing in space and Crank-Nicolson in time. This function takes as input the number of grid points, the CFL number, the diffusion number, and the values at the current time level stored in the array *uold*. This function updates the array *unew* with the new values at the next time level.

Parameter	Value
Method	Crank-Nicolson/Central Difference
Interval	[-3, 3]
N	31
Δt	0.2
C	1.0
D	0.05
Initial Condition	$\Theta(x,0) = e^{-2x^2}$
Boundary Conditions	$\Theta(-3,t) = 0; \frac{\partial \Theta}{\partial x}(3,t) = 0$

Table 8.9: Parameters for the solutions of figure 8.23.

Remark: Notice the use of '&' in the code above. Recall what this means – the '&' operator gives us "the address of." Hence, in the expression above, we are passing to the ThomasAlgorithm_per routine the *address* of the second entry of the unew array. The following would have been equally valid:

```
ThomasAlgorithm_per(N-1,c2,1.0e0+DN,c1,unew+1,q);
```

which would use pointer arithmetic to pass the address held by unew incremented by one addressable unit, which is the address of unew[1].

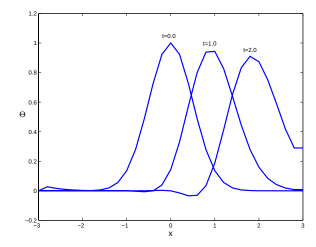


Figure 8.23: Crank-Nicolson/central difference at various times t for the parameters of table 8.9 (short domain).

Parameter	Value
Method	Crank-Nicolson/Central Difference
Interval	[-3, 6]
N	46
Δt	0.2
C	1.0
D	0.05
Initial Condition	$\Theta(x,0) = e^{-2x^2}$
Boundary Conditions	$\Theta(-3,t) = 0; \frac{\partial \Theta}{\partial x}(6,t) = 0$

Table 8.10: Parameters for the solutions of figure 8.24.

8.3 MPI: Non-Blocking Communications

In chapter 5 we presented MPI implementations for finite-differencing similar to the central differencing function presented above. To accomplish the parallel implementation, we used either MPI_Send/MPI_Recv or $MPI_Sendrecv_replace$ commands to exchange information. As was discussed, these functions require some level of synchronization for associating the corresponding sends and receives on the appropriate processes. MPI_Send and MPI_Recv are blocking communications, which means that they will not return until it is safe to modify or use the contents of the send/recv buffer respectively. MPI also provides non-blocking versions of these functions called MPI_Isend and MPI_Irecv , where the "I" stands for immediate. These functions allow a process to post that it wants to send to or receive from a process, and then later is allowed to call a function (MPI_Wait) to complete the sending/receiving. These functions are useful in that they allow the programmer to appropriately stagger computation and communication to minimize the total waiting time due to communication.

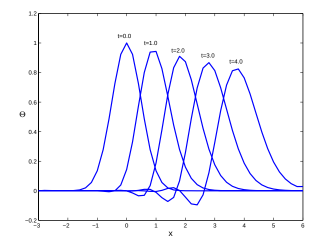


Figure 8.24: Crank-Nicolson/central difference at various times t for the parameters of table 8.10 (long domain).

The basic idea behind MPI_Isend and MPI_Irecv is as follows. Suppose process 0 needs to send information to process 1, but due to the particular algorithms that these two processes are running, the programmer knows that there will be a mismatch in the synchronization of these processes. Process 0 initiates an MPI_Isend to process 1 (posting that it wants to send a message), and then continues to accomplish things which do not require the contents of the buffer to be sent. At the point in the algorithm where process 0 can no longer continue without being guaranteed that the contents of the sending buffer can be modified, process 0 calls MPI_Wait to wait until the transaction is completed. On process 1, a similar situation occurs, with process 1 posting via MPI_Irecv that it is willing to accept a message. When process 1 can no longer continue without having the contents of the receive buffer, it too calls MPI_Wait to wait until the transaction is complete. At the conclusion of the MPI_Wait , the sender may modify the send buffer without compromising the send, and the receiver may use the data contained within the receive buffer; see figure 8.25 for a diagram of this process.

To further expand on these two functions, we will now present the function call syntax, argument list explanation, usage examples and some remarks.

MPI_Isend/MPI_Irecv/MPI_Wait

Function Call Syntax

```
int MPI_Isend(
         void*
                         message
                                        in
         int
                         count
                                        in
         MPI_Datatype
                         datatype
                                        in
                         dest
                                        in
        int
                                        in
                         tag
         MPI_Comm
                         comm
                                        in
         MPI_Request*
                                        out
                         request
int MPI_Irecv(
         void*
                         message
                                        out
         int
                         count
                                        in
         MPI_Datatype
                         datatype
                                        in
                         source
         int
                                        in
         int
                                        in
                         tag
         MPI_Comm
                                        in
                         comm
         MPI_Request*
                         request
                                        out
int MPI_Wait(
         MPI_Request*
                         request
                                      in/out
                                               */)
         MPI_Status*
                                      out
                         status
```

Understanding the Argument Lists

- message starting address of the send/recv buffer.
- *count* number of elements in the send/recv buffer.
- $\bullet \ \ datatype$ data type of the elements in the send buffer.
- $\bullet \ \ source$ process rank to send the data.
- $\bullet \ dest$ process rank to receive the data.
- taq message tag.
- comm communicator.
- request communication request.
- *status* status object.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be sent/recv
int sender; // process number of the sending process
```

```
int receiver; // process number of the receiving process
              // integer message tag
int tag;
                     // variable to contain status information
MPI_Status status;
MPI_Request request; // variable to maintain
                     //
                           isend/irecv information
MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
// Determine datasize
double * databuffer = new double[datasize];
// Fill in sender, receiver, tag on sender/receiver processes,
// and fill in databuffer on the sender process.
if(mynode==sender)
 MPI_Isend(databuffer,datasize,MPI_DOUBLE,receiver,tag,
            MPI_COMM_WORLD,&request);
if(mynode==receiver)
 MPI_Irecv(databuffer,datasize,MPI_DOUBLE,sender,tag,
            MPI_COMM_WORLD,&request);
// The sender/receiver can be accomplishing various things
// which do not involve the databuffer array
MPI_Wait(&request,&status); //synchronize to verify
                            // that data is sent
// Send/Recv complete
```

Remarks

- In general, the *message* array for both the sender and receiver should be of the same type and both of size at least *datasize*.
- In most cases the *sendtype* and *recvtype* are identical.

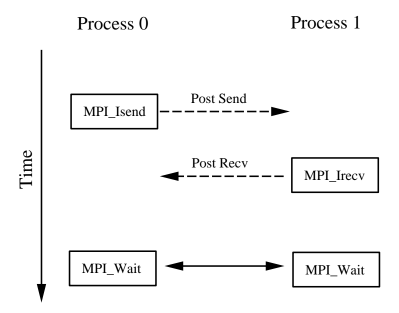


Figure 8.25: MPI_Isend/MPI_Irecv schematic demonstrating the communication between two processes.

- After the MPI_Isend call and before the MPI_Wait call, the contents of message should not be changed.
- After the MPI_Irecv call and before the MPI_Wait call, the contents of message should not be used.
- An MPI_Send can be received by an MPI_Irecv/MPI_Wait.
- An MPI_Recv can obtain information from an MPI_Isend/MPI_Wait.
- The tag can be any integer between 0-32767.
- MPI_Irecv may use for the tag the wildcard MPI_ANY_TAG. This allows an MPI_Irecv to receive from a send using any tag.
- MPI_Isend cannot use the wildcard MPI_ANY_TAG. A specific tag must be specified.
- MPI_Irecv may use for the source the wildcard MPI_ANY_SOURCE. This allows an MPI_Irecv to receive from a send from any source.
- MPI_Isend must specify the process rank of the destination. No wildcard exists.

8.4 Homework Problems

1. Use von Neumann analysis to demonstrate that leap-frog time differencing combined with *central-differencing* of

$$\frac{\partial\Theta}{\partial t} + \frac{\partial\Theta}{\partial x} = 0$$

(with periodic boundary conditions) leads to a conditionally stable scheme. Can you draw the same conclusion from the corresponding equivalent differential equation?

2. Use von Neumann analysis to analyze the stability of leap-frog time differencing combined with *upwind-differencing* of

$$\frac{\partial\Theta}{\partial t} + \frac{\partial\Theta}{\partial x} = 0$$

(with periodic boundary conditions). Compare your findings with heuristic arguments derived from the corresponding differential equation.

- 3. Use a third-order Adams-Bashforth/Central Difference scheme with parameters similar to the ones in table 8.6 but with N=30 as well as N=100 points to obtain solutions to the one-dimensional linear advection equation. Compare your solutions with the solutions of figure 8.14 corresponding to the second-order Adams-Bashforth/Central Difference scheme.
- 4. Use a leap-frog/Central Difference scheme with C=0.2 and N=30 to obtain a solution of the one-dimension advection equation for periodic as well as non-periodic boundary conditions. Are both solutions stable?
- 5. Solve numerically the one-dimensional advection-diffusion equation

$$\Theta_t + \Theta_x = \frac{1}{100} \Theta_{xx}$$

I.C.:
$$\Theta(x, t = 0) = e^{-2x^2}$$

B.C.:
$$\Theta(-3, t) = 0$$

$$\frac{\partial\Theta}{\partial x}(L,t) = 0$$

for
$$L = 3$$
 and $L = 10$.

Use center-differencing for spatial derivatives and Euler explicit in time. You can choose your grid so that

- (a) $\Delta x = 0.01$,
- (b) $\Delta x = 0.1$.

Compare your solutions at time t = 2 and time t = 5.

6. Nonlinear Lax-Friedrichs scheme and Tadmor's correction: In this problem we are interested in solving the inviscid Burgers equation

$$\frac{\partial \Theta}{\partial t} + \frac{\partial F(\Theta)}{\partial x} = 0,$$

where here $F(\Theta) = \frac{1}{2}\Theta^2(x,t)$ and the initial conditions are

$$\Theta(x,0) = \sin(\pi x), \quad x \in [-1,1].$$

The solution of this problem develops a shock discontinuity at time $t_s \approx 0.31$, and we are interested in comparing the numerical solutions obtained with the Lax-Friedrichs scheme and the Tadmor correction before and after the shock is formed at times t = 0.2 and t = 0.4, respectively. Specifically, we want to obtain both the L_1 -norm and the L_2 -norm of errors for grids with N = 50, 100, 200 and 400 points.

The nonlinear versions of the above schemes are given below.

The standard Lax-Friedrichs scheme for the above nonlinear hyperbolic law is:

$$\Theta_j^{n+1} = \frac{1}{2}(\Theta_{j+1}^n + \Theta_{j-1}^n) - \frac{C}{2}[F(\Theta_{j+1}^n) - F(\Theta_{j-1}^n)].$$

The second-order version introduced in [71] consists of a predictor

$$\Theta_j^{n+1/2} = \Theta_j^n - \frac{1}{2}CF_j' \quad (predictor)$$

and a corrector

$$\Theta_j^{n+1} = \frac{1}{2} [\Theta_{j+1}^n + \Theta_{j-1}^n] + \frac{1}{4} [\Theta_{j-1}' - \Theta_{j+1}'] - \frac{C}{2} [F(\Theta_{j+1}^{n+1/2}) - F(\Theta_{j-1}^{n+1/2})].$$

In section 8.1.3 we have defined all quantities except the derivative of the flux which is chosen so that the scheme is free of spurious oscillations. This is given by

$$F_j' = \frac{\partial F}{\partial x}|_j \Theta_j'.$$

We ask you to obtain the convergence rate for each scheme and identify the free parameter α , see section 8.1.3, which gives the best results in the aforementioned norms.

7. Re-write the $EF_FirstOrderUpwind$ function to be a parallel function using MPI_Isend and MPI_Irecv . Verify your code by replicating the numerical example given in the text. Modify your function to solve $u_t + au_x = 0$ where a is a non-zero constant which can be positive or negative. What must change in your MPI code if the direction of the wave propagation changes (i.e., should things change based on whether a is positive or negative)?

- 8. Modify the CrankNicolson_CentralDifference function so that instead of Crank-Nicolson it uses Euler-backward. Repeat the example from table 8.7 for the new Euler-backward routine and compare results with Crank-Nicolson.
- 9. Create a new Crank-Nicolson C++ function which uses a fourth-order compact scheme for the space differentiation (instead of the second-order central difference scheme presented in the text). Repeat the example from table 8.7. How much of a difference does the extra accuracy in space make?
- 10. (Gaussian-cone problem) Consider the advection-diffusion equation with spatially varying advection velocity given by

$$u = +y, \qquad v = -x,$$

and initial condition

$$u(x, y, 0) = e^{-[(x-x_0)^2 + (y-y_0)^2]/2\lambda^2}.$$

The exact solution is

$$u(x, y, t) = \frac{\lambda^2}{\lambda^2 + 2\nu t} e^{-[\hat{x}^2 + \hat{y}^2]/2(\lambda^2 + 2\nu t)},$$

where

$$\hat{x} = x - x_0 \cos t - y_0 \sin t,$$
 $\hat{y} = y + x_0 \sin t - y_0 \cos t.$

The constants are fixed at $\lambda = \frac{1}{8}$; $\nu = 10^{-4}$; and $(x_0, y_0) = (-\frac{1}{2}, 0)$.

Consider second-order central differencing on a grid consisting of 100×100 grid points, and perform time-integration for *one revolution* corresponding to $t = 2\pi$.

- (a) Obtain a numerical solution following the Eulerian approach and Adams-Bashforth/Crank-Nicolson time-stepping (second-order). Use Courant number C=0.01 and diffusion number D=0.5.
- (b) Repeat the above using a semi-Lagrangian approach with backward integration based on the explicit mid-point rule.
- (c) Repeat (b) but with time step Δt ten and twenty times larger, and compare the new results with the results in (b). What do you observe?
- 11. Repeat the previous problem but with a new initial condition, i.e.,

$$u(x, y, t = 0) = \begin{cases} -16 \left[r_0^2 - \frac{1}{16} \right] & \text{if } r_0 < \frac{1}{4} \\ 0 & \text{elsewhere} \end{cases},$$

where $r_0^2 = (x - x_0)^2 + (y - y_0)^2$ and (x_0, y_0) is the initial center position of the cone. The transport velocity field is the same as used in the previous example. The initial condition is a parabolic cone, which has a discontinuity in the derivatives, unlike the previously tested smooth Gaussian cone.

Compare the results from the Eulerian and semi-Lagrangian approaches in terms of contour plots and in terms of the L_2 error. What do you observe?

Chapter 9

Fast Linear Solvers

We have already discussed how to solve tridiagonal linear systems of equations using direct solvers (the Thomas algorithm) in chapter 6 and some iterative solvers (Jacobi, Gauss-Seidel, SOR and multigrid) in chapter 7. We have also discussed solutions of nonlinear and linear systems and have introduced the conjugate gradient method in chapter 4. In the current chapter we revisit this subject and present general algorithms for the *direct* and *iterative* solution of large linear systems. We start with the classical Gaussian elimination (which is a fast solver!) and then proceed with more sophisticated solvers and preconditioners for symmetric and non-symmetric systems.

In parallel computing, we introduce the broadcasting command MPI_Bcast , and demonstrate its usefulness in the context of Gaussian elimination. In addition, we reiterate the use of MPI_Send , MPI_Recv , $MPI_Allgather$, and $MPI_Allreduce$ through example implementations of algorithms presented in this chapter.

Gaussian Elimination 9.1

Gaussian elimination is one of the most effective ways to solve the linear system

$$\mathbf{A}\mathbf{x} = \mathbf{b}$$
.

The Thomas algorithm, see section 6.1.4, is a special case of Gaussian elimination for tridiagonal systems.

The computational complexity of Gaussian elimination is associated with the size and structure of the $n \times n$ matrix A, and so is its accuracy. It is based on the "superposition principle" for linear systems, i.e., the fact that we can replace equations of the original system with equivalent equations formed as linear combinations of the rows of A and corresponding values of **b**. In its simplest form it states:

• Take each row and subtract a multiple of it from subsequent rows in order to zero out the element of **A** below the diagonal.

We demonstrate this by the following example.

Example: Consider the 3×3 system

$$x_1$$
 + $\frac{1}{2}x_2 + \frac{1}{3}x_3 = 3$ (9.1)

$$\boxed{\frac{1}{2}x_1} + \frac{1}{3}x_2 + \frac{1}{4}x_3 = 2 \tag{9.2}$$

$$\boxed{\frac{1}{3}x_1} + \frac{1}{4}x_2 + \frac{1}{5}x_3 = 1 \tag{9.3}$$

We will solve this (3×3) system in two stages of elimination.

Stage 1: In the first stage we target the first term placed in a box in the equation above. To this end, we select the pivot $a_{11} = 1$ and also the multipliers:

$$\ell_{21}^{(1)} = \frac{a_{21}}{a_{11}} = \frac{1/2}{1} = 1/2$$
 and $\ell_{31}^{(1)} = \frac{a_{31}}{a_{11}} = \frac{1}{3}$.

We then multiply equation (9.1) by $\ell_{21}^{(1)}$ and subtract it from equation (9.2). We also multiply equation (9.1) by $\ell_{31}^{(1)}$ and subtract it from equation (9.3). The resulting two new equations replace (9.2) and (9.3), i.e.,

$$x_{1} + \frac{1}{2}x_{2} + \frac{1}{3}x_{3} = 3$$

$$\boxed{\frac{1}{12}x_{2}} + \frac{1}{12}x_{3} = 1/2$$

$$\boxed{\frac{1}{12}x_{2}} + \frac{4}{45}x_{3} = 0$$
(9.4)

$$\boxed{\frac{1}{12}x_2} + \frac{4}{45}x_3 = 0 \tag{9.5}$$

Stage 2: Next, we target the x_2 term, and choose a new pivot and new multipliers, respectively, as:

$$a_{22}^{(1)} = \frac{1}{12}; \qquad \ell_{32}^{(2)} = \frac{1/12}{1/12} = 1,$$

and proceed as before by multiplying equation (9.4) by $\ell_{32}^{(2)}$ and subtracting it from equation (9.5), we obtain

$$x_{1} + \frac{1}{2}x_{2} + \frac{1}{3}x_{3} = 3$$

$$\frac{1}{12}x_{2} + \frac{1}{12}x_{3} = \frac{1}{2}$$

$$\frac{1}{180}x_{3} = -\frac{1}{2}$$
(9.6)

We now see that the system of equations (9.1), (9.6) and (9.7) can be solved easily by back substitution starting from equation (9.7), then equation (9.6) and finally equation (9.1), to obtain:

$$x_3 = -90$$

 $x_2 = 12\left[\frac{1}{2} - \frac{1}{12}(-90)\right] = 96$
 $x_1 = 3 - \frac{1}{2}(96) - \frac{1}{3}(-90) = -15$

In matrix form, the system of equation (9.7) is

$$\begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ 0 & \frac{1}{12} & \frac{1}{12} \\ 0 & 0 & \frac{1}{180} \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 3 \\ \frac{1}{2} \\ -\frac{1}{2} \end{bmatrix}$$

so the coefficient matrix is upper triangular; we will denote this matrix by **U**. We also collect all the multipliers $\ell_{ij}^{(k)}$ we have calculated to form the following lower triangular matrix **L**

$$\mathbf{L} = \begin{bmatrix} 1 & 0 & 0 \\ \frac{1}{2} & 1 & 0 \\ \frac{1}{3} & 1 & 1 \end{bmatrix},$$

where we have places 1's in the diagonal. We can verify that

$$\mathbf{A} = \begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ \frac{1}{2} & \frac{1}{3} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{4} & \frac{1}{5} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ \frac{1}{2} & 1 & 0 \\ \frac{1}{3} & 1 & 1 \end{bmatrix} \begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ 0 & \frac{1}{12} & \frac{1}{12} \\ 0 & 0 & \frac{1}{180} \end{bmatrix} = \mathbf{L} \cdot \mathbf{U}$$

Therefore, the two first stages of Gaussian elimination resulted in the *factorization* of **A** into an **LU** product. Both **L** and **U** are special matrices, and this leads to substantial reduction in computational complexity.

Remark: The matrix **A** employed in this example is a special matrix that has elements $a_{ij} = \frac{1}{i+j-1}$ so the n^{th} row is the vector

$$\left(\frac{1}{n}, \frac{1}{n+1}, \frac{1}{n+2}, \dots, \frac{1}{2n-1}\right)^T$$
.

For n large, e.g., n=1000, the entries of the last row are about three orders of magnitude smaller than the entries of the first row. This large disparity leads to many difficulties because of the *ill-conditioning* of this matrix. This matrix was first introduced by the famous mathematician David Hilbert, and it is called the *Hilbert matrix*; it is an example of an ill-conditioned matrix. Its condition number is large, e.g., greater than 10^5 for $n \ge 5$.

9.1.1 LU Decomposition

We now generalize the Gaussian elimination procedure to an $n \times n$ system

$$a_{11}x_1 + a_{12}x_2 + \dots + a_{1n}x_n = b_1$$

 $a_{21}x_1 + a_{22}x_2 + \dots + a_{2n}x_n = b_2$
 $\vdots \qquad \vdots \qquad \vdots \qquad \vdots$
 $a_{n1}x_1 + a_{n2}x_2 + \dots + a_{nn}x_n = b_n$

In the general case we need (n-1) stages of elimination in order to arrive at the upper triangular system. We will assume that all the pivots at every stage k are $a_{(ii)}^{(k)} \neq 0$, but we will remove this constraint later when we discuss algorithms that involve row and/or column pivoting.

The first stage of elimination leads to

$$a_{11}x_1 + a_{12}x_2 + \dots + a_{1n}x_n = b_1$$

$$a_{22}^{(1)}x_2 + \dots + a_{2n}^{(1)}x_n = b_2^{(1)}$$

$$\vdots \qquad \vdots \qquad \vdots$$

$$a_{n2}^{(1)}x_2 + \dots + a_{nn}^{(1)}x_n = b_n^{(1)}.$$

Here the intermediate coefficients $a_{ij}^{(1)}$ are defined by

$$a_{ij}^{(1)} = a_{ij} - a_{1j}\ell_{i1}^{(1)}; \quad \ell_{i1}^{(1)} = \frac{a_{i1}}{a_{11}}$$

and the entries on the right-hand-side are

$$b_i^{(1)} = b_i - b_1 \frac{a_{i1}}{a_{11}}.$$

Similarly, the second stage of elimination produces $a_{ij}^{(2)}$ and $b_i^{(2)}$, and so on, until the $(n-1)^{th}$ stage, where we obtain $a_{nn}^{(n-1)}$ and $b_n^{(n-1)}$.

This procedure is the forward substitution and gives both matrices \mathbf{L} and \mathbf{U} . In particular, we replace $\mathbf{A}\mathbf{x} = \mathbf{b}$ by

$$\mathbf{L}\underbrace{\mathbf{U}\mathbf{x}}_{\mathbf{y}}=\mathbf{b}\Rightarrow\mathbf{L}\mathbf{y}=\mathbf{b},$$

where

$$\mathbf{U}\mathbf{x} = \mathbf{y}$$
.

We can now summarize the solution procedure, which consists of three main steps, as follows:

- 1. LU decomposition: $\mathbf{A} = \mathbf{L} \cdot \mathbf{U}$
- 2. Forward Solve for y: Ly = b
- 3. Backward Solve for \mathbf{x} : $\mathbf{U}\mathbf{x} = \mathbf{y}$

The pseudo-code for steps (1) and (2) is:

for
$$k = 1, n - 1$$

for $i = k + 1, n$

$$\ell_{ik} = \frac{a_{ik}}{a_{kk}} \text{ (assuming } a_{kk} \neq 0)$$
for $j = k, n$

$$a_{ij} = a_{ij} - \ell_{ik} a_{kj}$$
endfor

$$b_i = b_i - \ell_{ik} b_k$$
endfor
endfor

Computational Cost: The operation count for the above code is obtained by considering first the innermost loop j, then the loop i, and finally adding operations from all elimination stages: k = 1 to n - 1. Thus, we have

$$W_{LU} = 2\sum_{k=1}^{n-1} (n-k) \cdot (n-k) = 2\sum_{m=1}^{n-1} m^2 = 2\frac{(n-1)n(2n-1)}{6} \approx \frac{2}{3}n^3,$$

where the factor 2 accounts for one addition and one multiplication. If only multiplications are counted then

$$W_{LU} \approx \frac{n^3}{3}$$
,

which is the operation count often quoted in the literature.

The third step in the solution is the backward substitution, which yields first

$$x_n = \frac{b_n^{(n-1)}}{a_{nn}^{(n-1)}}$$

and marching backward all the way to the first entry, we have

$$x_1 = \frac{b_1 - a_{12}x_2 \dots - a_{1n}x_n}{a_{11}}.$$

The following pseudo-code represents this back solve:

for
$$k = n, 1$$
 (reverse ordering)
$$x_k = b_k \text{ (initialization)}$$
for $i = k + 1, n$

$$x_k = x_k - a_{ki}x_i$$
endfor
$$x_k = x_k/a_{kk}$$
endfor

We conclude from the above that the operation count for the backward/forward substitution is $\mathcal{O}(n^2)$.

Remark 1: In the pseudo-code above, steps (1) and (2) are accomplished together for computational efficiency. Solving for \mathbf{y} amounts to adding the $(b_i = b_i - \ell_{ik}b_k)$ line in the appropriate place. At the conclusion of this algorithm, the matrix \mathbf{A} has been over-written with the upper triangular matrix \mathbf{U} , and the vector \mathbf{b} has been over-written with the solution of $\mathbf{L}\mathbf{y} = \mathbf{b}$. All that remains is to accomplish the backsolve for $\mathbf{U}\mathbf{x} = \mathbf{y}$.

Remark 2: Note that in both pseudo-codes above we have attempted to minimize the required memory by over-writing onto the same memory locations. However, this should be avoided in cases where we are interested in using the matrix **A** again somewhere else in our program. In the codes above both the entries of matrix **A** and the entries of the right-hand-side are over-written.

Remark 3: We have already discussed the *Thomas algorithm* in section 6.1.4, which is a subcase of the LU decomposition presented here with bandwidth m = 1. In general for a banded matrix with order n and (semi-) bandwidth m the operation count is

- LU decomposition: $\mathcal{O}(m^2n), m \ll n$
- Back Solve: $\mathcal{O}(mn), m \ll n$

Remark 4: The Gram-Schmidt QR factorization of a matrix presented in chapter 2 produces an upper triangular matrix \mathbf{R} but the \mathbf{Q} matrix is an orthogonal *full* matrix. The Gram-Schmidt algorithm costs $\mathcal{O}(2n^3)$ (including addition and multiplications), i.e., it is *three times* more expensive than the LU algorithm. However, the QR decomposition can also be

n = 20:

achieved by the *Householder method* which is only twice as expensive, i.e., it costs $\mathcal{O}(\frac{4}{3}n^3)$, see section 9.3 below.

Remark 5: (Cramer versus Gauss) We compare here the cost for solving $\mathbf{A}\mathbf{x} = \mathbf{b}$ using the Cramer method of determinants (time t_C) which is $\mathcal{O}(n!)$ versus the Gaussian elimination method (time t_G) which is $\mathcal{O}\left(\frac{2n^3}{3}\right)$. We assume that we use a processor with sustained speed of 1 Gflops¹.

$$t_C \approx \frac{3!}{10^9} \text{ seconds } \approx 6 \text{ nanoseconds}$$

$$n = 3:$$

$$t_G \approx \frac{2}{3} \frac{3^3}{10^9} \text{ seconds } \approx 18 \text{ nanoseconds}$$

$$t_C \approx \frac{10!}{10^9} \text{ seconds } \approx 3 \text{ milliseconds}$$

$$n = 10:$$

$$t_G \approx \frac{210^3}{3 \cdot 10^9} \text{ seconds } \approx 0.6 \text{ microseconds}$$

$$t_C \approx \frac{10!}{10^9} \text{ seconds } \approx 675,806 \text{ hours } \approx 28,1585 \text{ days } \approx 80 \text{ years}$$

$$t_G \approx \frac{2}{3} \frac{20^3}{10^9} \text{ seconds } \approx 5 \text{ microseconds}$$

Clearly, Gauss wins by years! In figure 9.1 we plot the growth in computational work of Cramer's method versus the Gaussian elimination method.

Remark 6: The Gaussian elimination offers an efficient way of computing the determinant of **A**, since

$$\det(\mathbf{A}) = \det(\mathbf{L}) \det(\mathbf{U})$$
$$= 1 \cdot [u_{11} \cdot u_{22} \dots u_{nn}]$$

where u_{ii} , i = 1, ..., n are the diagonal elements of the upper triangular matrix **U**. We recall that the determinant of any triangular matrix is simply the product of its diagonal elements.

Remark 7: The Gaussian elimination can be used to explicitly construct the inverse A^{-1} by setting the columns of the identity matrix $I = AA^{-1}$ as

$$\mathbf{b}_i = (0 \dots 0 \underbrace{1}_{index \ i} 0 \dots 0)^T$$

with only the i^{th} entry being non-zero, and solve

$$\mathbf{A}\mathbf{x}_i = \mathbf{b}_i, \ i = 1, \dots, n.$$

The solution vector \mathbf{x}_i forms the column i^{th} of the inverse \mathbf{A}^{-1} . Note that this involves only one LU decomposition and n back solves of $\mathcal{O}(n^2)$ and the total cost is $still\ \mathcal{O}(n^3)$.

¹The TeraHertz processor is already in the horizon; it will consist of about one billion transistors!

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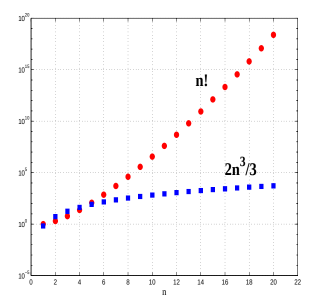


Figure 9.1: Comparison of the growth of computational work in Cramer's method, n!, versus Gaussian elimination method, $\frac{2}{3}n^3$.

9.1.2 To Pivot or Not to Pivot?

So far we have conveniently assumed that all the pivoting elements are non-zero, i.e.,

$$a_{ii}^{(k)} \neq 0$$

but this is not guaranteed for all problems! In practice, these pivots may be zero or very small numbers so that the multiplies $\ell_{ij}^{(k)}$ can potentially be very large numbers. To understand the effect of this, let us consider the 2×2 matrix

$$\mathbf{A} = \left[\begin{array}{cc} \epsilon & 1 \\ 1 & 1 \end{array} \right]$$

where $\epsilon \ll 1$. The condition number of **A** is $\kappa_2(\mathbf{A}) \to 2.6180$ as $\epsilon \to 0$ so this is a well conditioned matrix. We now obtain the LU decomposition of **A**:

The L matrix is

$$\mathbf{L} = \left[\begin{array}{cc} 1 & 0 \\ \epsilon^{-1} & 1 \end{array} \right]$$

and the U matrix is

$$\mathbf{U} = \left[\begin{array}{cc} \epsilon & 1 \\ 0 & 1 - \epsilon^{-1} \end{array} \right] .$$

For ϵ sufficiently small, $1 - \epsilon^{-1} \approx -\epsilon^{-1}$ and thus

$$\mathbf{U} \approx \left[\begin{array}{cc} \epsilon & 1 \\ 0 & -\epsilon^{-1} \end{array} \right],$$

SO

$$\mathbf{L} \cdot \mathbf{U} = \left[\begin{array}{cc} \epsilon & 1 \\ 1 & 0 \end{array} \right] ,$$

which is different than the original matrix A in the (2,2) entry, since

$$\mathbf{A} = \left[\begin{array}{cc} \epsilon & 1 \\ 1 & 1 \end{array} \right] .$$

We note that for any value of $a_{22} \neq 1$ (of order one) we get the same answer, which is obviously wrong! This is an example of a numerical instability. It is due to the fact that the condition number of **L** and **U** is extremely large unlike the condition number of the matrix **A** which is order one. This problem can be avoided if we simply reverse the order of the equations, i.e., interchange the rows, and work with the re-ordered matrix

$$\mathbf{A} = \left[\begin{array}{cc} 1 & 1 \\ \epsilon & 1 \end{array} \right]$$

as now the multiplier is $\epsilon < 1$.

We can generalize the result above and apply row interchange, which is also called *partial* pivoting, to obtain multipliers

$$|\ell_{ij}| < 1$$
.

The following pseudo-code describes Gaussian elimination with partial pivoting. It is an extension of the code we described earlier in the forward solve.

```
for k = 1, n - 1

|a_{mk}| = \max\{|a_{kk}|, |a_{k+1k}|, \dots |a_{nk}|\}

p = m

for q = k, n

c = a_{kq}

a_{kq} = a_{pq}

a_{pq} = c

endfor

for i = k + 1, n

\ell_{ik} = a_{ik}/a_{kk}

for j = k + 1, n

a_{ij} = a_{ij} - \ell_{ik}a_{kj}

endfor

endfor
```

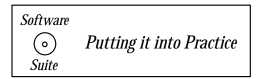
We note that for the system $\mathbf{A}\mathbf{x} = \mathbf{b}$ we also need to interchange appropriately the right-hand-side. Unlike the standard Gauss elimination, in the partial pivoting case we have

$$A \neq LU$$

but $\mathbf{A} = \mathbf{P}^{-1}\mathbf{L}\mathbf{U}$ is true, where \mathbf{P} is a permutation matrix describing the partial pivoting.

While partial (row-pivoting) works effectively in practice, there are a few pathological cases where even this may breakdown. In these cases, we can perform an additional similar pivoting by columns, searching for a maximum pivot along both rows and columns. In general, indications of an ill-conditioned matrix are provided by the small magnitude of the pivot or the large magnitude of the solution compared to the right-hand-side, although there are matrices which do not have these properties but they are still ill-conditioned.

Remark 1: For bounded matrices, (e.g., the tridiagonal systems involved in Thomas algorithms; see section 6.1.4), partial pivoting and complete pivoting result in increasing the bandwidth and even producing *full* matrices. Therefore, the computational work instead of being linear as in the Thomas algorithm may become $\mathcal{O}(n^2)$ or $\mathcal{O}(n^3)$ for row or row/column pivoting, respectively.



The function below is an implementation of both the pivot and non-pivot versions of Gaussian elimination by LU decomposition. Three arguments are required: the matrix \mathbf{A} , the vector \mathbf{b} , and an integer parameter pivotflag denoting whether the pivoting should be enabled (zero for no-pivoting and one for pivoting).

```
void GaussElimination(SCMatrix &A, SCVector &b, int pivotflag){
  int pivot;
  int N = A.Rows();
  /* NOTE: The values contained in both the matrix A and
     the vector b are modified in this routine.
     returning, A contains the upper triangular matrix
     obtained from its LU decomposition, and b contains
     the solution of the system Ax=b*/
  // Steps (1) and (2) (decomposition and solution of Ly = b)
  switch(pivotflag){
  case 1: // Case in which pivoting is employed
    for(int k=0; k < N-1; k++){
      pivot = A.MaxModInColumnindex(k,k);
      A.RowSwap(pivot,k);
      Swap(b(pivot),b(k));
      for(int i=k+1; i<N; i++){
        double l_{ik} = A(i,k)/A(k,k);
        for(int j=k; j<N; j++)
          A(i,j) = A(i,j) - l_ik*A(k,j);
        b(i) = b(i) - l_ik*b(k);
```

```
}
    }
    break;
           // Case O/default in which no pivoting is used
  default:
    for(int k=0; k < N-1; k++){
      for(int i=k+1;i<N;i++){</pre>
        double l_{ik} = A(i,k)/A(k,k);
        for(int j=k; j<N; j++)
          A(i,j) = A(i,j) - l_ik*A(k,j);
        b(i) = b(i) - l_ik*b(k);
      }
   }
  }
  // Step (3) (backsolving to solve Ux=y)
  b(N-1) = b(N-1)/A(N-1,N-1);
  for(int k=N-2; k>=0; k--){
    for(int j=k+1; j<N; j++)</pre>
      b(k) = A(k,j)*b(j);
    b(k) = b(k)/A(k,k);
  }
}
```

Remark 2: In the implementation above we use a *switch* statement to partition the different cases. Two advantages are: code readability and the ease of adding another condition. Suppose that we decided to implement a new pivoting algorithm; within the current function we could merely add another case to denote the new pivoting functionality.

Notice that it is not required that case '0' go before case '1'. Recall that the switch statement evaluates the validity of the switch in the order given in the code, hence in this case the switch statement first checks to see if case '1' is valid, and if not checks for case '0'. Also notice that we use a 'default' case. If neither a zero or a one is given in the switch input, the default will be executed. In this example, the default case and case '0' are identical.

Remark 3: In the code above, we utilize two *Matrix class* methods:

- 1. Matrix::MaxModInColumnindex(...), and
- 2. Matrix::Rowswap(...).

The first method implements the maximum modulus in a column operation needed for pivoting. The second method swaps two rows of the matrix A. The advantage of using these two methods is that from this level in the code the implementation details of how these two functions are accomplished are not our concern. At the level of this code we merely need to know the input, output, and contract for the methods.

Remark 4: Notice that in the code above we both declare and initialize the iterating value (such as int k=0) within the for loops. Recall that C++ does not require us to declare all variables at the beginning of the function. In this function we have made liberal use of this ability by declaring and initializing each iterating value with its respective for loop.

The need for pivoting is dictated by the properties of the matrix \mathbf{A} . There are two special categories of matrices for which we do not need to pivot. These matrices are:

- Diagonally-dominant, or
- Positive-definite.

A strictly diagonally-dominant matrix has the property

$$|a_{ii}| > \sum_{j=1}^{n} |a_{ij}|, i = 1, \dots, n; j \neq i$$

and it is guaranteed to be non-singular.

A positive-definite matrix is defined by the condition

$$\forall \mathbf{x} \neq 0, \ \mathbf{x}^T \mathbf{A} \mathbf{x} > 0,$$

which guarantees that the matrix A is non-singular. In addition, the above properties guarantee that there will be no numerical instabilities in systems where such matrices are involved.

Fortunately, many of the algebraic systems resulting from PDEs that describe physical phenomena (presented in chapters 6 and 7) have these desirable properties. In particular, the matrix obtained from the discretization of d^2u/dx^2 (see chapter 6) is tridiagonal with diagonals (1, -2, 1). Although it does not satisfy strictly the diagonal dominance condition, it is an *irreducible matrix*, i.e., its associated directed graph is strongly connected, and this condition is equivalent to diagonal dominance.

The special matrices we have described are guaranteed to be non-singular but they are also far from being even approximately singular. Clearly, if a matrix \mathbf{A} is singular, then Gaussian elimination cannot be applied as \mathbf{A}^{-1} does not exist. However, in practice many matrices are almost singular, and this is the condition we should investigate, as it leads to numerical instabilities. The value of the determinant, if the matrix is scaled properly, can give us an indication if the matrix is almost singular or ill-conditioned. However, computing determinants is costly and at least equivalent in computational complexity to an LU decomposition, the stability of which we investigate in first place! Only for special matrices the computation of determinant may be employed.

Another approach is to employ the condition number $\kappa(\mathbf{A})$ for matrix \mathbf{A} , i.e.,

$$\kappa(\mathbf{A}) \equiv \parallel \mathbf{A} \parallel \cdot \parallel \mathbf{A}^{-1} \parallel .$$

As we have seen in chapter 2 it relates the perturbation of data to the changes in the solution, i.e.,

$$\frac{\parallel A\mathbf{x} \parallel}{\parallel x \parallel} \le \kappa(\mathbf{A}) \frac{\parallel A\mathbf{b} \parallel}{\parallel b \parallel}.$$

The condition number is always computed with respect to some norm. We will use the notation $\kappa_i(\mathbf{A})$ to denote the condition number of \mathbf{A} with respect to the $\|\cdot\|_i$ norm. When no subscript is given, we assume that the $\|\cdot\|_2$ norm is used. We can define relationships between different norm-based condition numbers using the equivalence relations between the norms.

By definition the condition number $\kappa(\mathbf{A})$ is greater or equal to one, but we are interested in extremely large values of $\kappa(\mathbf{A})$, as it acts as an amplifier in the propagation of disturbance (noise) from the input to output (solution).

For symmetric matrices we have that

$$\kappa_2(\mathbf{A}) = \frac{|\lambda_{\text{max}}|}{|\lambda_{\text{min}}|},$$

and we have computed the eigenvalues λ_i for several cases in chapter 6; see also chapter 10. For general matrices, however, it is difficult to compute the condition number economically so approximate estimation algorithms are employed. Here we will present the estimator proposed by Hager [52]; see also [26]. The algorithm² obtains a lower bound on the inverse matrix in the one-norm. However, we note that \mathbf{A}^{-1} is not constructed explicitly as this is costly, in fact it requires an $\mathcal{O}(n^3)$ operation! Instead, the matrix-vector products $\mathbf{A}^{-1}\mathbf{x}$ are computed on-the-fly:

Initialize:
$$\mathbf{x} : \| \mathbf{x} \|_1 = 1$$

Begin Loop: $\mathbf{y} = \mathbf{A}^{-1}\mathbf{x}$

$$\mathbf{z} = sign(\mathbf{y})$$

$$\mathbf{q} = (\mathbf{A}^{-1})^T\mathbf{z}$$
if $\| \mathbf{q} \|_{\infty} \le \mathbf{q}^T\mathbf{x}$ return $\| \mathbf{y} \|_1$
elseif $\mathbf{x} = \mathbf{e}_j sign(q_j)$ with $|q_j| = \| \mathbf{q} \|_{\infty}$
endif

End Loop

Here $\mathbf{e}_j = (0, 0, \dots, 1, \dots 0)^T$ is the j^{th} column of the identity matrix. The condition number is then estimated from

$$\kappa \approx \parallel \mathbf{A} \parallel_1 \cdot \parallel y \parallel_1$$
.

We note that $\|\mathbf{y}\|_1$ is a local maximum to $\|\mathbf{A}^{-1}\mathbf{x}\|_1$, and that this method is based on computing the gradient of $f(x) \approx \|\mathbf{A}^{-1}\mathbf{x}\|_1$; for a detailed explanation of the algorithm see [52].

Remark 5: The accuracy degradation of the solution, if the condition number is large, can be estimated as follows: Assuming that the condition number is $\kappa(\mathbf{A}) \approx 10^p$, then the number of accurate digits in the solution is (q-p) if the solution is computed in q-digit arithmetic.

²This algorithm is available in LAPACK, routines **sgesvx** and **slacon**.

9.1.3 Parallel LU Decomposition

The efficient parallel implementation of decomposing a non-singular matrix \mathbf{A} into its LU factorization requires that we address two main issues:

- 1. How to split the matrix **A** among the processors.
- 2. How to organize the triple loop so that efficient BLAS operations can be employed.

We consider here distributed memory computers so only parts of the matrix **A** are stored in each processor. The obvious ways to split the matrix **A** are by rows or by columns, but it may also be beneficial to split it in blocks. A better layout, often used in practice, is *interleaved storage* either by row or by column. We examine these different cases in some detail in the following.

With regards to organizing the triple nested loop – recall that we deal with an $\mathcal{O}(n^3)$ operation – there are six ways of permutating the indices (ijk), just like in matrix-matrix multiply that we discussed in chapter 2. In table 9.1 we present all six versions and basic operations involved.

We have defined the basic operation in table 9.1 as the operation involved in the innermost loop. In four versions this is a daxpy operation (i.e., double a (scalar) \mathbf{x} (vector) plus \mathbf{y} (vector), while in the other two versions (ijk and jik) it is a ddot (i.e., double dot product). These are both BLAS1 operations and cannot easily take advantage of $cache\ blocking$ or $data\ re-use$, see section 2.2.6. Thus, appropriate modifications of these basic loops are required to be able to employ BLAS2 and BLAS3 routines. This will depend on the specific way we layout the matrix \mathbf{A} in order to also maximize parallel efficiency. We examine this issue in more detail next.

Access By Rows

Let us first assume that the matrix **A** is accessed by rows as in the (kij) loop of table 9.1; see figure 9.2 for a schematic explanation. Let us also assume that the first processor P_1 holds the first row \mathbf{a}_1^T , P_2 holds \mathbf{a}_2^T and so on.

During the *first* elimination stage, the processor P_1 needs to send its row to all other processors so that processors $P_2 \dots P_n$ will simultaneously update their columns. Therefore, the operations

$$\begin{cases} \ell_{i1} = \frac{a_{i1}}{a_{11}} \\ a_{ij} = a_{ij} - \ell_{i1} a_{1j} \end{cases} \begin{cases} j = 2, \dots n \\ P_2, \dots P_n \end{cases}$$

can be performed in parallel. During this first stage, processor P_1 remains essentially idle after it communicates with the rest of the processor.

The second stage also starts with a communication step as P_2 needs to broadcast its new row to all other processors $P_3
ldots P_n$. It too remains idle after that, while $P_3
ldots P_n$ update their rows in parallel, and so on for the remaining stages. The computations of the multiplies as well as the updates are done in parallel but after the k^{th} stage, k processors $(P_1
ldots P_k)$ remain idle. This approach reduces significantly the parallel efficiency.

```
1. ijk Loop: A - by column (ddot)
                                              2. ikj Loop: A- by row (daxpy)
 for i=2, n
     for j = 2, i
          \ell_{i,j-1} = a_{i,j-1}/a_{j-1,j-1}
                                                for i=2,n
          for k = 1, j - 1
                                                   for k = 1, i - 1
              a_{ij} = a_{ij} - \ell_{ik} a_{ki}
                                                         \ell_{ik} = a_{ik}/a_{kk}
         endfor
                                                         for j = k + 1, n
     endfor
                                                             a_{ij} = a_{ij} - \ell_{ik} a_{kj}
     for j = i + 1, n
                                                         endfor
          for k = 1, i - 1
                                                    endfor
             a_{ij} = a_{ij} - \ell_{ik} a_{kj}
                                                endfor
         endfor
     endfor
 endfor
3. jik Loop: A - by column (ddot)
                                              4. jki Loop: A - by column (daxpy)
 for j=2,n
     for p = j, n
          \ell_{p,j-1} = a_{p,j-1}/a_{j-1,j-1}
                                                for j=2,n
     endfor
                                                   for p = j, n
     for i=2, j
                                                         \ell_{p,j-1} = a_{p,j-1}/a_{j-1,j-1}
          for k = 1, i - 1
                                                    endfor
              a_{ij} = a_{ij} - \ell_{ik} a_{kj}
                                                   for k = 1, j - 1
          endfor
                                                        for i = k + 1, n
     endfor
                                                            a_{ij} = a_{ij} - \ell_{ik} a_{kj}
     for i = j + 1, n
                                                        endfor
          for k = 1, j - 1
                                                    endfor
              a_{ij} = a_{ij} - \ell_{ik} a_{kj}
                                                endfor
          endfor
     endfor
 endfor
                                              6. kji Loop: A - by column (daxpy)
5. kij Loop: A - by row (daxpy)
                                               for k = 1, n - 1
                                                    for p = k + 1, n
 for k = 1, n - 1
                                                         \ell_{pk} = a_{pk}/a_{kk}
     for i = k + 1, n
          \ell_{ik} = a_{ik}/a_{kk}
                                                    endfor
                                                    for j = k + 1, n
          for j = k + 1, n
                                                         for i = k + 1, n
              a_{ij} = a_{ij} - \ell_{ik} a_{kj}
          endfor
                                                             a_{ij} = a_{ij} - \ell_{ik} a_{kj}
                                                         endfor
     endfor
                                                    endfor
 endfor
                                                endfor
```

Table 9.1: Six different ways of writing the LU triple loops.

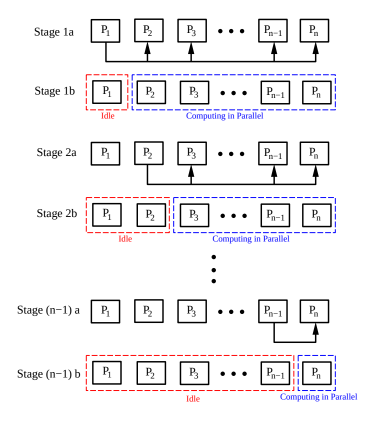


Figure 9.2: Schematic of communication (a) and computation (b) pattern when a matrix is partitioned such that each processor contains one row.

Some improvements can be made by overlapping communication with computation. For example, the broadcasting of the row of processor P_k can be done immediately after it is computed, so that the other processors receive it while they are updating their rows during the k^{th} stage of elimination. This overlapping of computation and communication is called send-ahead operation and it is quite common in distributed memory parallel computers. Another improvement would come about if we manage to increase the work done in parallel relative to communications. In general, the parallel efficiency improves as the ratio of computations to communications becomes larger. Thus, for given required communication the local work is increased as many rows are stored in each processor, so we have one, say, send-ahead operation every 100 rows on a P = 10 processor system and a matrix of n = 1000.

Blocked Layout

The row-blocked layout of the matrix **A** suggests that the practical way of storing **A** is by rows, and it extends also to a column-blocked layout, e.g., in the kji algorithm described in table 9.1. In this case, the multiplies ℓ_{ij} are computed on the processor P_j (or its corresponding number for blocked storage), and then ℓ_{ij} are broadcasted to all other (active) processors.

Partial pivoting in either the row-blocked or column-blocked parallel versions is handled differently. In the latter version, a search for the maximum pivot is performed within the processor while in the former (kij loop) the searching is across processors; this requires a fan-in algorithm for the max operator, as discussed in chapter 2.

Another improvement of the row-blocked or column-blocked layout schemes is to introduce a *cyclic* or *interleaved* layout, see figure 9.3. For example, assuming that we have available k = 10 processors for a 100×100 matrix **A**, then we pursue the following storage scheme:

$$P_1$$
: rows $1, 11, 21, ..., 91$
 P_2 : rows $2, 12, 22, ..., 92$
 \vdots
 P_{10} : rows $10, 20, ..., 100$

and similarly in a column-oriented storage scheme. Examining figure 9.3, we observe that during stage 1a processor one communicates row one to all other processors, and then in stage 1b all processors are active accomplishing row reduction. In comparison to the previous block setup in which rows one through ten would be assigned to processor one, rows 11 through 20 to processor two, etc., row two is now located on processor two. Hence in stage 2a, processor two communicates row two to all other processors, and in stage 2b all processors accomplish row reduction concurrently. In this manner, for the first m=90 stages all the processors remain active both in communication and computation. Some inefficiencies may occur during the last few stages, however, as the final row assigned to a processor is eliminated, hence retiring the processor from service for the elimination.

An even better option is to employ a block of rows (or columns) and assign these blocks in a cyclic or interleaved manner. The advantages of this approach are that no processor retires early and that BLAS2 and BLAS3 routines can be used because of the blocking. All processors see roughly the same amount of work, proportioned to 1/P, although the first processors work less, e.g., after they compute their first block.

Finally, a combination of row and column block interleaved storage of matrix **A** can be pursued. In this case we can imagine a mapping of b size blocks of **A** in a cyclic manner (of cycle length C) onto a *mesh-type* parallel computer consisting of $P = P_r \times P_c$ processors. Schematically, this arrangement is shown in figure 9.4.

This mapping can be established by setting

$$S_i = F(i, b)$$
 and $S_i = F(j, b)$,

with i = 0, ..., n-1 and j = 0, ..., n-1 (N being the size of **A**) where the value of the (S_i, S_j) pair defines the processor in the mesh architecture. The function F is defined as

function
$$F(i, b)$$

floor (i/b) modulo C
return;

where we assume that we deal with a square matrix and that $C^2 = P$. In the example of figure 9.4, we have P = 4 and C = 2. We have also assumed here for simplicity that the computer (mesh array) is symmetric but that may not be advantageous in practice, i.e, we

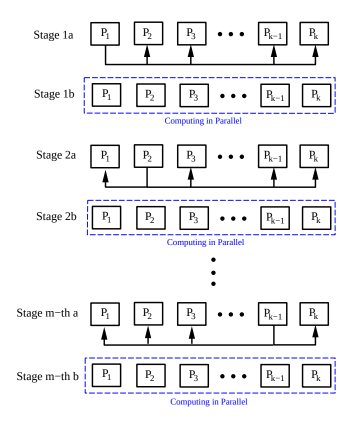


Figure 9.3: Schematic of communication (a) and computation (b) pattern when a matrix is partitioned with an interleaved layout.

may want to have $P_r > P_c$, that is more rows than columns. The advantage of this approach, in addition to its good parallel performance, is that it can make use of the BLAS3 routines, which provide high efficiency. The main steps of computing pivots, send-ahead, and parallel work on each processor that we presented earlier are also utilized here.

The plot of figure 9.5 shows a graphical representation of LU decomposition using BLAS3. The new entries to be computed are in the shaded area in the right lower corner of the matrix \mathbf{A} . The submatrix \mathbf{A}_s that is currently shown to be computed has a size $b \times b$, i.e., the size of the block of rows and columns. This implementation is included in ScaLAPACK software for distributed computers, see

www.netlib.org/scalapack

9.1.4 Parallel Back Substitution

We first present an *ideal* algorithm for parallel back substitution that provides a lower bound for its computational complexity. Although the lower bound may be unattainable given hardware constraints, it provides us with the "best-case" scenario that the algorithm can provide given unrestricted resources. By understanding the concept which yields the lower bound

1	2	1	2	1	2	1	2
3	4	3	4	3	4	3	4
1	2	1	2	1	2	1	2
3	4	3	4	3	4	3	4
1	2	1	2	1	2	1	2
3	4	3	4	3	4	3	4
1	2	1	2	1	2	1	2
3	4	3	4	3	4	3	4

Figure 9.4: 2D block interleaved mapping of a 16×16 matrix on P = 4 processors; here b = 2 and C = 2, and the processor id used above is given by $S_i + 2S_j + 1$. Here S_i and S_j have the values 0 or 1.

result, we hope that after incorporating relevant constraints we will obtain a reasonable algorithm. The ideal algorithm for parallel back substitution is based on a *divide-and-conquer* algorithm for inverting triangular matrices.

Let us consider the lower triangular matrix \mathbf{L} , which we decompose into submatrices of half size \mathbf{L}_1 \mathbf{L}_2 and \mathbf{L}_3 as follows

$$\mathbf{L} = \left[\begin{array}{cc} \mathbf{L}_1 & 0 \\ \mathbf{L}_2 & \mathbf{L}_3 \end{array} \right] \, .$$

We assume that $n = 2^k$ and that k is an integer. This is shown schematically in the plot of figure 9.6.

It is clear that \mathbf{L}_1 and \mathbf{L}_2 are themselves lower triangular matrices. We can prove that

$$\mathbf{L}^{-1} = \begin{bmatrix} \mathbf{L}_1^{-1} & 0 \\ \\ -\mathbf{L}_3^{-1} \mathbf{L}_2 \mathbf{L}_1^{-1} & \mathbf{L}_3^{-1} \end{bmatrix},$$

and take advantage of this equation to set up a divide-and-conquer algorithm for inverting the matrix \mathbf{L} of size n.

The main steps are shown in the following pseudo-code:

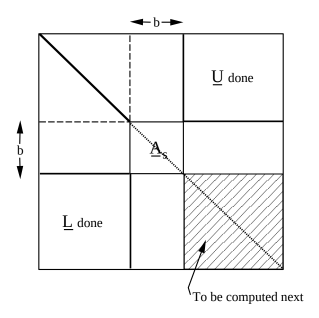


Figure 9.5: LU decomposition of a matrix **A** using 2D block interleaved storage. BLAS2 and BLAS3 can be effectively employed in this algorithm.

```
Function InvTriangular(L)
If size(L)= 1 return 1/L
Else
Set L1 top triangular part of L
Set L2 square part of L
Set L3 bottom triangular part of L

InvL1 =InvTriangular(L1)
InvL3 =InvTriangular(L3)
UpdateL2 = - InvL3 * L2 * InvL1

return L = \begin{vmatrix} InvL1 & 0 \\ UpdateL2 & InvL3 \end{vmatrix}
Endif
```

We can perform the inversion of L_1 and L_2 in parallel, so the cost C is

$$C[\operatorname{InvTriangular}(n)] = C[\operatorname{InvTriangular}(n/2) + C[mxm(n/2)],$$

where by mxm we denote the matrix-multiply. The ideal time for this matrix-multiply (assuming we employ n^3 processors) is $\mathcal{O}(\log n)$, and thus the ideal cost for inverting triangular matrices is $\mathcal{O}(\log^2 n)$. Unfortunately, this cost is impossible to realize in practice.

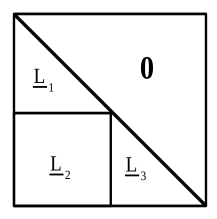


Figure 9.6: Decomposition of a lower triangular matrix L. Matrices L_1 and L_3 are also lower triangular.

The $\mathcal{O}(\log^2 n)$ time estimate is based on the equation

$$t(n) = t(n/2) + \mathcal{O}(\log n).$$

We set $k = \log_2 n$, and thus

$$t(k) = t(k-1) + \mathcal{O}(k)$$

$$= t(k-2) + \mathcal{O}(k) + \mathcal{O}(k-1)$$

$$= t(1) + \mathcal{O}(k) + \mathcal{O}(k-1) + \dots + 1$$

$$\approx \frac{k(k-1)}{2} \sim \mathcal{O}(k^2/2) \sim \mathcal{O}(\log^2 n).$$

Note that the $\mathcal{O}(\log n)$ ideal estimate for matrix multiply is based on the fact that all entries of the product matrix can be computed in parallel on $P = n^3$ processors, and then summed up using a fan-in algorithm which is $\mathcal{O}(\log n)$. This is of course unattainable at the moment since the n = 1000 size (ideally parallel) matrix-multiply would require more than one billion processors, more than what we currently have on this planet!

We now discuss how the *back substitution* step $\mathbf{U}\mathbf{x} = \mathbf{y}$, which is also a triangular system, can be performed in parallel. The algorithm proceeds by computing first

$$x_n = \frac{b_n^{(n-1)}}{a_{nn}^{(n-1)}}$$

while all other unknowns are computed from

$$\forall i = n-1, \dots 1: x_i = \frac{b_i^{(i-1)} - a_{ii}^{(i-1)} x_{i+1}}{a_{i,i}^{i-1}} = \frac{b_i - u_{i,i+1} x_{i+1} - u_{i,i+2} x_{i+2} - \dots u_{in} x_{in}}{u_{ii}},$$

where u_{ij} denote entries of the matrix **U**.

The most obvious implementation of this is the following:

for
$$j = n, 1$$

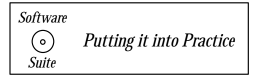
for $j = i + 1, n$
 $b_i = b_i - u_{ij}x_j$
endfor
 $x_i = b_i/u_{ii}$
endfor

This assumes that \mathbf{U} is stored by rows, which are accessed in the innermost loop j. More specifically, the matrix \mathbf{U} is stored in blocks of rows assigned to each processor. The corresponding operation employs the BLAS1 ddot routine, which implements the inner (dot) product of the innermost loop. However, if \mathbf{U} is stored in blocks of columns per processor, then the (ij) loop of the ddot product implementation above has to be reversed as follows

for
$$j = n, 1$$

 $x_j = b_j/u_{jj}$
for $i = 1, j - 1$
 $b_i = b_i - u_{ij}x_j$
endfor
endfor

Here the innermost loop (i) generates updates of the numerators for the final answer x_j while the outer loop (j) sweeps the matrix \mathbf{U} by columns starting from right to left. This is sometimes referred to as *left-looking* access of \mathbf{U} , and it involves *daxpy* operations in the innermost loop.



Prior to presenting a parallel implementation of Gaussian elimination, we will first discuss one new MPI function not introduced previously: MPI_Bcast . This function allows us to distribute to all processes within the communicator an identical piece of data. In the case of Gaussian elimination, MPI_Bcast will allow us to "broadcast" to all processes a particular row being used in the elimination. We will now present the function call syntax, argument list explanation, usage example and some remarks.

MPI_Bcast:

Function Call Syntax

```
int MPI_Bcast(
    void* buffer /* in/out */,
    int count /* in */,
    MPI_Datatype datatype /* in */,
    int root /* in */,
    MPI_Comm comm /* in */)
```

Understanding the Argument List

- buffer starting address of the send buffer.
- count number of elements in the send buffer.
- datatype data type of the elements in the send buffer.
- root rank of the process broadcasting its data.
- comm communicator.

Example of Usage

```
int mynode, totalnodes;
int datasize; // number of data units to be broadcast
int root; // process which is broadcasting its data

MPI_Init(&argc,&argv);
MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

// Determine datasize and root

double * databuffer = new double[datasize];

// Fill in databuffer array with data to be broadcast

MPI_Bcast(databuffer,datasize,MPI_DOUBLE,root,MPI_COMM_WORLD);

// At this point, every process has received into the

// databuffer array the data from process root
```

Remarks

• Each process will make an identical call of the MPI_Bcast function. On the broadcasting (root) process, the *buffer* array contains the data to be broadcast. At the conclusion of the call, all processes have obtained a copy of the contents of the *buffer* array from process root, see figure 9.7.

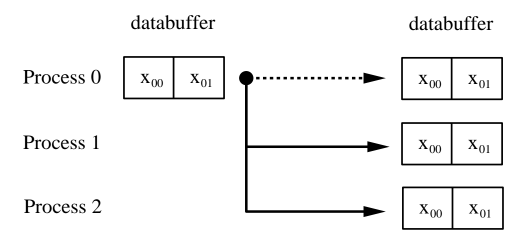


Figure 9.7: *MPI_Bcast* schematic demonstrating a broadcast of two data objects from process zero to all other processes.

We now present a parallel implementation of Gaussian elimination with back substitution. As a model problem, we solve for the interpolating polynomial of the Runge function (see section 3.1.4) by forming a Vandermonde matrix based on the Chebyshev points. Recall that the goal is to find the polynomial coefficients by solving the system $\mathbf{A}\mathbf{x} = \mathbf{b}$ where \mathbf{A} is the Vandermonde matrix and \mathbf{b} is the function of interest evaluated at the interpolation points.

To better explain the code, we have broken the entire program into six parts, labeled part one through part six. The six parts break down the code as follows:

- 1. Part 1 MPI initialization/setup and initial memory allocations.
- 2. Part 2 Generation of the matrix rows local to each process.
- 3. Part 3 Gaussian elimination of the augmented matrix.
- 4. Part 4 Preparation for back substitution.
- 5. Part 5 Back substitution to find the solution.
- 6. Part 6 Program finalization and clean-up.

For each part, we will first present the code and then present a collection of remarks elucidating the salient points within each part.

Part 1 - MPI initialization

```
#include <iostream.h>
#include <iomanip.h>
#include "SCmathlib.h"
#include "SCchapter3.h"
#include<mpi.h>

void ChebyVandermonde(int npts, double *A, int row);
```

```
// Global variable to set size of the system
const int size = 10;

int main(int argc, char *argv[]){
  int i,j,k,index;
  int mynode, totalnodes;
  double scaling;
  MPI_Status status;

MPI_Init(&argc,&argv);
  MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
  MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

int numrows = size/totalnodes;
  double **A_local = new double*[numrows];
  int * myrows = new int[numrows];
```

Remark 1: Notice that for this program we use a global constant variable to denote the size of the matrix system. By placing the variable declaration outside of the main function, the declaration is global to all functions (including the main function).

Remark 2: We have made the assumption that the size of the matrix is evenly divisible by the number of processors we are using. If this were not the case, we would have to properly take this into account by having different numbers of rows per processor.

Part 2 - Generation of matrix rows

```
/* PART 2 */
double * xpts = new double[size];
ChebyshevPoints(size,xpts);

for(i=0;i<numrows;i++){
    A_local[i] = new double[size+1];
    index = mynode + totalnodes*i;
    myrows[i] = index;
    ChebyVandermonde(size,A_local[i],index);

    // Set-up right-hand-side as the Runge function
    A_local[i][size] = 1.0/(1.0+25.0*xpts[index]*xpts[index]);
}
delete[] xpts;

double * tmp = new double[size+1];
double * x = new double[size];</pre>
```

Remark 1: Notice that we allocate for each row (size+1) columns. Recall that in Gaussian elimination it is necessary for us to act on the right-hand-side (the vector \mathbf{b}) as we do the row reduction. We can eliminate the extra communication cost that would come by handling the right-hand-side separately by forming an augmented matrix as shown in figure 9.8.

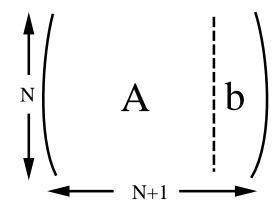


Figure 9.8: The augmented matrix consists of the original matrix $\bf A$ with the right-hand-side vector $\bf b$ appended as an additional column.

As you will see in the next section, all row reduction steps will be accomplished on (size + 1) columns so that both the matrix and right-hand-side are updated properly.

Remark 2: To keep track of which rows a processor possesses, we store the indices of the rows in an array named *myrows*. This array contains *numrows* entries, each entry denoting for which rows in the matrix the processor is responsible.

Part 3 - Gaussian elimination

```
for(j=cnt;j<numrows;j++){
    scaling = A_local[j][i]/tmp[i];
    for(k=i;k<size+1;k++)
        A_local[j][k] = A_local[j][k] - scaling*tmp[k];
}
</pre>
```

Remark 1: We use the integer variable *cnt* to keep track of how many rows *on each processor* have been reduced. Recall that each processor has its own copy of the variable *cnt*, and hence on each processor it can be used to keep track of what is the active row.

Remark 2: We have chosen the *cyclic distribution* discussed earlier. A schematic of the communication and computation pattern for a four processor run executed on a size = 12 system is shown in figure 9.9.

Part 4 - Preparation for back substitution

```
/* PART 4 */
/* On each processor, initialize the value of x as equal to
    the modified (by Gaussian elimination) right-hand-side if
    that information is on the processor, otherwise initialize
    to zero.
*/

cnt = 0;
for(i=0;i<size;i++){
    if(i==myrows[cnt]){
        x[i] = A_local[cnt][size];
        cnt++;
    }
    else
        x[i] = 0;
}</pre>
```

Remark: To accomplish the back substitution, we first initialize the solution by setting the array entry x[i] equal to the last column of the augmented matrix (which contains the modified right-hand-side) for those rows for which a processor is responsible, and equal to zero for all rows for which a particular processor is not responsible.

Part 5 - Back substitution

```
/* PART 5 */
/* Backsolve to find the solution x */
cnt = numrows-1;
for(i=size-1;i>0;i--){
```

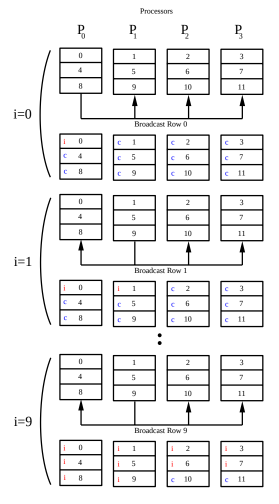


Figure 9.9: Schematic of the communication and computation pattern for a four processor run executed on a size = 12 system. Iterations i = 0, i = 1, and i = 9 are shown. The letter **i** denotes an idle row, while a c denotes a computing row.

```
if(cnt>=0){
    if(i == myrows[cnt]){
      x[i] = x[i]/A_local[cnt][i];
      MPI_Bcast(x+i,1,MPI_DOUBLE,mynode,MPI_COMM_WORLD);
      cnt--;
    }
    else
      MPI_Bcast(x+i,1,MPI_DOUBLE,i%totalnodes,MPI_COMM_WORLD);
  }
  else
    MPI_Bcast(x+i,1,MPI_DOUBLE,i%totalnodes,MPI_COMM_WORLD);
  for(j=0;j<=cnt;j++)
    x[myrows[j]] = x[myrows[j]] - A_local[j][i]*x[i];
}
if(mynode==0){
 x[0] = x[0]/A_local[cnt][0];
 MPI_Bcast(x,1,MPI_DOUBLE,0,MPI_COMM_WORLD);
}
else
 MPI_Bcast(x,1,MPI_DOUBLE,0,MPI_COMM_WORLD);
```

Remark 1: Observe that the variable cnt is initialized to (numrows-1) and is decremented in this loop. Recall that we want to traverse back up the rows (hence the name back substitution). As a processor computes the solution for a row for which it is responsible, it then broadcasts the result to all the other processors.

Remark 2: Why do we check for $cnt \geq 0$? If you examine this loop carefully, you will notice that at some point the variable cnt is equal to -1. This occurs when all the solution components on a processor have been computed. We must verify that $cnt \geq 0$ prior to attempting to access myrows[cnt], otherwise we are performing an illegal memory access, and hence the process may fail. We know, however, that once a processor's value of cnt is equal to -1 it merely needs to obtain the updated values from other processors, and hence we can immediately proceed to an MPI_Bcast call to obtain the solution from another processor.

Remark 3: Observe in the MPI_Bcast arguments that we use pointer arithmetic for updating the address which is passed to the function. Recall that x is a pointer (and hence has as its value an address). The expression (x + i) is equivalent to the expression &x[i], which can be read as "the address of the array element x[i]."

Part 6 - Program finalization

```
/* PART 6 */
if(mynode==0){
```

```
for(i=0;i<size;i++)</pre>
      cout << x[i] << endl;</pre>
  }
  delete[] tmp;
  delete[] myrows;
  for(i=0;i<numrows;i++)</pre>
    delete[] A_local[i];
  delete[] A_local;
  MPI_Finalize();
}
void ChebyVandermonde(int npts, double *A, int row){
  int i,j;
  double * x = new double[npts];
  ChebyshevPoints(npts,x);
  for(j=0;j<npts;j++)</pre>
    A[j] = pow(x[row], j);
  delete[] x;
}
```

Remark: We conclude the program by printing the solution from the first processor (mynode = 0). If we were not to put the *if* statement there, all processors would print the solution.

9.1.5 Gaussian Elimination and Sparse Systems

Many linear systems that arise from discretization of partial differential equations, as we have seen in chapters 6 and 7, are sparse and more specifically bandeded. Assuming that we deal with symmetrically banded systems of (semi)-bandwidth m, we can modify accordingly the (ijk) loops of table 9.1 to account for this sparsity. For example, the kij loop is modified as follows:

```
for k = 1, n - 1

for i = k + 1, \min(k + m, n)

\ell_{ik} = a_{ik}/a_{kk}

for j = k + 1, \min(k + m, n)

a_{ij} = a_{ij} - \ell_{ik}a_{kj}

endfor

endfor

endfor
```

The computational complexity of this algorithm is significantly less on a serial computer than on a parallel computer. The operation count for LU decomposition is about $\mathcal{O}(nm^2)$ for an $n \times n$ matrix with bandwidth m and for the backsolve is $\mathcal{O}(nm)$. However, on a parallel computer a straightforward implementation of the above algorithm would result in large inefficiencies. For example, the row- (column-) blocked interleaved scheme discussed earlier becomes very inefficient when m < P, where P is the total number of processors as only m processors are effectively used. Clearly, the case with m = 1, i.e., the tridiagonal system, is the most difficult case and needs to be handled differently; we study this case next.

9.1.6 Parallel Cyclic Reduction for Tridiagonal Systems

Several algorithms have been developed over the years for the parallel solution of tridiagonal linear systems including recursive doubling [82], cyclic reduction [13], domain decomposition [90], and their many variants. Here we present the cyclic reduction method, which has been one of the most successful approaches.

The main idea of cyclic reduction is to group the unknowns in *even* and *odd-numbered* entries, just like in black/red Gauss-Seidel (see section 7.2.4), and successively eliminate the odd-numbered entries. Most of the operations in this process can be done in parallel. We present here some details by considering a specific small system in order to illustrate how we manipulate the equations.

Let us consider the tridiagonal system

$$a_i x_{i-1} + b_i x_i + c_i x_{i+1} = F_i, \quad i = 1, \dots, n$$
 (9.8)

where a_i , b_i , c_i , and F_i are given, and we also assume that $n = 2^p - 1$. If $n \neq 2^p - 1$ then we add additional trivial equations of the form $x_i = 0$, $i = n + 1, \ldots, 2^p - 1$.

The key idea is then to combine linearly the equations in order to eliminate the oddnumbered unknowns

$$x_1, x_3, x_5, \ldots, x_n$$

in the first stage. We then re-order (i.e., re-number) the unknowns and repeat this process until we arrive at a single equation with one unknown. Upon solution of that equation, we march backwards to obtain the rest of the unknowns. To do this we combine the equations in triplets.

Next, we demonstrate this for the case of $n=7=2^3-1$ unknowns for which we have three triplets. We start by forming the first triplet from the first three equations. To this end, we multiply by the parameters α_2 , β_2 , γ_2 to get

$$\alpha_2 b_1 x_1 + \alpha_2 c_1 x_2 = \alpha_2 F_1$$

$$\beta_2 a_2 x_1 + \beta_2 b_2 x_2 + \beta_2 c_2 x_3 = \beta_2 F_2$$

$$\gamma_2 a_3 x_2 + \gamma_2 b_3 x_3 + \gamma_2 c_3 x_4 = \gamma_2 F_3.$$

In order to eliminate x_1 and x_3 , we add the equations and choose:

$$\beta_2 = 1$$

$$\alpha_2 b_1 + \beta_2 a_2 = 0$$

$$\beta_2 c_2 + \gamma_2 b_3 = 0.$$

The resulted equation (sum of the above three equations) is

$$\underbrace{(\alpha_2 c_1 + \beta_2 b_2 + \gamma_2 a_3)}_{\hat{p}_2} x_2 + \underbrace{\gamma_2 c_3}_{\hat{c}_2} x_4 = \underbrace{\alpha_2 F_1 + \beta_2 F_2 + \gamma_2 F_3}_{\hat{F}_2}.$$

Similarly, combining again equations, i.e., the third, fourth and fifth equations obtained from equation (9.8) we form the second triplet from which we obtain

$$\underbrace{\alpha_4 a_3}_{\hat{a}_4} x_2 + \underbrace{(\alpha_4 c_3 + \beta_4 b_4 + \gamma_4 a_5)}_{\hat{b}_4} x_4 + \underbrace{\gamma_4 c_5}_{\hat{c}_4} x_6 = \underbrace{a_4 F_3 + \beta_4 F_4 + \gamma_4 F_5}_{\hat{F}_4},$$

and α_4 , β_4 and γ_4 are determined from

$$eta_4 = 1$$
 $lpha_4 b_3 + eta_4 a_4 = 0$
 $eta_4 c_4 + \gamma_4 b_5 = 0.$

Finally, for the third triplet we obtain, as before, the only surviving equation

$$\hat{a}_6 x_4 + \hat{b}_6 x_6 = \hat{F}_6$$

where the parameters α_6 , β_6 , γ_6 involved in the definition of \hat{a}_6 , \hat{b}_6 and \hat{F}_6 are determined by solving

$$\alpha_6 b_5 + \beta_6 a_6 = 0$$

$$\beta_6 c_6 + \gamma_b b_7 = 0.$$

We see that the three resulted equations also form a tridiagonal system, i.e.,

$$\hat{b}_2 x_2 + \hat{c}_2 x_4 = \hat{F}_2 \tag{9.9}$$

$$\hat{a}_4 x_2 + \hat{b}_4 x_4 + \hat{c}_4 x_6 = \hat{F}_4 \tag{9.10}$$

$$\hat{a}_6 x_4 + \hat{b}_6 x_6 = \hat{F}_6. \tag{9.11}$$

We can repeat the same elimination process as before, i.e., first multiply by α'_4 , β'_4 and γ'_4 respectively the above equations and choose

$$\alpha'_{4}\hat{b}_{2} + \beta'_{4}\hat{a}_{4} = 0 \beta'_{4}\hat{c}_{4} + \gamma'_{4}\hat{b}_{6} = 0$$

that leads to only one equation

$$\alpha_4^* x_4 = F_4^*.$$

Using back substitution, after we obtain x_4 from above, we can compute x_2 from the reduced equation (9.9) and x_6 from equation (9.11). Finally, we use the original equations to obtain x_1 , x_3 , x_5 and x_7 .

In summary, we perform the following steps:

1. Compute

$$(\alpha_2, \beta_2, \gamma_2)$$
$$(\alpha_4, \beta_4, \gamma_4)$$
$$(\alpha_6, \beta_6, \gamma_6)$$

2. Compute

$$(\hat{b}_2, \hat{c}_2, \hat{F}_2)$$

 $(\hat{a}_4, \hat{b}_4, \hat{c}_4, \hat{F}_4)$
 $(\hat{a}_6, \hat{b}_6, \hat{F}_6)$

3. Compute

$$(\alpha'_4, \beta'_4, \gamma'_4)$$

 (a_4^*, F_4^*)

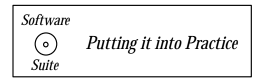
4. Solve for x_4 , x_2 , x_6 , x_1 , x_3 , x_5 and x_7 .

Remark: There exist non-singular matrices for which the above method will terminate early with the solution. The number of levels used above is the worst-case scenario. For some matrices, the depth of reduction can be truncated due to the solution being obtained for one of the variables. Immediately back-substitution can begin.

The operation count is $\mathcal{O}(13n)$ multiplications compared to $\mathcal{O}(5n)$ for the standard LU decomposition. However, this is the serial work but many of the above computations can be done in parallel. For example, let us assume that we have

$$P = \frac{n-1}{2}$$

processors and we store each triplet on one processor. The work for eliminating the oddnumbered unknowns can be done in parallel, and at the end of this stage each processor holds one reduced equation. To proceed, the processors need to exchange data. This can be done by nearest neighbor communications, e.g., P_2 will receive data from P_1 and P_3 , P_4 will receive data from P_3 and P_5 , and so on. This implies that about half of the processors (e.g., all the odd-numbered $P_1, P_3 ...$) will remain idle. After (p-1) reduction stages (where $n=2^p-1$) only one processor will solve the final equation. However, the rest of the processors are not quite retired yet, as in the back substitution they will all be called on duty again!



We now present implementations of cyclic reduction for tridiagonal systems. Due to the complex nature of the indexing involved, we will present both the serial and then the parallel version. As pointed out earlier, the serial version of this algorithm is rarely used because it

is actually more expensive than standard LU; we present it however because it is easier to understand the complex indexing in the serial setting without the additional complexity of the parallelization.

As a model problem, we solve for the system given in figure 9.10. Notice that the matrix **A** is of the form found in discretizations of the Laplace's equation with second-order finite difference schemes.

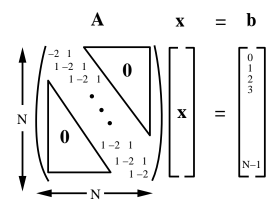


Figure 9.10: Tridiagonal system used as a model problem for cyclic reduction.

To better explain the code, we have broken the entire program into multiple parts. The serial code is broken down into three parts as follows:

- 1. Part 1 Memory allocation and generation of the matrix **A**.
- 2. Part 2 Cyclic reduction stages.
- 3. Part 3 Cyclic reduction back substitution to recover the solution.

For each part, we will first present the code and then present a collection of remarks elucidating the salient points within each part.

Serial

```
#include <iostream.h>
#include <iomanip.h>
#include "SCmathlib.h"

const int size = 15;

void main(){
  int i,j,k;
  int index1,index2,offset;
  double alpha,gamma;
```

Part 1 - Memory allocation and generation of matrix

```
/* Part 1 */
double * x = new double[size];
for(i=0;i<size;i++)</pre>
  x[i] = 0.0;
double * F = new double[size];
double ** A = new double*[size];
for(i=0;i<size;i++){</pre>
  A[i] = new double[size];
  for(j=0;j<size;j++)
    A[i][j] = 0.;
 F[i] = (double)i;
}
A[0][0] = -2.0; A[0][1] = 1.0;
A[size-1][size-2] = 1.0; A[size-1][size-1] = -2.0;
for(i=1;i<size-1;i++){
  A[i][i] = -2.0;
  A[i][i-1] = 1.0;
 A[i][i+1] = 1.0;
}
```

Remark 1: Observe that in this program we use more memory than necessary. Because we know that we are operating on a tridiagonal system, the memory required is greatly reduced from that which would be needed in **A** were full. Here we allocate memory as if **A** is a full matrix; we leave it as an exercise to modify this program to use only the necessary amount of memory.

Part 2 - Cyclic reduction

```
/* Part 2 */
for(i=0;i<log2(size+1)-1;i++){
  for(j=pow(2,i+1)-1;j<size;j=j+pow(2,i+1)){
    offset = pow(2,i);
    index1 = j - offset;
    index2 = j + offset;

    alpha = A[j][index1]/A[index1][index1];
    gamma = A[j][index2]/A[index2][index2];

  for(k=0;k<size;k++){</pre>
```

```
A[j][k] -= (alpha*A[index1][k] + gamma*A[index2][k]);
}
F[j] -= (alpha*F[index1] + gamma*F[index2]);
}
```

Remark 2: In the above code section, the first *for* loop (over *i*) iterates through the levels of reduction which occur, while the second *for* loop (over *j*) indexes which rows at each level are to be acted upon. Notice that both loops and the variables *index1* and *index2* are based on powers of two.

Part 3 - Back substitution

```
/* Part 3 */
int index = (size-1)/2;
x[index] = F[index]/A[index][index];
for(i=log2(size+1)-2;i>=0;i--){
  for(j=pow(2,i+1)-1;j<size;j=j+pow(2,i+1)){
    offset = pow(2,i);
    index1 = j - offset;
    index2 = j + offset;
    x[index1] = F[index1];
    x[index2] = F[index2];
    for(k=0;k\leq ize;k++){
      if(k!= index1)
        x[index1] -= A[index1][k]*x[k];
      if(k!= index2)
        x[index2] -= A[index2][k]*x[k];
    }
    x[index1] = x[index1]/A[index1][index1];
    x[index2] = x[index2]/A[index2][index2];
  }
}
for(i=0;i<size;i++){
  cout << x[i] << endl;</pre>
}
delete[] x;
delete[] F;
for(i=0;i<size;i++)</pre>
  delete[] A[i];
```

```
delete[] A;
}
```

Remark 3: Once the full reduction has occurred, we must traverse back up the reduction tree. Note that the two *for* loops accomplish this traversal.

Parallel

We now present the parallel version of the serial code presented above. We assume that given P processors, we will accomplish cyclic reduction on a matrix of size $2^{\log_2(P+1)+1} - 1$. This amounts to associating three rows per processor during the first stage. The parallel code is broken down into four parts as follows:

- 1. Part 1 MPI initialization and both memory allocation and generation of the matrix **A**.
- 2. Part 2 Parallel cyclic reduction stages.
- 3. Part 3 Parallel cyclic reduction back substitution to distribute necessary information.
- 4. Part 4 Solution for the odd rows for which each process is responsible.

For each part, we will first present the code and then present a collection of remarks elucidating the salient points within each part.

Part 1 - MPI initialization

```
#include <iostream.h>
#include <iomanip.h>
#include "SCmathlib.h"
#include<mpi.h>

int main(int argc, char *argv[]){
   int i,j,k,size,index;
   int index1,index2;
   int mynode, totalnodes;
   double alpha,gamma;
   const int numrows = 5;
   MPI_Status status;

MPI_Init(&argc,&argv);
   MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
   MPI_Comm_rank(MPI_COMM_WORLD, &mynode);

size = (int) pow(2,log2(totalnodes+1)+1)-1;
```

```
double ** A = new double*[numrows];
for(i=0;i<numrows;i++){</pre>
  A[i] = new double[size+1];
  for(j=0;j<size+1;j++)</pre>
    A[i][j] = 0.0;
}
if(mynode==0){
  A[0][0] = -2.0; A[0][1] = 1.0;
  A[1][0] = 1.0; A[1][1] = -2.0; A[1][2] = 1.0;
  A[2][1] = 1.0; A[2][2] = -2.0; A[2][3] = 1.0;
}
else if(mynode==(totalnodes-1)){
  index = 2*mynode;
  A[0][index-1] = 1.0; A[0][index] = -2.0;
  A[0][index+1] = 1.0;
  index = 2*mynode+1;
  A[1][index-1] = 1.0; A[1][index] = -2.0;
  A[1][index+1] = 1.0;
  A[2][size-2] = 1.0; A[2][size-1] = -2.0;
}
else{
  for(i=0;i<3;i++){
    index = i + 2*mynode;
    A[i][index-1] = 1.0;
    A[i][index]
                = -2.0;
    A[i][index+1] = 1.0;
  }
}
for(i=0;i<3;i++)
  A[i][size] = 2*mynode+i;
int numactivep = totalnodes;
int * activep = new int[totalnodes];
for(j=0;j<numactivep;j++)</pre>
  activep[j] = j;
for(j=0;j<size+1;j++){</pre>
  A[3][j] = A[0][j];
 A[4][j] = A[2][j];
}
```

Remark 1: Just as in the parallel Gaussian elimination code, we augment the matrix **A** with the right-hand-side (appending **A** with an extra column). This helps to minimize the communication by allowing us to communicate both the row and right-hand-side information simultaneously.

Part 2 - Cyclic reduction

```
/* Part 2 */
for(i=0;i<log2(size+1)-1;i++){}
   for(j=0;j<numactivep;j++){</pre>
     if(mynode==activep[j]){
       index1 = 2*mynode + 1 - pow(2,i);
       index2 = 2*mynode + 1 + pow(2,i);
       alpha = A[1][index1]/A[3][index1];
       gamma = A[1][index2]/A[4][index2];
       for(k=0;k\leq ize+1;k++)
         A[1][k] = (alpha*A[3][k] + gamma*A[4][k]);
       if(numactivep>1){
         if(j==0){
  MPI_Send(A[1],size+1,MPI_DOUBLE,activep[1],0,
                    MPI_COMM_WORLD);
 }
 else if(j==numactivep-1){
  MPI_Send(A[1],size+1,MPI_DOUBLE,activep[numactivep-2],
                    1,MPI_COMM_WORLD);
 }
 else if(j\%2==0){
  MPI_Send(A[1],size+1,MPI_DOUBLE,activep[j-1],
                    1,MPI_COMM_WORLD);
  MPI_Send(A[1],size+1,MPI_DOUBLE,activep[j+1],
                    O, MPI_COMM_WORLD);
}
 else{
  MPI_Recv(A[3],size+1,MPI_DOUBLE,activep[j-1],0,
                    MPI_COMM_WORLD,&status);
  MPI_Recv(A[4],size+1,MPI_DOUBLE,activep[j+1],1,
                    MPI_COMM_WORLD,&status);
}
       }
     }
```

```
numactivep = 0;
for(j=activep[1];j<totalnodes;j=j+pow(2,i+1)){
   activep[numactivep++]=j;
}</pre>
```

Remark 2: The communication is accomplished through a series of MPI_Send and MPI_Recv calls. Each processor is communicating (either sending or receiving) from at most two other processors. To keep track of whom is to be sending/receiving, we maintain an active processor list within the integer array *activep*. A communication schematic for cyclic reduction using seven processors is given in figure 9.11.

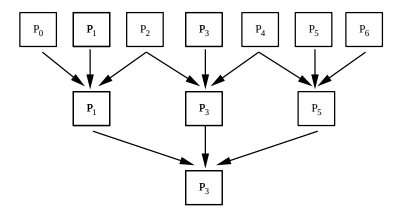


Figure 9.11: Cyclic reduction communication pattern for seven processor case.

Part 3 - Back substitution

```
/* Part 3 */
double * x = new double[totalnodes];
for(j=0;j<totalnodes;j++)
    x[j] = 0.0;

if(mynode==activep[0]){
    x[mynode] = A[1][size]/A[1][(size-1)/2];
}

double tmp;</pre>
```

```
for(i=log2(size+1)-3;i>=0;i--){
  tmp = x[mynode];
 MPI_Allgather(&tmp,1,MPI_DOUBLE,x,1,MPI_DOUBLE,
                 MPI_COMM_WORLD);
 numactivep = 0;
  for(j=activep[0]-pow(2,i);j<totalnodes;j=j+pow(2,i+1)){</pre>
    activep[numactivep++]=j;
  }
  for(j=0;j<numactivep;j++){</pre>
    if(mynode == activep[j]){
      x[mynode] = A[1][size];
      for(k=0;k<totalnodes;k++){</pre>
        if(k!=mynode)
          x[mynode] -= A[1][2*k+1]*x[k];
      }
      x[mynode] = x[mynode]/A[1][2*mynode+1];
    }
 }
}
tmp = x[mynode];
MPI_Allgather(&tmp,1,MPI_DOUBLE,x,1,MPI_DOUBLE,
              MPI_COMM_WORLD);
```

Remark 3: A schematic for the backward solve communication is given in figure 9.12. Notice that is varies slightly from that of the forward part of the reduction. After a processor has found the solution for its row and has communicated that information to the appropriate processors, it is no longer active. This can be observed in figure 9.12 - observe that processor P_3 no longer has things to compute in the second and third levels.

Remark 4: We use the $MPI_Allgather$ command so that at any given level all the processors have the available solution up to that point. This all inclusive communication could be replaced by MPI_Send/MPI_Recv pairs where only those processors requiring particular information would be updated.

Part 4 - Solving for odd rows

```
/* Part 4 */
for(k=0;k<totalnodes;k++){
   A[0][size] -= A[0][2*k+1]*x[k];
   A[2][size] -= A[2][2*k+1]*x[k];
}
A[0][size] = A[0][size]/A[0][2*mynode];
A[1][size] = x[mynode];</pre>
```

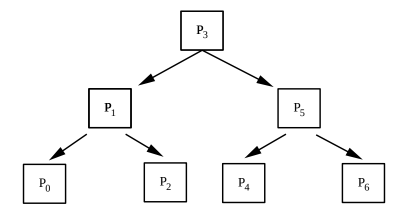


Figure 9.12: A schematic for the backward solve communication when seven processors are used.

```
A[2][size] = A[2][size]/A[2][2*mynode+2];

delete[] activep;
for(i=0;i<numrows;i++)
   delete[] A[i];
delete[] A;
delete[] x;</pre>

MPI_Finalize();
}
```

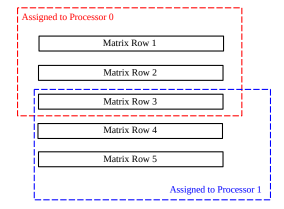


Figure 9.13: Overlap in row solutions for two processor case.

Remark 5: The program concludes with each processor computing the solution for the odd rows for which it was responsible. If the total solution vector were needed on all processors, $MPI_Allgather$ could be used to collect the solution on each processor. Note that some additional logic would be necessary to properly take into account the overlap in row distribution (i.e., both processor zero and processor one solve for the solution of matrix row number three) as shown in figure 9.13.

9.2 Cholesky Factorization

A special case of the LU decomposition for a $symmetric\ positive-definite\ matrix\ {\bf A}$ can be obtained in the form

$$\mathbf{A} = \mathbf{L} \; \mathbf{L}^T$$
.

Here, the matrix \mathbf{U} is equal to the transpose of \mathbf{L} due to symmetry. There are many possibilities for \mathbf{L} but this factorization is unique if we require that all diagonal elements of \mathbf{L} be positive. We note that in this case \mathbf{L} is different than the matrix we obtain in the LU decomposition, where all diagonal elements are equal to one.

Instead of following the standard LU decomposition, we can obtain explicitly the elements ℓ_{ij} of **L** by setting:

$$\begin{bmatrix} a_{11} & a_{12} & \dots & a_{1n} \\ \vdots & & & & \\ \vdots & & & & \\ \vdots & & & & \\ a_{n1} & \dots & \dots & a_{nn} \end{bmatrix} = \begin{bmatrix} \ell_{11} & & & & & \\ \ell_{12} & \ell_{22} & & & & \\ \vdots & \vdots & \ddots & & & \\ \vdots & \vdots & \ddots & & & \\ \ell_{n1} & \ell_{n2} & \dots & & \ell_{nn} \end{bmatrix} \begin{bmatrix} & \ell_{11} & \ell_{21} & \dots & & \ell_{n1} \\ & & \ell_{22} & & & \ell_{n2} \\ & & & \ddots & & \vdots \\ & & & & \ddots & & \\ \ell_{n1} & \ell_{n2} & \dots & & \ell_{nn} \end{bmatrix}$$

Next, we equate elements on both sides to obtain

$$a_{i1} = \ell_{i1}\ell_{11}$$
 and $a_{11}^2 = \ell_{11}^2$

for the entire first column (i = 1, ..., n), and similarly for the other columns. The following pseudo-code summarizes the Cholesky algorithm:

$$for \ j=1,n$$

$$\ell_{ij} = \sqrt{a_{ij} - \sum_{k=1}^{i-1} \ell_{jk}^2}$$

$$for \ i=j+1,n$$

$$\ell_{ij} = \frac{a_{ij} - \sum_{k=1}^{j-1} \ell_{ik}\ell_{jk}}{\ell_{jj}}$$

$$end for$$

$$end for$$

Remark 1: The Cholesky factorization algorithm is stable and thus it does not require pivoting [25].

Remark 2: The Cholesky algorithm requires about half the memory and about half the operations of the LU decomposition.

Remark 3: The positive-definite property is important in obtaining the ℓ_{ij} without partial pivoting. In fact, partial pivoting can destroy the symmetry of a matrix **A**.

Remark 4: In some cases an *incomplete* or *approximate* Cholesky decomposition is required, e.g. as a preconditioner in accelerating the convergence of iterative solvers, see section 9.4.2. This is achieved by simply filling in with zeros the entries of **L**, which have corresponding zero entries in the original (presumably sparse) matrix **A**.

9.3 QR Factorization and Householder Transformation

The LU factorization is not the only way of factorizing a matrix \mathbf{A} . The Householder transformation we present here is the basis of an efficient factorization of a general matrix

$$\mathbf{A} = \mathbf{Q}\mathbf{R}$$
,

where \mathbf{Q} is an orthogonal matrix and \mathbf{R} is an upper triangular matrix. We have already studied in section 2.2.9 how to achieve such a QR decomposition by orthogonalizing vectors via the Gram-Schmidt procedure, but the procedure we present here is always stable and much more efficient.

We start by considering the following important operation in scientific computing:

• How to take a full vector and produce a special vector with only one of its entries non-zero.

This is accomplished efficiently in terms of the (orthogonal) Householder matrix, which is defined by

$$\mathbf{H} = \mathbf{I} - 2 \frac{\mathbf{w} \mathbf{w}^T}{\mathbf{w}^T \mathbf{w}} \quad \forall \mathbf{w} \neq 0.$$

Of particular interest is the vector

$$\alpha \mathbf{e}_1 = (\alpha, 0, 0, \dots 0)^T$$

which can be created from an arbitrary vector \mathbf{x} if the vector \mathbf{w} is computed appropriately. To this end, we set \mathbf{w} such that:

$$\mathbf{H} \mathbf{x} = \left(\mathbf{I} - 2\frac{\mathbf{w}\mathbf{w}^T}{\mathbf{w}^T\mathbf{w}}\right) \mathbf{x} = \begin{bmatrix} \alpha \\ 0 \\ 0 \\ \vdots \\ 0 \end{bmatrix} = \alpha \mathbf{e}_1.$$

The solution to this problem is simple, and it is given by

$$\mathbf{w} = \mathbf{x} + sign(x_1) \cdot ||\mathbf{x}||_2 \mathbf{e}_1,$$

where x_1 is the first entry of the vector \mathbf{x} . This transformation from $\mathbf{H} \mathbf{x} \to (\alpha, 0, 0, \dots, 0)^T$ is called *Householder transformation*.

First, we summarize the algorithm that describes the Householder transformation in the following pseudo-code:

$$x_m = \max\{|x_1|, |x_2|, \dots, |x_n|\}$$

for $k = 1, n$
 $w_k = x_k/x_m$
endfor
 $\alpha = sign(w_1)[w_1^2 + w_2^2 + \dots + w_n^2]^{1/2}$
 $w_1 = w_1 + \alpha$
 $\alpha = -\alpha x_m$

Then, the desired vector is $(\alpha, 0, 0, \dots 0)^T$. The number of operations is proportional to $\mathcal{O}(n)$, that is it takes *linear work* only to accomplish this important operation.

Example: Consider the vector

$$\mathbf{x} = \begin{pmatrix} 1 \\ 2 \\ -3 \end{pmatrix},$$

then $x_m = 3$, and the intermediate values of the components of **w** are:

$$w_1 = \frac{1}{3}; \quad w_2 = \frac{2}{3}; \quad w_3 = -1.$$

Also, the intermediate value of α is:

$$\alpha = +\sqrt{\frac{1}{3^2} + \frac{2^2}{3^2} + (-1)^2} = 1.2472.$$

The updated values are then

$$w_1 = \frac{1}{3} + 1.2472 = 1.5805;$$
 $w_2 = \frac{2}{3};$ $w_3 = -1$

and

$$\alpha = -1.2472 \cdot 3 = -3.7416$$

while the desired vector is

$$\mathbf{H}\mathbf{x} = \left(\begin{array}{c} -3.7416 \\ 0 \\ 0 \end{array} \right).$$

Remark 1: The matrix-vector product with a Householder matrix

$$\mathbf{H} = \mathbf{I} - 2 \frac{\mathbf{w} \mathbf{w}^T}{\mathbf{w}^T \mathbf{w}}$$

is only an $\mathcal{O}(n)$ operation compared to the $\mathcal{O}(n^2)$ for a general matrix-vector product. This can be achieved from the relation

$$\mathbf{H} \mathbf{x} = \mathbf{x} - \beta \mathbf{w} (\mathbf{w}^T \mathbf{x}),$$

where $\beta^{-1} = \mathbf{w}^T \mathbf{w}/2$ is a scalar. We note that the right-hand-side is computed within one loop

$$x_i = x_i - \beta \cdot \gamma \cdot w_i, i = 1, \dots, n$$

where $\gamma = \mathbf{w}^T \mathbf{x}$ is also scalar.

Clearly, we do not need to construct explicitly **H** in this case – that cost would lead to an $\mathcal{O}(n^2)$ operation!

To accomplish the QR factorization of a square general $n \times n$ matrix **A**, we consider its columns and apply successively the Householder transformation in order to zero out the subdiagonal entries of each column. We do this in (n-1) stages, just like in LU decomposition.

Stage 1: We consider the *first column* of **A** and determine a Householder matrix \mathbf{H}_1 so that:

$$\mathbf{H}_1 \left[\begin{array}{c} a_{11} \\ a_{21} \\ \vdots \\ a_{n1} \end{array} \right] = \left[\begin{array}{c} \alpha_1 \\ 0 \\ \vdots \\ 0 \end{array} \right].$$

To determine \mathbf{H}_1 we simply need to apply the Householder transformation algorithm to obtain \mathbf{w}_1 of length n. After the first stage we overwrite \mathbf{A} by \mathbf{A}_1 where

$$\mathbf{A}_{1} = \begin{pmatrix} \alpha_{1} & a_{12}^{*} & \dots & a_{1n}^{*} \\ 0 & a_{22}^{*} & \dots & \dots \\ \vdots & \vdots & & & \\ 0 & a_{n2}^{*} & \dots & a_{nn}^{*} \end{pmatrix} = \mathbf{H}_{1}\mathbf{A},$$

which has all new elements (denoted by star) after the first column.

Stage 2: Next we consider the second column of the updated matrix

$$A \equiv A_1 = H_1 A$$

and take only the part below the diagonal, to obtain

$$\mathbf{H}_{2}^{*} \begin{bmatrix} a_{22}^{*} \\ a_{32}^{*} \\ \vdots \\ a_{n2}^{*} \end{bmatrix} = \begin{bmatrix} \alpha_{2} \\ 0 \\ \vdots \\ 0 \end{bmatrix},$$

which yields a vector \mathbf{w}_2 of length (n-1). This vector defines uniquely the Householder matrix

$$\mathbf{H}_2^* = \mathbf{I} - 2 \frac{\mathbf{w}_2 \mathbf{w}_2^T}{\mathbf{w}_2^T \mathbf{w}_2}.$$

Unlike \mathbf{H}_1 , here we first need to "inflate" \mathbf{H}_2^* to \mathbf{H}_2 and then overwrite \mathbf{A} by $\mathbf{H}_2\mathbf{A}_1$, where

$$\mathbf{H}_2 = \left(\begin{array}{ccc} 1 & \cdots & 0 \\ 0 & & \mathbf{H}_2^* \end{array}\right).$$

Stage k: In the k-stage of the QR procedure we produce a vector \mathbf{w}_k of length (n-k+1) by solving

$$\mathbf{H}_{k}^{*} \begin{bmatrix} a_{kk}^{*} \\ a_{k+1,k}^{*} \\ \vdots \\ a_{nk}^{*} \end{bmatrix} = \begin{bmatrix} \alpha_{k} \\ 0 \\ \vdots \\ 0 \end{bmatrix}.$$

Here again, we overwrite \mathbf{A} by

$$\mathbf{A}_k = \mathbf{H}_k \mathbf{A}_{k-1} \,,$$

and subsequently we "inflate" \mathbf{H}_{k}^{*} as

$$\mathbf{H}_k = \left[\begin{array}{cc} \mathbf{I}_{k-1} & 0 \\ 0 & \mathbf{H}_k^* \end{array} \right].$$

Remark 2: The efficiency of the Householder algorithm is based on the efficient multiplication

$$\mathbf{A}_{k+1} = \mathbf{H}_{k+1} \cdot \mathbf{A}_k \,,$$

which should not be performed explicitly but rather using the $\mathcal{O}(n)$ matrix-vector product algorithm we presented above; that is we compute

$$(\mathbf{I} - \beta \mathbf{w}_{n-k} \mathbf{w}_{n-k}^T) [\mathbf{a}_{kj}]$$

where

$$\beta^{-1} = \mathbf{w}_{n-k}^T \cdot \mathbf{w}_{n-k} / 2,$$

and \mathbf{a}_{kj} denotes the columns of \mathbf{A}_k with $j = k + 1, \dots, n - k$. The determination of \mathbf{H}_k^* requires (n - k) operations because the unknown vector \mathbf{w} is of (n - k) length.

Throughout the following section, the subscript for the vector \mathbf{w} will denote the current size of the vector; hence \mathbf{w}_{n-k} denotes the vector of size (n-k). In the algorithm to be presented, the notation w_{ij} denotes the i^{th} entry of the vector \mathbf{w}_{i} .

After a total of (n-1) stages we obtain an upper triangular matrix **R** with diagonal elements the α_k $(k=1,\ldots,n)$ and the other entries computed from:

$$r_{ij} = a_{ij} - \gamma w_{ik}$$

where $n \geq i$, $j \geq k$, and

$$\gamma = \beta \sum_{i=k}^{n} w_{ik} a_{ij}$$
$$\beta^{-1} = (\mathbf{w}_{n-k+1}^{T} \cdot \mathbf{w}_{n-k+1})/2$$

We note that the above formulas compute all the entries above the diagonal but also the r_{nn} entry.

Then, the matrix \mathbf{R} is:

$$\mathbf{R} = \begin{bmatrix} \alpha_1 & r_{12} & r_{13} & \dots & \dots & r_{1n} \\ & \alpha_2 & r_{23} & \dots & \dots & r_{2n} \\ & & & \alpha_3 & \dots & \dots & r_{3n} \\ & & & & \ddots & & \\ & & & & & \alpha_{n-1} & \\ & & & & & & r_{nn} \end{bmatrix}.$$

This matrix can be constructed by forming an equivalent Householder matrix \mathbf{H}_k of size n at each stage from the \mathbf{H}_k^* which has order (n-k+1). To this end, we simply set

$$\mathbf{H}_k = \left[\begin{array}{cc} \mathbf{I}_{k-1} & O \\ O & \mathbf{H}_k^* \end{array} \right],$$

and we also compute:

$$\mathbf{A}_k = \mathbf{H}_k \mathbf{A}_{k-1}$$

with

$$\mathbf{A}_1 \equiv \mathbf{A}$$

The upper triangular matrix is then

$$R = A_{n-1} = H_{n-1}A_{n-2} = \ldots = H_{n-1}H_{n-2}\ldots H_1A$$
.

We can invert the above equation if we set

$$\mathbf{Q}^T = \mathbf{H}_{n-1} \mathbf{H}_{n-2} \dots \mathbf{H}_1 \,,$$

then

$$\mathbf{Q}^{-1} = \mathbf{Q}^T$$

because \mathbf{H}_k are all orthogonal. Thus, $\mathbf{Q} \cdot \mathbf{R} = \mathbf{Q} \mathbf{Q}^T \mathbf{A} \Rightarrow \mathbf{A} = \mathbf{Q} \mathbf{R}$. We can now compute the orthogonal matrix \mathbf{Q} , i.e., $\mathbf{Q} = \mathbf{H}_1^T \mathbf{H}_1^T \dots \mathbf{H}_{n-1}^T$.

We have thus obtained a QR decomposition of an $n \times n$ matrix \mathbf{A} , similar to the Gram-Schmidt procedure but at reduced cost, since we are operating with shorter and shorter vectors in each stage. We first provide the pseudo-code for the Householder QR decomposition and then we will compute the exact operation count. The pseudo-code below returns only the value of \mathbf{R} . The matrix \mathbf{Q} can be formed from the above equation using the matrix \mathbf{H}_i formed by its vector \mathbf{w} from the Householder transformation.

Householder Algorithm

Begin Loop:
$$k = 1, ..., n-1$$
 (number of stages)
$$\bullet Solve \mathbf{H}_{k}^{*} \begin{bmatrix} a_{kk}^{*} \\ a_{k+1,k}^{*} \\ \vdots \\ a_{k}^{*} \end{bmatrix} = \begin{bmatrix} \alpha_{k} \\ 0 \\ \vdots \\ 0 \end{bmatrix}$$
(Obtain \mathbf{w}_{n-k+1})

•
$$r_{kk} = \alpha_k$$

• $Compute \ \beta^{-1} = \mathbf{w}_{n-k+1}^T \cdot \mathbf{w}_{n-k+1}/2$
 $Zero \ a_{ik} \colon i = k+1, \dots, n$
 $Begin \ Loop \colon j = k+1, \dots, n$
 $\gamma_j = 0$
 $Begin \ Loop \colon q = k, \dots n$
 $\gamma_j = \gamma_j + \beta w_{qk} a_{qj}$
 $End \ Loop$
 $Begin \ Loop \colon i = k, \dots, n$
 $r_{ij} = a_{ij} - \gamma_j w_{ik}$
 $a_{ij} = r_{ij}$
 $End \ Loop$
 $End \ Loop$
 $End \ Loop$
 $End \ Loop$

Example: Let us consider the 3×3 Hilbert matrix

$$\mathbf{A} = \begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ \frac{1}{2} & \frac{1}{3} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{4} & \frac{1}{5} \end{bmatrix}$$

and apply the Householder QR algorithm.

• In the first stage (k = 1) we solve

$$\mathbf{H}_{1} \begin{bmatrix} 1 \\ \frac{1}{2} \\ \frac{1}{3} \end{bmatrix} = \begin{bmatrix} \alpha_{1} \\ 0 \\ 0 \end{bmatrix} \Rightarrow \mathbf{w}_{1} = \begin{bmatrix} 2.1666 \\ 0.5 \\ 0.3333 \end{bmatrix}$$

and also $r_{11} = \alpha_1 = -1.1666$ and

$$\beta = \frac{2}{\mathbf{w}_1^T \mathbf{w}_1} = 0.3956.$$

Then, for j = 2, we calculate:

$$\gamma_2 = \beta[w_{11}a_{12} + w_{21}a_{22} + w_{31}a_{32}] = 0.5274$$

and thus

$$a_{12} := r_{12} = a_{12} - \gamma_2 w_{11} = -0.6429$$

 $a_{22} := r_{22} = a_{22} - \gamma_2 w_{21} = 0.0696$
 $a_{32} := r_{32} = a_{32} - \gamma_2 w_{31} = 0.0795$.

In the next iteration, j = 3, we calculate similarly:

$$\gamma_3 = \beta[w_1 a_{13} + w_2 a_{23} + w_3 a_{33}] = 0.3615$$

and thus

$$a_{13} := r_{13} = a_{13} - \gamma_3 w_{11} = -0.4500$$

 $a_{23} := r_{23} = a_{23} - \gamma_3 w_{21} = 0.0692$
 $a_{33} := r_{33} = a_{33} - \gamma_3 w_{31} = 0.0795$.

Also, we have that $a_{21} = a_{31} = 0$.

• In the second stage (k = 2) we solve

$$\mathbf{H}_{2}^{*} \left[\begin{array}{c} 0.0696 \\ 0.0795 \end{array} \right] = \left[\begin{array}{c} \alpha_{2} \\ 0 \end{array} \right] \Rightarrow \mathbf{w}_{2} = \left[\begin{array}{c} 2.3095 \\ 1.0000 \end{array} \right].$$

Also, $r_{22} = \alpha_2 = -0.1017$, and

$$\beta = 2/(\mathbf{w}_2^T \cdot \mathbf{w}_2) = 0.315759.$$

Then, for j = 3, we calculate:

$$\gamma_3 = \beta[w_{22}a_{23} + w_{32}a_{33}] = 0.0756$$

$$r_{23} = a_{23} - \gamma_3 w_{22} = -0.1053$$

$$r_{33} = a_{33} - \gamma_3 w_{32} = 0.0039,$$

where we note that a_{23} and a_{33} are the updated values, which were modified in the first stage.

At the conclusion of this example, we now have the resulting \mathbf{R} matrix of the $\mathbf{Q}\mathbf{R}$ decomposition, and we also have the Householder transformation vectors \mathbf{w} from which we can form \mathbf{Q} .

Computational Cost: The computational complexity of the QR decomposition, as described above, is determined by the cost of computing the matrix \mathbf{H}_k^* which is $\mathcal{O}(n-k)$, and also of computing \mathbf{A}_k from $\mathbf{A}_k = \mathbf{H}_k \mathbf{A}_{k-1}$ which requires $\mathcal{O}((n-k)^2)$ operations. Thus, the combined cost is

$$\sum_{k=1}^{n-1} (n-k) + \sum_{k=1}^{n-1} (n-k)^2 = \frac{(n-1)(n)}{2} + \frac{(n-1)(n)(2n-1)}{6} \approx \frac{n^3}{3}.$$

Calculating more carefully the constant factor and accounting for both additions and multiplications we have

$$\mathcal{O}\left(\frac{4}{3}n^3\right)$$
 QR versus $\mathcal{O}\left(\frac{2}{3}n^3\right)$ LU decomposition.

Remark 3: After we obtain the QR factorization of **A** we can solve a linear system $\mathbf{A}\mathbf{x} = \mathbf{b}$ as follows

$$Ax = b \Rightarrow QRx = b \Rightarrow Q^TQRx = Q^Tb \Rightarrow Rx = Q^Tb$$

which is an upper triangular system and can be solved by back substitution.

Remark 4: It can be shown that the QR decomposition based on the Householder transformation is always stable, see [47].

Remark 5: The QR-Householder decomposition is about twice as expensive as the LU decomposition and it also expands the bandwidth of a sparse matrix \mathbf{A} . In contrast, the LU decomposition preserves the bandwidth but it may be susceptible to numerical instabilities as demonstrated earlier. However, even with partial pivoting the LU decomposition is more efficient than the QR decomposition for large matrices \mathbf{A} .

Remark 6: (*Givens Rotations*) A third way of obtaining the QR decomposition of a matrix **A** (in addition to Gram-Schmidt and Householder) is to employ the Givens rotation matrix

$$\mathbf{R}(\theta) \equiv \left[\begin{array}{cc} \cos \theta & \sin \theta \\ -\sin \theta & \cos \theta \end{array} \right] \,,$$

which is orthonormal. By setting

$$\mathbf{R}(\theta) \cdot \left[\begin{array}{c} x \\ y \end{array} \right] = \left[\begin{array}{c} (x^2 + y^2)^{1/2} \\ 0 \end{array} \right]$$

we obtain

$$\cos \theta = \frac{x}{\sqrt{x^2 + y^2}}; \quad \sin \theta = \frac{y}{\sqrt{x^2 + y^2}}.$$

Based on this idea we construct a general rotation matrix

$$\mathbf{R}(\theta; i, j) = \begin{bmatrix} 1 & & & & & \\ & \ddots & & & & \\ & & \cos \theta & \sin \theta & & O \\ & & & -\sin \theta & \cos \theta & & \\ & O & & \ddots & & \\ & & & & 1 \end{bmatrix}$$

that can be employed to zero out *one entry* in each iteration cycle instead of a column as in the Householder transformation. However, the Givens rotation is *twice more expensive* than the Householder and *four times* more expensive than the LU. It is not used in solving *square* linear systems but it is used in solutions of *least squares* linear systems and also in eigensolvers.

Remark 7: The stability of the Householder method (also the Givens rotation) is due to the fact that the **Q** matrix is a product of orthogonal matrices, which have condition number equal to one.

9.3.1 Hessenberg and Tridiagonal Reduction

We now consider the transformation of a matrix A to an upper triangular which also has the first lower diagonal *non-zero*. Specifically, this matrix has the form

and it is called upper Hessenberg matrix.

The Householder transformation procedure can also be used to obtain this for general matrices. If the matrix is also *symmetric* then the resulted matrix is tridiagonal.

The reduction algorithm involves (n-2) stages of elimination. We want to obtain

$$\mathbf{H_e} = \mathbf{H} \cdot \mathbf{A} \cdot \mathbf{H}^T,$$

where the matrix $\mathbf{H} = \mathbf{H}_{n-2} \cdot \mathbf{H}_{n-1} \dots \mathbf{H}_1$. These are Householder matrices, which are computed from zeroing out sub-columns of \mathbf{A} and its updated versions, i.e.,

$$\mathbf{H}_1 = \left[\begin{array}{cc} \mathbf{I}_1 & 0 \\ 0 & \mathbf{H}_1^* \end{array} \right] \,,$$

where

$$\mathbf{H}_{1}^{*} \begin{bmatrix} a_{21} \\ a_{31} \\ \vdots \\ a_{n1} \end{bmatrix} = \begin{bmatrix} \alpha_{1} \\ 0 \\ \vdots \\ 0 \end{bmatrix},$$

and

$$\mathbf{A}_1 = \mathbf{H}_1 \mathbf{A} \mathbf{H}_1^T$$

and so on ...

Clearly, **A** has a new first column with $a_{11} \neq 0$ and $a_{21} \neq 0$ but all other entries are equal to zero.

Example: We consider again the 3×3 Hilbert matrix

$$\mathbf{A} = \begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ \frac{1}{2} & \frac{1}{3} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{4} & \frac{1}{5} \end{bmatrix}$$

which we transform into an upper Hessenberg matrix in one (3-2) stage. Since A is also symmetric we expect a tridiagonal matrix H_e .

• In the first stage (k = 1) we solve

$$\mathbf{H}_{1}^{*} \begin{bmatrix} \frac{1}{2} \\ \frac{1}{3} \end{bmatrix} = \begin{bmatrix} \alpha \\ 0 \end{bmatrix}$$

which gives

$$\mathbf{w}_2 = \left[\begin{array}{c} 2.2019 \\ 0.6666 \end{array} \right]$$

and $\alpha = -0.6009$.

We then obtain the new entries of **A** using $a_{ij} = a_{ij} - \gamma_i w_j$:

$$\mathbf{H_e} = \mathbf{H}_1 \mathbf{A} \mathbf{H}_1^T = \begin{bmatrix} 1 & 0 & 0 \\ \\ 0 & -0.8321 & -0.5547 \\ \\ 0 & -0.5547 & 0.8321 \end{bmatrix}$$

$$\begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ \frac{1}{2} & \frac{1}{3} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{4} & \frac{1}{5} \end{bmatrix} \begin{bmatrix} 1 & 0 & 0 \\ 0 & -0.8321 & -0.5547 \\ 0 & -0.5547 & 0.8321 \end{bmatrix} = \begin{bmatrix} 1.0 & -0.6009 & 0.0 \\ -0.6009 & 0.5231 & -0.0346 \\ 0.0 & -0.0346 & 0.0103 \end{bmatrix}.$$

Remark 1: The solution of the Hessenberg linear system

$$H_e \cdot x = b$$

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can be obtained with Gaussian elimination with partial pivoting, and it is guaranteed to be stable [94]. The computational complexity of this is $\mathcal{O}(n^2)$ including the partial pivoting cost.

We now present a function for computing the upper Hessenberg matrix given a matrix A. The function takes as input the matrix A and upon completion returns the upper Hessenberg matrix in place of the matrix A.

```
void Hessenberg(SCMatrix &A){
  int i,j,k,q;
  double beta, gamma;
  int N = A.Rows();
  SCVector *x = new SCVector(N),
    *w = new SCVector(N);
  for(k=0;k<N-2;k++){
    A.GetColumn(k,*x,k+1);
    A(k+1,k) = HouseholderTrans(*x,*w);
    beta = 2.0/dot(N-k-1,*w,*w);
    for(i=k+2;i<N;i++)</pre>
      A(i,k) = (*w)(i-k);
    for(j=k+1; j<N; j++){
      gamma = 0.0;
      for(q=k+1;q<N;q++)
        gamma += beta*(*w)(q-k-1)*A(q,j);
      for(i=k+1;i<N;i++){</pre>
        A(i,j) = A(i,j) - gamma*(*w)(i-k-1);
      }
    }
    for(i=0;i<N;i++){
      gamma = 0.0;
      for(q=k+1;q<N;q++)
        gamma += beta*(*w)(q-k-1)*A(i,q);
      for(j=k+1; j<N; j++){</pre>
        A(i,j) = A(i,j) - gamma*(*w)(j-k-1);
    }
  }
  delete x;
  delete w;
```

}

Remark 2: Observe that we pass the reference to the matrix **A** (denoted by the "SCMatrix &A" in the argument list.) We do this because we want to replace the values in **A** with the new upper Hessenberg matrix. If we were to omit the "&" and hence pass by value, the modifications made to the matrix **A** within the function would be lost when the function returns to the calling program.

Remark 3: Inside the function, we dynamically allocate two new SCVectors and assign them to the two pointers w and x. To use the "()" operator associated with the SCVector class, we first use the unitary operator '*' to retrieve the object to which the pointer points. Hence, the expression (*w) yields the object to which the pointer w points. The extra parentheses around this expression are used to guarantee that the '*' is carried out before the "()" operator.

9.4 Preconditioned Conjugate Gradient Method - PCGM

9.4.1 Convergence Rate of CGM

We have already introduced the Conjugate Gradient Method (CGM) in section 4.1.7 for the linear system

$$Ax = b$$
.

By defining the residual of the k^{th} iteration

$$r_{\mathbf{k}} \equiv \mathbf{b} - \mathbf{A}\mathbf{x}_k,$$

the solution and search directions are computed from

$$\mathbf{x}_{k+1} = \mathbf{x}_k + \alpha_k \mathbf{p}_k \tag{9.12}$$

$$\mathbf{p}_{k+1} = \mathbf{r}_{k+1} + \beta_k \mathbf{p}_k \tag{9.13}$$

while the residual can also be computed iteratively, i.e.,

$$\mathbf{r}_{k+1} = \mathbf{r}_k - \alpha_k \mathbf{A} \mathbf{p}_k. \tag{9.14}$$

We can derive a three-term recurrence formula by substituting in equation (9.12) the vector \mathbf{p}_k from equation (9.13) and also using the residual definition, to obtain

$$\mathbf{x}_{k+1} = (1 + \gamma_k)\mathbf{x}_k + \alpha_k(\mathbf{b} - \mathbf{A}\mathbf{x}_k) - \gamma_k\mathbf{x}_{k-1}$$

where we have defined

$$\gamma_k \equiv \frac{\alpha_k \beta_{k-1}}{\alpha_{k-1}}.$$

This is sometimes referred to as the *Rutishauser* formula. Symmetry and orthogonality together lead to the familiar three-term *magic formula* as we have seen many times in this book!

We have mentioned in section 4.1.7 that the CG method is equivalent to minimizing a properly defined quadratic form. In fact, it can be proved that if \mathbf{s} is the exact solution, then the conjugate gradient iterate \mathbf{x}_k minimizes the norm $||\mathbf{s} - \mathbf{x}||_A$ over the Krylov subspace of dimension k. This space is defined based on powers of \mathbf{A} with the orthogonal directions, i.e.,

$$\mathcal{K}_k(\mathbf{A}, \mathbf{p}) = \operatorname{span} \{ \mathbf{p}, \mathbf{A}\mathbf{p}, \mathbf{A}^2\mathbf{p} \dots, A^{k-1}\mathbf{p} \}.$$

The Rutishauser formula is similar to the Lanczos three-term formula, see section 10.3.6, and so CGM and Lanczos are related - they both use the same Krylov subspace and have three-term recurrence formulas. More specifically, a tridiagonal matrix \mathbf{T}_j can be constructed for CGM from

$$\mathbf{T}_j = \mathbf{P}^T \mathbf{A} \mathbf{P}.$$

Here $\mathbf{P} = \mathbf{R}\mathbf{D}^{-1}$, where \mathbf{D} is a diagonal matrix containing the magnitudes of residuals. Also, \mathbf{R} is the product of two matrices, the first formed by the columns of the orthogonal search directions \mathbf{p}_j while the second is a bidiagonal matrix with 1's in the diagonal and the scalars β_j above the main diagonal.

With regards to convergence, Reid [76] has observed that in practice CGM produces very good answers even before the total number of iterations reaches n, where $n \times n$ is the size of the symmetric positive-definite matrix \mathbf{A} . We recall that the fundamental theorem on conjugate directions, stated in section 4.1.7, guarantees that the exact solution will be achieved after n iterations. More specifically, if matrix \mathbf{A} has only $m \le n$ distinct eigenvalues then convergence will be achieved in m iterations. Round-off and corresponding loss of orthogonality is responsible for deviation from the theory, although round-off is not as severe as in the Lanczos method, as we discuss in section 10.3.6.

In practice, we always use a *stopping criterion* for convergence instead of having a loop with n iterations. To this end, it is important that the tolerance ϵ in the convergence test be proportional to the relative reduction of the initial residual or in some cases to be normalized with the right hand-side, e.g.,

$$|| r_{k+1} ||_2 \geq \epsilon || b ||_2$$
.

Here $\epsilon \approx 10^{-d}$ where d is the number of digits of desired accuracy. The computation of $\|\mathbf{r}_{k+1}\|^2 = (\mathbf{r}_{k+1}, \mathbf{r}_{k+1})$ requires no extra work as this quantity is used in the numerator of the formula for β_k . Also, we note that the residual $\|\mathbf{r}_{k+1}\|_2$ may not be decreasing monotonically although the solution error $\|\mathbf{s} - \mathbf{x}_k\|_2$ decreases monotonically. This is because in the minimization procedure the error of the solution is targeted directly but not the residual. Direct minimization of the residual is more common in solvers for non-symmetric systems, as we discuss in section 9.5. However, the \mathbf{A}^{-1} -residual norm, $(\mathbf{r}^T\mathbf{A}^{-1}\mathbf{r})$, decreases monotonically.

Because of finite arithmetic, the convergence rate of CGM is controlled by the condition number of matrix \mathbf{A} , which we denote by $\kappa_2(\mathbf{A})$. Specifically, the following estimate holds, (see [26, 46])

$$\frac{\parallel \mathbf{s} - \mathbf{x}_k \parallel_2}{\parallel \mathbf{s} - \mathbf{x}_0 \parallel_2} \le 2\gamma^k \sqrt{\kappa_2(\mathbf{A})},$$

where

$$\kappa_2(\mathbf{A}) = \frac{\lambda_{\max}}{\lambda_{\min}} \text{ and } \gamma = \frac{\sqrt{\kappa_2(\mathbf{A})} - 1}{\sqrt{\kappa_2(\mathbf{A})} + 1}.$$

For large values of κ_2 we have that $\gamma \to 1$, and thus the number of iterations for convergence of CGM is proportional to $\sqrt{\kappa_2(\mathbf{A})}$. For example, for the Poisson equation discretized on an N-by-N grid using second-order finite difference discretization (see chapter 6) we have that $\kappa_2(\mathbf{A}) \propto N^2$, and thus the number of iterations is proportional to N. The computational cost of CGM is then similar to SOR assuming for the latter an optimum relaxation parameter.

9.4.2 Preconditioners

In order to accelerate the convergence of CGM we employ preconditioners, see also 7.2.8. That is, we transform the linear system $\mathbf{A}\mathbf{x} = \mathbf{b}$ to

$$\mathbf{M}^{-1}\mathbf{A}\mathbf{x} = \mathbf{M}^{-1}\mathbf{b}$$

by multiplying by the preconditioner (non-singular) matrix M. We have already seen in section 7.2.8 some of the desired properties of the preconditioner: It should be:

• Spectrally close to matrix A so that the condition number

$$\kappa_2(\mathbf{M}^{-1}\mathbf{A}) \ll \kappa_2(\mathbf{A})$$

and also,

• Inexpensive to invert since the solution of $\mathbf{M}\mathbf{x} = \mathbf{b}$ will be required.

In addition, since we consider here a symmetric positive-definite matrix A, then M has to also be symmetric and positive-definite.

The objective is to modify the original CG algorithm only slightly in order to "correct" the search directions but not to increase the computational complexity significantly. To this end, we first need to symmetrize the preconditioned system as $\mathbf{M}^{-1}\mathbf{A}$ is not a symmetric matrix. We then express \mathbf{M} in terms of its eigenvectors and eigenvalues, i.e.,

$$\mathbf{M} = \mathbf{V}\Lambda\mathbf{V}^T \Rightarrow \mathbf{M}^{1/2} \equiv \mathbf{V}\Lambda^{1/2}\mathbf{V}^T.$$

Next we multiply $\mathbf{M}^{-1}\mathbf{A}\mathbf{x} = \mathbf{M}^{-1}\mathbf{b}$ by $\mathbf{M}^{1/2}$ to arrive at

$$(\mathbf{M}^{-1/2}\mathbf{A}\mathbf{M}^{-1/2})(\mathbf{M}^{1/2}\mathbf{x}) = \mathbf{M}^{-1/2}\mathbf{b}$$

$$\Rightarrow \mathbf{B}\mathbf{y} = \mathbf{f}, \text{ where } \mathbf{B} \equiv \mathbf{M}^{-1/2} \mathbf{A} \mathbf{M}^{-1/2},$$

and,

$$\mathbf{y} \equiv \mathbf{M}^{1/2} \mathbf{x}$$
 and $\mathbf{f} \equiv \mathbf{M}^{-1/2} \mathbf{b}$.

The above defines the new system (i.e., $\mathbf{B}, \mathbf{y}, \mathbf{f}$) to be solved using the original CGM. We note that

$$\mathbf{M}^{-1/2}\mathbf{B}\mathbf{M}^{1/2} = \mathbf{M}^{-1/2}(\mathbf{M}^{-1/2}\mathbf{A}\mathbf{M}^{-1/2})\mathbf{M}^{1/2} = \mathbf{M}^{-1}\mathbf{A}$$

and thus \mathbf{B} and $\mathbf{M}^{-1}\mathbf{A}$ are similar so they have the same eigenvalues.

The following PCG algorithm is derived by applying the standard CG algorithm to the new system

$$\mathbf{B}\mathbf{v} = \mathbf{f}$$
.

as defined above. The important thing is that we do not need to explicitly take the square root of M – this would have been costly!

Preconditioned Conjugate Gradient Algorithm

- Initialize:
 - Choose $\mathbf{x}_0 \Rightarrow \mathbf{r}_0 = \mathbf{b} \mathbf{A}\mathbf{x}_0$
 - Solve $\mathbf{M}\tilde{\mathbf{r}}_0 = \mathbf{r}_0 \Rightarrow \mathbf{p}_0 = \tilde{\mathbf{r}}_0$

• Begin Loop: for k = 1, ...

$$\begin{array}{ccccc} \alpha_k & = & \frac{\left(\tilde{\mathbf{r}}_k, \mathbf{r}_k\right)}{\left(\mathbf{p}_k, \mathbf{A}\mathbf{p}_k\right)} \\ \mathbf{x}_{k+1} & = & \mathbf{x}_k + \alpha_k \mathbf{p}_k \\ \mathbf{r}_{k+1} & = & \mathbf{r}_k - \alpha_k \mathbf{A}\mathbf{p}_k \\ \mathbf{M}\tilde{\mathbf{r}}_{k+1} & = & \mathbf{r}_{k+1} \\ & \text{If} & \left(\tilde{\mathbf{r}}_{k+1}, \mathbf{r}_{k+1}\right) \leq \epsilon \\ & \text{If} & \left(\mathbf{r}_{k+1}, \mathbf{r}_{k+1}\right) \leq \epsilon \\ & \text{return} \\ & \beta_k & = & \frac{\left(\tilde{\mathbf{r}}_{k+1}, \mathbf{r}_{k+1}\right)}{\left(\tilde{\mathbf{r}}_k, \mathbf{r}_k\right)} \\ & \mathbf{p}_{k+1} & = & \tilde{\mathbf{r}}_{k+1} + \beta_k \mathbf{p}_k \end{array}$$

endfor

• End Loop

Remark 1: The *stopping criterion* is based on the actual residual \mathbf{r} and not on the modified residual $\tilde{\mathbf{r}}$. However, checking the latter first saves some computational time as it is readily available while the former requires explicit calculation.

The question is still open as to what is the *best preconditioner* since in most cases it is problem-dependent. For matrices with diagonal elements that are very different in magnitude, using

$$\mathbf{M} = diag(a_{11}, a_{22}, \dots, a_{nn})$$

is very effective as it reduces the condition number of \mathbf{B} by a factor of n of its minimum value [26]. This is called *diagonal scaling* and corresponds to Jacobi preconditioning. An extension of this idea is to build \mathbf{M} as a block-diagonal matrix out of block submatrices of \mathbf{A} . Similarly, the Gauss-Seidel method can be used as preconditioner but it needs to be symmetrized first. To this end, the symmetric SOR (SSOR) we presented in section 7.2.6 can be a more effective block preconditioner.

Diagonal scaling fails if all diagonal elements are equal such as in the matrices resulted from finite difference discretization of diffusion problems on uniforms grids, see chapter 7, or other Toeplitz type matrices (see next section). One of the most effective and popular preconditioners for such problems is based on the **incomplete Cholesky factorization** of the matrix $\bf A$. We have already presented the Cholesky factorization of $\bf A$ in section 9.2, which is feasible for symmetric positive-definite matrices, like the ones we consider here. The problem with Cholesky is that it fills-in the zero entries of $\bf A$ in the LL^T decomposition, and in some cases it may totally destroy the possible sparsity initially present in $\bf A$.

To obtain an incomplete Cholesky factorization of $\bf A$ we can simply suppress the fill-in entries, i.e., zeroing out the entries corresponding to the zero entries of the original matrix. Alternatively, to avoid computing these entries in $\bf L$ and simply place zeros at the corresponding locations, we can modify the Cholesky algorithm so that

• if $a_{ij} \neq 0$, compute l_{ij} ,

• elseif $l_{ij} = 0$.

We note that the two approaches are not the same as can be easily verified in the following example. Let us consider the (4×4) matrix

$$\mathbf{A} = \begin{bmatrix} 4 & -1 & -1 & 0 \\ -1 & 4 & 0 & -1 \\ -1 & 0 & 4 & -1 \\ 0 & -1 & -1 & 4 \end{bmatrix}.$$

We first perform a standard Cholesky decomposition, which yields the following triangular matrix

$$\mathbf{L}_c = \begin{bmatrix} 2.0000 & 0 & 0 & 0 \\ -0.5000 & 1.9365 & 0 & 0 \\ -0.5000 & -0.1291 & 1.9322 & 0 \\ 0 & -0.5164 & -0.5521 & 1.8516 \end{bmatrix}.$$

We note that the (3,2) entry which was initially zero in **A** has become non-zero in \mathbf{L}_c . The first version of incomplete Cholesky would then be:

$$\mathbf{L}_{i1} = \begin{bmatrix} 2.0000 & 0 & 0 & 0 \\ -0.5000 & 1.9365 & 0 & 0 \\ -0.5000 & 0 & 1.9322 & 0 \\ 0 & -0.5164 & -0.5521 & 1.8516 \end{bmatrix}.$$

However, the second version, where we don't compute the l_{ij} at all if the corresponding $a_{ij} = 0$, is

$$\mathbf{L}_{i2} = \begin{bmatrix} 2.0000 & 0 & 0 & 0 \\ -0.5000 & 1.9365 & 0 & 0 \\ -0.5000 & 0 & 1.9365 & 0 \\ 0 & -0.5164 & -0.5164 & 1.8619 \end{bmatrix},$$

which is different than L_{i1} in the entries (3,3), (4,3) and (4,4).

In practice, we can replace the second code statement above by setting a *threshold value* on the Cholesky entry as follows

• elseif $|l_{ij}| \le \epsilon$ then $l_{ij} = 0$,

so that only significantly large entries are retained. Clearly, the incomplete Cholesky decomposition can be performed only initially, i.e., before the iteration loop, and store $\mathbf{M} = \mathbf{L}\mathbf{L}^T$ or simply store \mathbf{L} .

Remark 2: While Cholesky factorization of a symmetric positive-definite matrix is always feasible, incomplete factorization may not be possible in some cases. This complication may manifest itself as a square root of a negative number to compute the l_{ii} . In this case, a positive large number should over-write the appropriate element, which can be chosen either arbitrarily or as a sum of the adjoint diagonal entries or even as a sum of absolute values of the rest of the entries in the row; the latter will ensure diagonal dominance.

Remark 3: Domain decomposition is another way of preconditioning partial differential equations. The idea is to break up the domain of interest to subdomains, which can also overlap, and subsequently solve the PDE approximately but fast in each of the subdomains. This can be also done independently for each domain in an embarrassingly parallel fashion. The preconditioner matrix M is constructed by piecing together the solutions to subproblems leading to a block diagonal M if the subdomains do not overlap or a product of block diagonal submatrices if the subdomains overlap.

9.4.3 Toeplitz Matrices and Circulant Preconditioners

A special preconditioner is very effective for Toeplitz matrices, the circulant matrix. Topeplitz matrices have constant diagonals and are encountered in signal processing and in a wide range of other problems which are invariant in time and space. A circular matrix \mathbf{C} is a Toeplitz matrix but it is entries also satisfy

$$c_k = c_{k+n}$$

where $n \times n$ is the size of the matrix \mathbf{C} .

A general Toeplitz matrix has (2n-1) independent entries, which are determined by the first row and column. A general circulant matrix on the other hand has only n independent entries. Assuming symmetry, then a Toeplitz matrix has n independent entries (the first column) whereas a circulant matrix has $\lfloor n/2 \rfloor + 1$. More clearly, the differences between a general Toeplitz and a general circulant matrix are shown below.

$$\mathbf{A} = \begin{bmatrix} a_0 & a_{-1} & . & . & a_{1-n} \\ a_1 & a_0 & a_{-1} & . & . \\ . & a_1 & a_0 & . & . \\ . & . & . & . & a_{-1} \\ a_{n-1} & . & . & a_1 & a_0 \end{bmatrix},$$

and

$$\mathbf{C} = \begin{bmatrix} c_0 & c_{n-1} & . & . & c_1 \\ c_1 & c_0 & c_{n-1} & . & . \\ . & c_1 & c_0 & . & . \\ . & . & . & . & c_{n-1} \\ c_{n-1} & . & . & c_1 & c_0 \end{bmatrix}.$$

Assuming we want to solve the system $\mathbf{A}\mathbf{x} = \mathbf{b}$ where \mathbf{A} is a symmetric Toeplitz matrix, the suggestion, first proposed in [83], is to precondition it with a corresponding circulant matrix. To this end, we recall that in most applications the main diagonal and its neighbors are strongly dominant and thus we can use them to construct an appropriate circulant preconditioner \mathbf{C} . The entries $c_1 = a_1$ appear both next to the main diagonal but also in the extreme corners as shown above. It turns out that these relatively large entries control the eigenspectrum of $\mathbf{C}^{-1}\mathbf{A}$, which in turn controls the convergence rate of PCGM. The rest of the eigenvalues are clustered around one; this is demonstrated in homework problems, see 10.7. Typically, we copied a few of the main diagonals from the Toeplitz matrix onto the circulant matrix but of course not all! In the homework problems of section 9.8 we ask

you to experiment with different circulant preconditioners. As you can see, the speed-up is substantial!

9.4.4 Parallel PCGM

The parallelization of the preconditioned conjugate gradient is fairly straight-forward. Assuming that the matrix **A** has been distributed by rows across processes, the conjugate gradient component of the algorithm can be accomplished with only four MPI calls: three calls to MPI_Allreduce to accomplish dot products, and one call to MPI_Allgather. We can eliminate one of the reduction calls if we decide to use the dot product of the residual with the modified residual for the stopping criterion. Depending on the choice of preconditioner, however, additional MPI calls may be required to accomplish the preconditioning. In the case of a diagonal preconditioner, no MPI calls are necessary since diagonal preconditioning can be accomplished locally on all rows contained in a process. Other preconditioners such as incomplete Cholesky may require additional MPI calls, the cost of which should be considered when determining what preconditioner to use. In figure 9.14 we provide a schematic of the iterative part of the parallel PCG algorithm with annotations denoting what BLAS and MPI operations should be used. Note that in figure 9.14 we use the modified residual for the stopping criterion, and hence we only have three MPI calls.

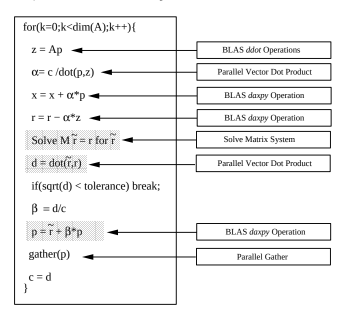


Figure 9.14: Iterative part of the preconditioned conjugate gradient (PCG) algorithm.



We now present an MPI program demonstrating the preconditioned conjugate gradient method with a diagonal preconditioner. As a sample problem, we will solve (using second-order finite differences) the following equation:

$$\frac{d^2u(x)}{dx^2} - c_2e^{c_1(x-0.5)^2}u(x) = -\sin(2\pi x)(4\pi^2 - c_2e^{c_1(x-0.5)^2})$$
(9.15)

in the interval $x \in [0, 1]$ with boundary conditions u(0) = u(1) = 0 and constants $c_1 = 20.0$ and $c_2 = 1000.0$. The exact solution is $u(x) = \sin(2\pi x)$.

We have chosen the constants c_1 and c_2 so that there is a large disparity in the values along the diagonal. This will allow us to see the difference between conjugate gradient (CGM) and preconditioned conjugate gradient (PCGM) with a diagonal preconditioner.

Since we have chosen to use a second-order finite difference method to approximate the derivative operator, we expect that the error should decrease by a factor of four when we double the number of grid points used. In figure 9.15 we show a log-log plot of the L_2 error versus the number of grid points used. The magnitude of the slope of the line is approximately two, consistent with the fact that our approximation is second order.

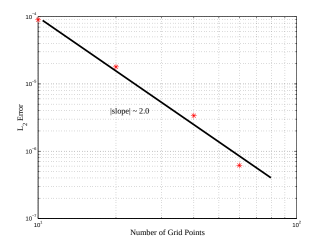


Figure 9.15: L_2 error versus the number of grid points for the PCGM example defined in equation (9.15).

In figure 9.16 we plot the inner product of the residual (r, r) for both the CGM and the PCGM. As predicted, PCGM converges much faster than the standard CGM.

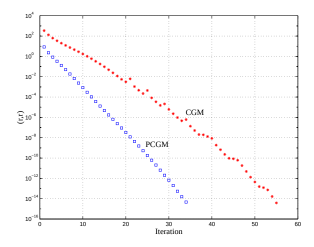


Figure 9.16: Inner product of the residual for both the CG and PCG methods applied to the PCGM example defined in equation (9.15).

To better explain the code, we have broken the entire program into four parts, labeled part one through part four. The four parts are as follows:

- 1. Part 1 MPI initialization, initial memory allocation, and generation of grid and right-hand-side vector.
- 2. Part 2 Memory allocation and generation of the matrix **A**.
- 3. Part 3 PCGM initialization.
- 4. Part 4 PCGM main iteration loop.

For each part, we will first present the code and then present a collection of remarks elucidating the salient points within each part.

Part 1 - MPI initialization

```
#include <iostream.h>
#include <iomanip.h>
#include "SCmathlib.h"
#include<mpi.h>
const int rows_per_proc = 40;
const double c1 = 20.0;
const double c2 = 1000.0;
const double tol = 1.0e-14;
int main(int argc, char *argv[]){
  int i, j, k;
  int mynode, totalnodes, totalsize, offset;
  MPI_Status status;
  double sum,local_sum,c,d,alpha,beta;
  double ** A, *q, *x, *grid;
  double *p,*z,*r,*mr;
  MPI_Init(&argc,&argv);
  MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
  MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
  totalsize = totalnodes*rows_per_proc;
  p = new double[totalsize];
  z = new double[rows_per_proc];
  r = new double[rows_per_proc];
 mr = new double[rows_per_proc];
```

```
x = new double[rows_per_proc];
q = new double[rows_per_proc];
grid = new double[rows_per_proc];

double dx = 1.0/(totalsize+1);
for(i=0;i<rows_per_proc;i++){
   grid[i] = dx*(1+rows_per_proc*mynode+i);
   q[i] = -dx*dx*sin(2.0*M_PI*grid[i])*
      (-4.0*M_PI*M_PI - c2*exp(c1*(grid[i]-0.5)*(grid[i]-0.5)));
   x[i] = 1.0;
}</pre>
```

Remark 1: We have four global variables in this program. The variable *rows_per_proc* gives the number of rows per processor. In this program we have decomposed the matrix by associating rows to processors. The two constants *c1* and *c2* are specific to the problem we are solving. The last global constant variable *tol* specifies the tolerance to which we should converge.

Remark 2: Observe that with the exception of the array p all other arrays need only be of size $rows_per_proc$ and not of size totalsize. Each processor need only maintain its part of the residual, modified residual and solution vector; however, each processor must have a copy of the entire p vector.

Part 2 - Memory allocation and generation of matrix

```
/* Part 2 */
A = new double*[rows_per_proc];
for(i=0;i<rows_per_proc;i++){</pre>
  A[i] = new double[totalsize];
  for(j=0;j<totalsize;j++)</pre>
    A[i][j] = 0.0;
}
if(mynode==0){
  A[0][0] = 2.0 + dx*dx*c2*exp(c1*(grid[0]-0.5)*(grid[0]-0.5));
  A[0][1] = -1.0;
  for(i=1;i<rows_per_proc;i++){</pre>
    A[i][i] = 2.0 + dx*dx*c2*exp(c1*(grid[i]-0.5)*
               (grid[i]-0.5));
    A[i][i-1] = -1.0;
    A[i][i+1] = -1.0;
 }
}
```

```
else if(mynode == (totalnodes-1)){
  A[rows_per_proc-1][totalsize-1] = 2.0 +
      dx*dx*c2*exp(c1*(grid[rows_per_proc-1]-0.5)*
      (grid[rows_per_proc-1]-0.5));
  A[rows_per_proc-1][totalsize-2] = -1.0;
  for(i=0;i<rows_per_proc-1;i++){</pre>
    offset = rows_per_proc*mynode + i;
    A[i][offset] = 2.0 + dx*dx*c2*exp(c1*(grid[i]-0.5)*
                   (grid[i]-0.5));
    A[i][offset-1] = -1.0;
    A[i][offset+1] = -1.0;
  }
}
else{
  for(i=0;i<rows_per_proc;i++){</pre>
    offset = rows_per_proc*mynode + i;
    A[i][offset] = 2.0 + dx*dx*c2*exp(c1*(grid[i]-0.5)*(grid[i]-0.5));
    A[i][offset-1] = -1.0;
    A[i][offset+1] = -1.0;
  }
}
```

Remark 3: We break the matrix setup into three cases. We have to carefully handle the first and last processors because they contain the first and last rows, respectively.

Part 3 - PCGM initialization

```
MPI_DOUBLE,MPI_COMM_WORLD);
```

Remark 4: We have chosen our initial vector as the vector of all ones. Because we have to calculate the initial residual, we require that all processors have the entire initial guess. Notice that we temporarily use the p array to accomplish this instead of allocating a new vector. Because p is not in use until after the modified residual is calculated, we can use the allocated space with no adverse effect.

Part 4 - PCGM main iteration loop

```
/* Part 4 */
for(k=0;k<totalsize;k++){</pre>
   for(i=0;i<rows_per_proc;i++)</pre>
     z[i] = dot(totalsize,A[i],p);
   local_sum = dot(rows_per_proc,z,p+offset);
  MPI_Allreduce(&local_sum,&sum,1,MPI_DOUBLE,MPI_SUM,
                 MPI_COMM_WORLD);
   alpha = c/sum;
   for(i=0;i<rows_per_proc;i++){</pre>
     x[i] = x[i] + alpha*p[offset+i];
     r[i] = r[i] - alpha*z[i];
   }
   /* Preconditioning Stage */
   for(i=0;i<rows_per_proc;i++)</pre>
     mr[i] = r[i]/A[i][offset+i];
   local_sum = dot(rows_per_proc,mr,r);
  MPI_Allreduce(&local_sum,&sum,1,MPI_DOUBLE,MPI_SUM,
                 MPI_COMM_WORLD);
  d = sum; //contains inner product of
            //residual and modified residual
   local_sum = dot(rows_per_proc,r,r);
  MPI_Allreduce(&local_sum,&sum,1,MPI_DOUBLE,MPI_SUM,
                 MPI_COMM_WORLD);
```

```
//sum now contains inner product of residual and residual
    if(mynode == 0){
      cout << k << "\t" << "dot(mr,r) = " << d << "\t";
      cout << "dot(r,r) = " << sum << endl;
    }
    if(fabs(d) < tol) break;
    if(fabs(sum) < tol) break;</pre>
    beta = d/c;
    for(i=0;i<rows_per_proc;i++)</pre>
      z[i] = mr[i] + beta*p[i+offset];
    MPI_Allgather(z,rows_per_proc,MPI_DOUBLE,p,rows_per_proc,
                   MPI_DOUBLE,MPI_COMM_WORLD);
    c = d;
 }
  delete[] p;
  delete[] z;
  delete[] r;
  delete[] mr;
  delete[] x;
  delete[] q;
  delete[] grid;
  for(i=0;i<rows_per_proc;i++)</pre>
    delete[] A[i];
  delete[] A;
 MPI_Finalize();
}
```

Remark 5: Observe that we only require four MPI calls: three $MPI_Allreduce$ and one $MPI_Allgather$. The three reductions are used to obtain the inner product across all processors; the Allgather is used so that all processors have the updated value of p.

Remark 6: Because we are using diagonal preconditioning, no additional communication is required for the preconditioner. If we were to use incomplete Cholesky as a preconditioner, additional communication similar to what was accomplished in the parallel Gaussian elimination solver would have to be used. The factorization would be accomplished before the

iteration loop, and only the cost of two parallel back solves would be incurred within the iteration loop.

9.5 Non-Symmetric Systems

The conjugate gradient algorithm assumes that the matrix A is symmetric and positive-definite. To deal with the non-symmetric system

$$\mathbf{A}\mathbf{x} = \mathbf{b}$$

we need to come up with new solvers or transform the non-symmetric system into a symmetric one as follows

$$\mathbf{A}^T \mathbf{A} \mathbf{x} = \mathbf{A}^T \mathbf{b}$$
 (CGNR).

Clearly, this transformation produces a symmetric positive-definite matrix $\mathbf{B} \equiv \mathbf{A}^T \mathbf{A}$ and thus we can apply PCGM to it. This is called the CGNR (conjugate gradient normal residual) algorithm and it is an acceptable method for matrices which are well-conditioned, although we have to pay extra to include matrix-vector multiplies with both \mathbf{A}^T and \mathbf{A} . On the other hand, the condition number of \mathbf{B} is the *square* of the condition number of \mathbf{A} and thus for ill-conditioned matrices an extremely large number of iterations is required for convergence.

We now compare the CG and CGNR algorithms by considering what quantity is that we *minimize* in these minimization-based solvers. We have already seen in section 4.1.7 that the solution with CG is equivalent to minimizing the quadratic form

$$P_{CG}(\mathbf{x}) = \frac{1}{2}(\mathbf{x}, \mathbf{A}\mathbf{x}) - (\mathbf{b}, \mathbf{x}).$$

We can re-write the above form in terms of the error $\mathbf{e} \equiv \mathbf{x} - \mathbf{s}$, where \mathbf{s} is the exact solution, as follows

$$2P_{CG}(\mathbf{x}) = (\mathbf{x}, \mathbf{A}\mathbf{x}) - 2(\mathbf{x}, \mathbf{A}\mathbf{s}) + (\mathbf{s}, \mathbf{A}\mathbf{s})$$

= $\mathbf{e}^{T}\mathbf{A}\mathbf{e}$
= $||\mathbf{e}||_{A}$.

The above simply states that searching for the solution using the CG algorithm is equivalent to minimizing the *error* in the A-norm. In contrast, the CGNR algorithm minimizes the residual in the L_2 -norm, and thus the corresponding quadratic form is

$$2P_{CGNR}(\mathbf{x}) = ||\mathbf{b} - \mathbf{A}\mathbf{x}||_{2}$$
$$= (\mathbf{b} - \mathbf{A}\mathbf{x}, \mathbf{b} - \mathbf{A}\mathbf{x})$$
$$= (\mathbf{A}\mathbf{x}, \mathbf{A}\mathbf{x}) - 2(\mathbf{b}, \mathbf{A}\mathbf{x}) + (\mathbf{b}, \mathbf{b}).$$

Therefore, the minimization of P_{CGNR} corresponds to applying CG to the system

$$\mathbf{B}\mathbf{x} = \mathbf{f}$$

where $\mathbf{B} = \mathbf{A}^T \mathbf{A}$ and $\mathbf{f} = \mathbf{A}^T \mathbf{b}$.

There are several conjugate residual algorithms in the literature, which are based on the minimization of the residual in different forms. A significant difference between such algorithms and the CG for symmetric systems is that the loss of symmetry results in the loss of the three-term magic recurrence formula, which keeps things sparse and leads to efficiency both in memory and cost. Instead of two or three vectors, in the non-symmetric conjugate algorithms we typically need to store the entire sequence of conjugate directions, which is prohibitively expensive for very large n. However, in practice a subset of the last k vectors may work or a restart of the algorithm after k steps can also be used.

In the following, instead of presenting all variants of conjugate gradient residual algorithms we concentrate on one that makes use of an iterative procedure to simplify the original matrix, namely the *Arnoldi iteration*. Based on this procedure, we can solve, relatively efficiently, non-symmetric systems as we demonstrate below. We can also compute eigenvalues of non-symmetric matrices, as we discuss in section 10.5 in the next chapter.

9.5.1 The Arnoldi Iteration

The Arnoldi method is an orthogonal projection onto a Krylov subspace \mathcal{K}_m for non-symmetric matrices $\mathbf{A}(n \times n)$; usually $m \ll n$. It reduces the matrix \mathbf{A} to a Hessenberg form, that is it accomplishes what Householder does for the entire matrix \mathbf{A} but here we have the option of only going half-way. Arnoldi, who introduced this algorithm in the early 1950s [3], suggested that it leads to good approximations for some of the eigenvalues of \mathbf{A} even if we terminate prematurely! In practice, it is a useful technique for obtaining the eigenvalues of large sparse matrices. It is useful to think of this as an extension of Lanczos method (see section 10.3.6) to non-symmetric matrices. It has the same iterative and approximate form - although a separate version of Lanczos exists for non-symmetric matrices [79].

Let us begin by considering the similarity transformation

$$\mathbf{A} = \mathbf{V}^T \mathbf{H} \mathbf{V}.$$

where we define the matrix V or V_m with orthonormal columns

$$\mathbf{V}_m = [\mathbf{v}_1 \,|\, \mathbf{v}_2 \,|\, \dots \,|\, \mathbf{v}_m\,],$$

and \mathbf{H}_m is the Hessenberg matrix $(m \times m)$. We also define the extended matrix $\tilde{\mathbf{H}}_m$ of dimension $(m+1) \times m$, as follows:

$$\tilde{\mathbf{H}}_{m} = \begin{bmatrix} h_{11} & h_{12} & \dots & h_{1m} \\ h_{21} & h_{22} & \dots & h_{2m} \\ & \ddots & \ddots & \vdots \\ & & h_{m,m-1} & h_{mm} \\ & & & h_{m+1,m} \end{bmatrix},$$

and we can write

$$\tilde{\mathbf{H}}_m = \mathbf{H}_m + h_{m+1,m} \mathbf{v}_{m+1} \mathbf{e}_m^T,$$

where \mathbf{v}_{m+1} , defined as orthonormal, i.e., it has unit norm.

The Arnoldi iteration process satisfies:

$$\mathbf{AV}_m = \mathbf{V}_{m+1}\tilde{\mathbf{H}}_m, \tag{9.16}$$

and correspondingly, the $m^{\rm th}$ column of this equation reads:

$$\mathbf{A}\mathbf{v}_m = h_{1m}\mathbf{v}_1 + h_{2m}\mathbf{v}_2 + \ldots + h_{mm}\mathbf{v}_m + h_{m+1,m}\mathbf{v}_{m+1},$$

which is an (m+1)-term recurrence formula, instead of the three-term formula in symmetric systems.

The basic algorithm of Arnoldi iteration implements the above formula in a straightforward manner, as follows:

Basic Arnoldi Algorithm

Initialize: Choose a vector
$$\mathbf{x}_0 \Rightarrow \mathbf{v}_1 = \frac{\mathbf{x}_0}{\|\mathbf{x}_0\|}$$

Begin Loop: for $j = 1, 2, \dots m$

for $i = 1, \dots, j$
 $h_{ij} = (\mathbf{A}\mathbf{v}_j, \mathbf{v}_i)$

endfor

 $\mathbf{w}_j = \mathbf{A}\mathbf{v}_j - \sum_{i=1}^j h_{ij}\mathbf{v}_i$
 $h_{j+1,j} = \|\mathbf{w}_j\|_2$
 $\mathbf{v}_{j+1} = \mathbf{w}_j/h_{j+1,j}$

End Loop:

The above algorithm is basically the standard Gram-Schmidt orthogonalization procedure applied to the Krylov space \mathcal{K}_m .

endfor

We note that in each iteration we compute the entire j column, including the entry $h_{j+1,j}$ below the main diagonal. In particular, if $h_{j+1,j} = 0$ then the Arnoldi iteration stalls, but that is in fact good news! This is because such a breakdown usually implies that we have achieved convergence and the iteration should be terminated. If we want to continue, we can restart the process with a new orthonormal vector \mathbf{v}_{j+1} , which can be selected arbitrarily.

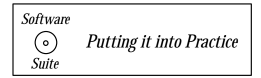
The implementation of the Arnoldi algorithm given above although straightforward it suffers from round-off errors, just as is the case of the basic Gram-Schmidt algorithm. To this end, we can apply the *more stable* modified Gram-Schmidt orthogonalization procedure to the Krylov space to come up with a better code for the Arnoldi iteration, as follows:

Modified Arnoldi Algorithm

Initialize: Choose
$$\mathbf{x}_0 \Rightarrow \mathbf{v}_1 = \frac{\mathbf{x}_0}{\|\mathbf{x}_0\|_2}$$
.

 $Begin\ Loop: \ \text{for}\ j=1,\ldots,m$
 $\mathbf{w} = \mathbf{A}\mathbf{v}_j$
 $\text{for}\ i=1,\ldots,j$
 $h_{ij} = (\mathbf{w},\mathbf{v}_i)$
 $\mathbf{w} = \mathbf{w} - h_{ij}\mathbf{v}_i$
 endfor
 $h_{j+1,j} = \|\mathbf{w}\|_2$
 $\mathbf{v}_{j+1} = \mathbf{w}/h_{j+1,j}$
 $End\ Loop: \ \text{endfor}$

This is a much more stable algorithm but even this version needs further treatment sometimes, so an extra orthogonalization may be required occasionally. For robustness, we can apply the Householder algorithm periodically for extra orthogonalization.



Below we present a function for computing the Arnoldi decomposition using the modified algorithm presented earlier. The function takes four arguments: the integer value m denoting the dimension of the Krylov space on to which we are projecting, a SCMatrix A of size $N \times N$ on which the decomposition is to be accomplished, and two SCMatrix variables H and V for storing the resulting $\tilde{\mathbf{H}}$ and $\tilde{\mathbf{V}}$ from the decomposition. Note that H must be a SCMatrix of size $(m+1) \times m$ and V must be a SCMatrix of size $N \times (m+1)$. The initial direction vector x defaults to the first unit vector.

```
w = A*v;
for(int i=0;i<=j;i++){
    V.GetColumn(i,v);
    H(i,j) = dot(w,v);
    w = w - H(i,j)*v;
}
H(j+1,j) = w.Norm_12();
v = w/H(j+1,j);
V.PutColumn(j+1,v);
}</pre>
```

Remark 1: Once again we pass all three SCMatrix variables by reference (denoted by the '&' symbol) as opposed to by the default case of passing by value. For the variables H and V, the reason is as before - we want to change the values within the matrices, and we want those changes to remain valid after the function has returned (i.e., not be lost in the "pass by value" copy of the variable discarded when the function returns). However, in the case of the matrix A, this is not the case. We do not want to modify the values of A. Why then do we pass by reference as opposed to passing by value? In this case, we do so because we assume that A is very large, and therefore we do not wish to allocate space for a copy of A when the function is called. We instead pass the matrix by reference so that no additional memory must be allocated; the original memory from the calling function is used. We do, however, want to guarantee that the matrix A does not change within the function, hence why we add the **const** in the appropriate place within the argument list. The "const SCMatrix &A" allows us to pass the matrix A by reference, but not to update its values within the function ModifiedArnoldi.

Remark 2: We introduced two new SCMatrix methods:

- 1. SCMatrix::PutColumn(int col, const SCVector &v), and
- 2. SCMatrix::GetColumn(int col, SCVector &v).

The first function copies the contents of the vector v into the col column of the matrix. The second function copies the contents of the col column from the matrix into the vector v.

Example: As an example of the use of the code above, we revisit our old friend the Hilbert matrix of size three:

$$\mathbf{A} = \begin{bmatrix} 1 & 1/2 & 1/3 \\ 1/2 & 1/3 & 1/4 \\ 1/3 & 1/4 & 1/5 \end{bmatrix}.$$

The goal is to find the decomposition when m=2. We input into our function m=2, the Hilbert matrix \mathbf{A} as given above, and two SCMatrix variables H and V which are of size 3×2 and 3×3 respectively. As output we obtain $\tilde{\mathbf{H}}$ and $\tilde{\mathbf{V}}$ contained within the SCmatrix variables H and V respectively. The decomposition for the 3×3 Hilbert matrix given above is:

$$\tilde{\mathbf{V}} = \begin{bmatrix} 1.0000 & 0.0000 & 0.0000 \\ 0.0000 & 0.8321 & -0.5547 \\ 0.0000 & 0.5547 & 0.83205 \end{bmatrix}$$

and

$$\tilde{\mathbf{H}} = \left[\begin{array}{ccc} 1.00000 & 0.60093 \\ 0.60093 & 0.52308 \\ 0.00000 & 0.03462 \end{array} \right] \, .$$

9.5.2 GMRES

GMRES stands for "generalized minimum residual" and is one of the most effective solvers for non-symmetric systems

$$\mathbf{A}\mathbf{x} = \mathbf{b}$$
,

where **A** is an $n \times n$ square but non-symmetric matrix; it is also assumed non-singular since we look for the solution $\mathbf{A}^{-1}\mathbf{b}$.

The main idea of GMRES is to minimize the residual $\|\mathbf{b} - \mathbf{A}\mathbf{x}_m\|_2$ at the m^{th} iteration. Specifically, \mathbf{x}_m is a vector in the Krylov space

$$\mathcal{K}_m = \{\mathbf{v}, \mathbf{A}\mathbf{b}, \dots, \mathbf{A}^{m-1}\mathbf{b}\}$$

and it can be determined by solving a least-squares problem; here m < n. At the point m = n we are attempting to minimize the residual $\|\mathbf{b} - \mathbf{A}\mathbf{x}_n\|_2$ and hence obtain the solution \mathbf{x} .

Such a minimization is equivalent to performing a QR decomposition to the matrix of least-squares coefficients \mathbf{C} (see section 3.1.7). We illustrate this point next.

Let us assume that in matrix form we seek to find

$$\parallel \mathbf{Ca} - \mathbf{f} \parallel_2 = \min \mathbf{mum}$$

with respect to **a**. Then, we need to QR-decompose **C** which is a non-square matrix $p \times q$, p > q.

The idea is to apply Householder transformation to the extended matrix \mathbf{C} of $p \times p$ but stop when we have completed zeroing out the entries in the first q columns. We then write:

$$\mathbf{C} = \mathbf{Q} \left[\begin{array}{c} \mathbf{R} \\ 0 \end{array} \right],$$

where **R** is a $q \times q$ upper triangular matrix. Since $\|\mathbf{Q}\| = \|\mathbf{Q}^{-1}\| = 1$, we also have

$$\parallel \mathbf{Ca} - \mathbf{f} \parallel_2 = \parallel \mathbf{Q}^{-1}(\mathbf{Ca} - \mathbf{f}) \parallel_2 = \parallel \tilde{\mathbf{R}}\mathbf{a} - \tilde{\mathbf{f}} \parallel_2,$$

where we have defined $\tilde{\mathbf{R}} = \begin{bmatrix} \mathbf{R} \\ 0 \end{bmatrix}$ and also $\tilde{\mathbf{f}} = \mathbf{Q}^{-1}\mathbf{f}$.

But

$$\tilde{\mathbf{R}}\mathbf{a} - \tilde{\mathbf{f}} = \begin{bmatrix} \mathbf{R} \\ 0 \end{bmatrix} \mathbf{a} - \begin{bmatrix} \tilde{\mathbf{f}}_1 \\ \tilde{\mathbf{f}}_2 \end{bmatrix} = \begin{bmatrix} \mathbf{R}\mathbf{a} - \tilde{\mathbf{f}}_1 \\ -\tilde{\mathbf{f}}_2 \end{bmatrix}$$

where we split $\tilde{\mathbf{f}}$ in $\tilde{\mathbf{f}}_1$ of length q and $\tilde{\mathbf{f}}_2$ of length (p-q). Then,

$$\| \mathbf{Ca} - \mathbf{f} \|_{2}^{2} = \| \mathbf{Ra} - \tilde{\mathbf{f}}_{1} \|_{2}^{2} + \| \tilde{\mathbf{f}}_{2} \|_{2}^{2}.$$

Therefore, the least-squares problem is equivalent to solving

$$\mathbf{Ra} = \tilde{\mathbf{f}}_1$$
,

which minimizes the entire residual, since $\tilde{\mathbf{f}}_2$ is fixed with respect to \mathbf{a} .

The problem, however, in applying directly this QR procedure to

$$\|\mathbf{A}\mathbf{x}_m - \mathbf{b}\|_2 = \min$$

involved in GMRES is that it leads to numerical instabilities. This is exactly where the Arnoldi iteration comes in! The idea is to replace

$$\mathbf{x}_m = \mathbf{V}_m \mathbf{y}$$

and seek to find

$$\| \mathbf{A} \mathbf{V}_m \mathbf{y} - \mathbf{b} \|_2 = \text{minimum}.$$

Here V_m is the matrix containing as columns the orthonormal vectors produced by the Arnoldi iteration. Notice that we still use the Krylov iteration but we basically orthonormalize them first via Arnoldi.

We can simplify the above problem since $\mathbf{AV}_m = \mathbf{V}_{m+1}\tilde{\mathbf{H}}_m$, where $\tilde{\mathbf{H}}_m$ is the extended Hessenberg matrix we have encountered before. Therefore, we seek to find

$$\|\mathbf{V}_{m+1}\tilde{\mathbf{H}}_{m}\mathbf{y} - \mathbf{b}\|_{2} = \min$$

and after multiplying by \mathbf{V}_{m+1}^T (whose norm is unity), we have

$$\|\tilde{\mathbf{H}}_{m}\mathbf{y} - \mathbf{V}_{m+1}^{T}\mathbf{b}\|_{2} = \text{minimum}.$$

Finally, since $\mathbf{V}_{m+1}^T\mathbf{b} = \parallel \mathbf{b} \parallel_2 \mathbf{e}_1$ by construction of the first orthogonal vector \mathbf{v}_1 , the minimization problem is:

$$\|\tilde{\mathbf{H}}_{m}\mathbf{y} - \|\mathbf{b}\|_{2} \mathbf{e}_{1}\|_{2} = \text{minimum}.$$

Applying QR to this problem now involves the Hessenberg matrix $\tilde{\mathbf{H}}_m$, so

$$\mathbf{Q}_m \tilde{\mathbf{H}}_m = \tilde{\mathbf{R}}_m = \left[egin{array}{c} \mathbf{R}_m \\ 0 \end{array}
ight] \, ,$$

where \mathbf{R}_m is an $m \times m$ upper triangular matrix, and \mathbf{Q}_m is an $(m+1) \times (m+1)$ orthogonal matrix.

We can exploit the structure of the Hessenberg matrix and use Givens rotations for accomplishing the decomposition. Recall that the Hessenberg matrix contains an off-diagonal

below the main diagonal. This set of entries can easily by modified to zero by the appropriate Givens rotations yielding an upper-triangular matrix \mathbf{R} .

Following the discussion above on least-squares, we can compute \mathbf{y}_m from

$$\mathbf{R}_m \mathbf{y} = \parallel \mathbf{b} \parallel \mathbf{q}_1$$
,

where \mathbf{q}_1 is the *first column* of \mathbf{Q}_m excluding the last entry, i.e., \mathbf{q}_1 is a vector of length m. Finally, we compute \mathbf{x}_m from

$$\mathbf{x}_m = \mathbf{V}_m \mathbf{y}$$
.

For convergence purposes we need to compute the residual \mathbf{r}_m . This is done efficiently as follows:

$$\| \mathbf{r}_{m} \|_{2} = \| \tilde{\mathbf{H}}_{m} \mathbf{y} - \| \mathbf{b} \|_{2} \mathbf{e}_{1} \|_{2}$$

$$= \| \mathbf{Q}_{m} (\tilde{\mathbf{H}}_{m} \mathbf{y} - \| \mathbf{b} \|_{2} \mathbf{e}_{1}) \|_{2}$$

$$= \| \mathbf{R}_{m} \mathbf{y} - \| \mathbf{b} \| \tilde{\mathbf{q}}_{1} \|_{2}$$

$$= \| \underbrace{(\mathbf{R}_{m} \mathbf{y} - \| \mathbf{b} \| \mathbf{q}_{1})}_{0} + \| \mathbf{b} \| (\mathbf{q}_{1} - \tilde{\mathbf{q}}_{1}) \|_{2}$$

$$= \| \mathbf{b} \| \times (\text{last entry of} |\tilde{\mathbf{q}}_{1}|),$$

where $\tilde{\mathbf{q}}_1$ is the extended (m+1) vector produced by Arnoldi. This formula then gives a very inexpensive way to compute the residual and check convergence.

Remark 1: Unlike CGM, in GMRES the residuals decrease monotonically, i.e.,

$$\parallel \mathbf{r}_{m+1} \parallel_2 < \parallel \mathbf{r}_m \parallel_2$$

because the corresponding Krylov spaces are nested and the residual is minimized directly. We recall that in CGM the error is minimized instead, and thus reduction of the residual in the L_2 -norm is not guaranteed.

Remark 2: From Arnoldi, we can check the value of $h_{m+1,m}$ to tell us when the iteration can be terminated. When the condition $h_{m+1,m} = 0$ is satisfied, we know that the solution \mathbf{x} lies within the Krylov space \mathcal{K}_m . This being the case, the least-squares problem over the Krylov space will produce the exact solution.

Example: We now present an example to demonstrate the GMRES process. Consider the following matrix

$$\mathbf{A} = \begin{bmatrix} 1.0 & 2.0 & 3.0 \\ 2.0 & 5.0 & 7.0 \\ 3.0 & 8.0 & 9.0 \end{bmatrix}$$

and right-hand-side vector

$$\mathbf{b} = \left[\begin{array}{c} 0.0 \\ 1.0 \\ 2.0 \end{array} \right].$$

We will assume that our initial guess $\mathbf{x}_0 = \mathbf{0}$. Using our modified Arnoldi solver with m = 3 we obtain the extended Hessenberg matrix

$$\tilde{\mathbf{H}}_m = \begin{bmatrix} 14.2000 & 4.8652 & -0.8305 \\ 4.3081 & 1.0414 & 0.3855 \\ 0.0000 & 0.3855 & -0.2414 \\ 0.0000 & 0.0000 & 0.0000 \end{bmatrix}$$

and corresponding matrix

$$\mathbf{V}_{m+1} = \begin{bmatrix} 0.0000 & 0.8305 & 0.5571 & 0.4082 \\ 0.4472 & 0.4983 & -0.7428 & 0.8165 \\ 0.8944 & -0.2491 & 0.3714 & 0.4082 \end{bmatrix}.$$

Both of these matrices will be used. A linear combination of the first three columns of \mathbf{V}_{m+1} will be used to form the solution. To determine what the proper combination is, we now want to solve the following problem:

$$\|\tilde{\mathbf{H}}_{m}\mathbf{y} - \|\mathbf{b}\|_{2} \mathbf{e}_{1}\|_{2} = \text{minimum}.$$

where $\tilde{\mathbf{H}}_m$ is the matrix above and $\|\mathbf{b}\|_2 \mathbf{e}_1 = (\sqrt{5} \ 0 \ 0 \ 0)^T$.

To solve the minimization problem, we accomplish QR decomposition using Givens rotations. For this problem, the three rotation matrices G_0 , G_1 and G_2 are given below:

$$\mathbf{G}_0 = \left[\begin{array}{cccc} 0.0645 & 0.0196 & 0 & 0 \\ -0.0196 & 0.0645 & 0 & 0 \\ 0 & 0 & 1.0000 & 0 \\ 0 & 0 & 0 & 1.0000 \end{array} \right]$$

$$\mathbf{G}_1 = \left[\begin{array}{cccc} 1.0 & 0.0 & 0.0 & 0.0 \\ 0.0 & -0.1876 & 2.5802 & 0.0 \\ 0.0 & -2.5802 & -0.1876 & 0.0 \\ 0.0 & 0.0 & 0.0 & 1.0000 \end{array} \right]$$

$$\mathbf{G}_2 = \left[\begin{array}{cccc} 1.0 & 0.0 & 0.0 & 0.0 \\ 0.0 & 1.0 & 0.0 & 0.0 \\ 0.0 & 0.0 & -16.4509 & 0.0 \\ 0.0 & 0.0 & 0.0 & -16.4509 \end{array} \right].$$

Each successive Givens rotation matrix is formed based upon the updated matrix. Hence, \mathbf{G}_0 is formed based upon the entries of $\tilde{\mathbf{H}}_m$, \mathbf{G}_1 is formed based upon the entries of $\mathbf{G}_0\tilde{\mathbf{H}}_m$, and \mathbf{G}_2 is formed based upon the entries of $\mathbf{G}_1\mathbf{G}_0\tilde{\mathbf{H}}_m$. Applying the Givens rotations to both the right- and left-hand-sides yields a modified matrix

$$\mathbf{R}_m = \begin{bmatrix} 1.0000 & 0.3341 & -0.0460 \\ 0.0000 & 1.0000 & -0.6305 \\ 0.0000 & 0.0000 & 1.0000 \end{bmatrix}$$

and a modified right-hand-side vector

$$\mathbf{z} = \begin{bmatrix} 0.1442 \\ 0.0082 \\ -1.8569 \end{bmatrix}.$$

Here we have omitted the last row of both the matrix and right-hand-side vector because all the entries are zero as expected. We now solve the system $\mathbf{R}_m \mathbf{y} = \mathbf{z}$ using back substitution. We obtain the vector

$$\mathbf{y} = \begin{bmatrix} 0.4472 \\ -1.1626 \\ -1.8569 \end{bmatrix},$$

which provide us with the coefficients for the linear combination of the vectors of \mathbf{V}_{m+1} . Taking the linear combination of the first three columns of \mathbf{V}_{m+1} (since m=3), we obtain:

$$0.4472 \begin{bmatrix} 0.0000 \\ 0.4472 \\ 0.8944 \end{bmatrix} - 1.1626 \begin{bmatrix} 0.8305 \\ 0.4983 \\ -0.2491 \end{bmatrix} - 1.8569 \begin{bmatrix} 0.5571 \\ -0.7428 \\ 0.3714 \end{bmatrix} = \begin{bmatrix} -2.000 \\ 1.000 \\ 0.000 \end{bmatrix},$$

which is the exact solution.

9.5.3 GMRES(k)

The problem with the GMRES is that it requires storing all the vectors \mathbf{v}_m , which can become very expensive. To save storage and also computational cost we can run the GMRES process only for k steps and subsequently restart it with the vector \mathbf{x}_k as an initial guess. This is the so-called GMRES(k) version of the generalized minimum residual method. The choice of k is crucial, as a value too small may lead to divergence while a large value results in extra computations.



Below we present a serial implementation of GMRES(m). This function takes as input the integer m denoting the size of the Krylov subspace to be used, the matrix \mathbf{A} , the right-hand-side vector \mathbf{b} , and the result vector \mathbf{x} . We automatically set the initial direction to $(1,0,\ldots,0)^T$. Two constants can be found in the function: maxit which specifies the maximum number of iterations before terminating and tol which specifies the stopping tolerance.

```
void GMRES(int m, const SCMatrix &A, const SCVector &b,
           SCVector &x){
  int i,j,k,ll,nr;
  int N = A.Rows();
  SCMatrix H(m+1,m), V(N,m+1);
  SCVector w(N),r(N),y(m+1),z(N);
  double * c = new double[m+1];
  double * s = new double[m+1];
  const int maxit = 1000;
  const double tol = 1.0e-7;
  double delta, rho, tmp;
 x.Initialize(0.0);
 r = b - A*x;
 for(j=0;j<maxit;j++){</pre>
   y.Initialize(0.0);
   y(0) = r.Norm_12();
   r.Normalize();
   ModifiedArnoldi(m,r,A,H,V);
   /* Givens Rotation to accomplish QR factorization */
   for(i=0;i<m;i++){
      for(k=1;k<=i;k++){
        tmp = H(k-1,i);
        H(k-1,i) = c[k-1]*H(k-1,i) + s[k-1]*H(k,i);
       H(k,i) = -s[k-1]*tmp + c[k-1]*H(k,i);
      }
      delta = sqrt(H(i,i)*H(i,i)+H(i+1,i)*H(i+1,i));
      c[i] = H(i,i)/delta;
      s[i] = H(i+1,i)/delta;
      H(i,i) = c[i]*H(i,i) + s[i]*H(i+1,i);
      for(k=i+1;k< m+1;k++)
        H(k,i) = 0.0;
      y(i+1) = -s[i]*y(i);
      y(i) = c[i]*y(i);
      rho = fabs(y(i+1));
      if(rho < tol){</pre>
        nr = i;
```

```
break;
      }
    }
    /* Backsolve to obtain coefficients */
    z.Initialize(0.0);
    if(i>=(m-1)){
      nr = m;
      z(nr-1) = y(nr-1)/H(nr-1,nr-1);
    }
    for(k=nr-2;k>=0;k--){
      z(k) = y(k);
      for(ll=k+1;ll<nr;ll++)
        z(k) = H(k,11)*z(11);
      z(k) = z(k)/H(k,k);
    }
    /* Linear combination of basis vectors
       of the Krylov space
                                             */
    for(i=0;i<nr;i++){
      V.GetColumn(i,r);
      x = x + z(i)*r;
    }
    if(rho<tol)
      break;
    r = b - A*x;
  }
  delete[] c;
  delete[] s;
}
```

Remark 1: Observe that this function requires the storage of both \mathbf{H} and \mathbf{V} , both of which may be of the same size as the original matrix \mathbf{A} . This fact is one of the primary motivations for introducing the restart parameter so that smaller Krylov spaces (and hence less storage) can be used.

Remark 2: Instead of forming the Givens matrices explicitly as we did in the example above, we can take advantage of the structure of the matrices so that we need not store the rotation matrices. Instead, we can loop through the appropriate positions, updating as we go.

9.5.4 Preconditioning GMRES

GMRES is used in practice when the matrix \mathbf{A} is not well-conditioned. This means that convergence is typically slow and appropriate preconditioners should be employed. Similar to preconditioning of symmetric matrices where incomplete Cholesky was found to be effective, here we use *incomplete LU (ILU)*. Specifically, we employ the preconditioner

$$M = LU$$

where **L** and **U** are the lower and upper triangular matrices corresponding to **A** but with no fills-in at the entries $a_{ij} = 0$. Note that the preconditioner should not be constructed explicitly but rather be incorporated in the Arnoldi iteration process. To this end, we need to insert the following code

$$\begin{array}{rcl}
\vdots \\
\mathbf{M}\mathbf{y} &=& \mathbf{v}_{j} \\
\mathbf{w} &=& \mathbf{A}\mathbf{y} \\
& \text{for } i = 1, \dots, j \\
& h_{ij} &=& (\mathbf{w}, \mathbf{v}_{i}) \\
& \mathbf{w} &=& \mathbf{w} - h_{ij} \mathbf{v}_{i} \\
& \vdots
\end{array}$$

in the modified Arnoldi iteration algorithm presented above.

Remark: GMRES employs long vectors to obtain orthogonality unlike the three-term recurrence formula associated with symmetric systems. For non-symmetric systems it is also possible to use three-term recurrence formula as done in the Biconjugate Gradient (BiCG) method, employing two mutually orthogonal sequences of vectors. A more stable version of BiCG is the Quasi-Minimal Residual (QMR) method which avoids possible break-downs and converges faster, i.e., as fast as GMRES [39]. Both BiCG and QMR solve tridiagonal systems corresponding to the three-term recurrence sequences. Details of implementation for both methods can be found in [5].

9.5.5 Parallel GMRES

One immediate complication when attempting to parallelize the serial algorithm previously presented is attempting to parallelize the modified Arnoldi component. The brute-force implementation leads to very poor scalability. Similar to the parallel PCG algorithm, we can also use blocked operations to increase the efficiency and parallelism. One way proposed in [64] is to first produce the following basis for the Krylov space

$$\mathbf{v}_1, \mathbf{A}\mathbf{v}_1, \dots, \mathbf{A}^k\mathbf{v}_1$$

and subsequently to orthogonalize the entire set. In contrast, in the standard GMRES method each new vector is immediately orthogonalized to all previous vectors. This approach increases significantly data locality.

Another approach is to employ BLAS2 routines as much as possible in GMRES instead of the obvious BLAS1, which is the least efficient. An algorithm proposed in [79] is to replace the modified Gram-Schmidt with the standard Gram-Schmidt but apply it twice. The double orthogonalization has been shown to reduce the numerical instability associated with the classical Gram-Schmidt method. In the context of more efficient computation, we can now compute all the *dot products* in parallel.

9.6 What Solver to Choose?

The question of what solver and what preconditioner to choose is a complex one, and in many cases there are more than one good candidates. From the algorithmic point of view, we have to consider the properties of the matrix \mathbf{A} , and examine if:

- A is symmetric or non-symmetric.
- A is positive-definite.
- Both **A** and \mathbf{A}^T are available.
- A is sparse.
- A is ill-conditioned or not.
- Good preconditioner exists.

In addition, we have to consider the computational requirements and resources, namely:

- Parallel or serial computation.
- Vector or scalar processor.
- Multi-threading.
- Re-use of data in cache.
- Indirect addressing.
- Memory size.

The above lists are indicative but not exhaustive of the issues that need to be considered in the decision regarding the choice of solver. What is obviously a faster code for serial computations is not necessarily faster on a parallel computer. In table 9.2 we list the main operations in terms of BLAS routines of the iterative solvers we studied in this book. Algorithms that employ dgemv, i.e., the matrix-vector multiply, are typically more efficient as this operation can be done efficiently both in a serial and in a parallel environment.

For problems involving differential equations, the type of differential equation we have to deal with and the corresponding discretization we choose defines the linear system we solve. For Poisson and Helmholtz equations we obtain symmetric systems. A typical orderof-magnitude cost in computational work and storage for various direct and iterative solvers

Method	ddot	daxpy	dgemv	Storage
Jacobi			1	3n
SOR		1	1	2n
CGM	2	3	1	6n
GMRES	j+1	j+1	1	(j+5)n
QMR	2	12	2	16n

Table 9.2: Main operations and storage for iterative solvers; n is the matrix order and j denotes the iteration number. The storage shown does not include the matrix storage.

Method	Direct/Iterative	Work	Storage
Multigrid	Ι	n	n
SSOR/Chebyshev	I	$n^{5/4}$	n
SOR	I	$n^{3/2}$	n
CGM	I	$n^{3/2}$	n
Gauss-Seidel	Ι	n^2	n
Jacobi	Ι	n^2	n
Gauss Elimination/sparse	D	$n^{3/2}$	$n \cdot \log n$
Gauss Elimination/dense	D	n^3	n^2

Table 9.3: Computational work and storage for solution of a Poisson equation on a N-by-N finite difference grid; $n = N^2$. I stands for iterative and D for direct solver.

is shown in table 9.3. The lower bound is of the same order of magnitude as the multigrid method. This clear advantage of multigrid, however, can easily be lost on a parallel computer as very sparse systems that need to be solved at the coarsest level of multigrid are not easily parallelizable. Hence, multigrid is not necessarily the fastest and Jacobi is not the slowest as the estimate for the serial work suggests. Similarly, the most effective (in terms of its spectrum and serial cost) preconditioner may not be the best overall preconditioner. Typically, the more sophisticated preconditioners are more complex and *simplicity* is the rule in parallel computing.

For large size problems the associated memory considerations suggest the use of iterative solvers. A possible decision tree for this case is as follows, see figure 9.17:

- If the matrix is symmetric and positive-definite then preconditioned conjugate gradient may be the best candidate. The question then becomes which preconditioner is the best that is problem-dependent and computer-dependent.
- If **A** is not positive-definite use CGNR assuming the condition number is reasonable.
- On the other hand, if **A** is not symmetric then the first choice should be GMRES or GMRES(k) if memory is at a premium.

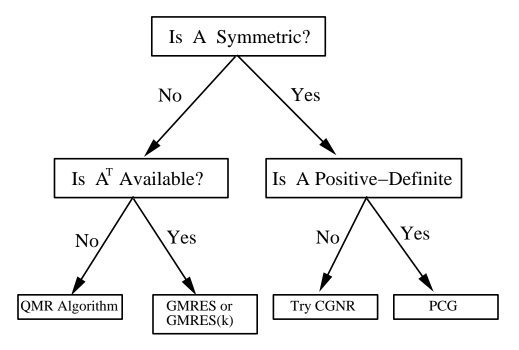


Figure 9.17: Decision tree for which algorithm to use based upon the properties of A.

ullet However, if ${f A}^T$ is not available then the QMR algorithm which is quite robust and as fast as GMRES would be a good candidate – it works effectively even for ill-conditioned matrices.

This is just one of the many possible scenarios – what makes this field interesting is that the choices are not unique!

9.7 Available Software for Fast Solvers

The basic algorithms we have presented in this chapter are relatively easy to program but more sophisticated versions with respect to preconditioning, restarts, orthogonalization, and parallel implementation are available in free software at:

• www.netlib.org

The direct solvers are part of ScaLAPACK and LAPACK++ while the iterative solvers are part of the Templates package [5].

Specifically in ScaLAPACK/LAPACK++, for LU type operations the routine **SGETRF** performs LU factorization with pivoting based on the BLAS3 routines and therefore it is very efficient. A similar routine **SGETF2** is based on BLAS2 routines, and it is also efficient. For the specific implementations involved see [26]. For symmetric positive-definite matrices the routines **SPOTRF** and **SPOTRS** perform Cholesky factorization of a matrix and solve a linear system, respectively. Finally, the routine **SPTTRF** performs an LDL^T factorization of a symmetric positive-definite matrix.

Also in ScaLAPACK/LAPACK++, for QR type operations the routine **SGEQRF** performs QR factorization and the routine **SGEQPF** performs QR factorization with column pivoting. The routine **SGERQF** performs RQ factorization while the routine **SGEHRD** reduces a general matrix to upper Hessenberg form.

The **cpptemplates** files available at www.netlib.org contain implementations of all the iterative solvers presented in this chapter for matrix-vector classes. In particular, the routines declared in **cg.h** and **cheby.h** implement the conjugate gradient method and the preconditioned Chebyshev method, respectively, for symmetric positive-definite systems. The routines declared in **cgs.h**, **gmres.h**, **qmr.h** and **bicg.h** are suitable for non-symmetric systems and their names indicate the corresponding algorithms.

9.8 Homework Problems

1. Find the condition number of the matrices

$$A = \begin{bmatrix} 0.001 & 1 \\ 1 & 1 \end{bmatrix} \text{ and } B = \begin{bmatrix} 7 & 6.990 \\ 4 & 4 \end{bmatrix}$$

2. Let $\mathbf{A} = \mathbf{L} \ \mathbf{D} \mathbf{L}^T$ be a symmetric positive-definite matrix, and $\mathbf{D} = \operatorname{diag}(d_{ii})$. Then, show that

$$\kappa_2(\mathbf{A}) \ge \frac{\max(d_{ii})}{\min(d_{ii})},$$

where $\kappa_2(\mathbf{A})$ is the condition number of \mathbf{A} in the L_2 -norm.

3. For what values of ϵ the matrix

$$\left[\begin{array}{cc} 1 & \epsilon \\ \epsilon & 1 \end{array}\right]$$

is ill-conditioned? How will you results be affected if you are to compute in *single* or *double* precision?

4. Let us assume that the matrix **A** is *strictly* diagonally dominant, i.e.,

$$\sum_{i\neq j} |a_{ij}| < |a_{ii}|.$$

Show that if you apply the LU factorization procedure to **A** with partial pivoting, it has no effect on the rows, that is no actual row exchange occurs. This proves what we discussed in section 9.1.2 that no pivoting is required for a strictly diagonally-dominant matrix.

5. (Almost triangular matrix - Hessenberg)

This matrix is defined by

$$a_{ij} = 0, i > j+1$$

 $a_{ij} \neq 0, i \leq j+1$

Estimate the operation count for the LU factorization and the backward solve for this matrix.

6. Consider the matrix **A** written in a block 2×2 form with submatrices \mathbf{A}_{ij} , i, j = 1, 2 of equal size $m \times m$. Show that the Schur complement defined as

$$S \equiv A_{22} - A_{21}A_{11}^{-1}A_{12}$$

overwrites the matrix A_{22} after m steps of Gaussian elimination without pivoting.

7. Apply Hager's algorithm (by-hand calculation) to a (3×3) matrix whose rows (\mathbf{r}_i , i = 1, 2, 3) are: $\mathbf{r}_i = i + j - 1$, j = 1, 2, 3. Compute also the exact condition number in the L_1 -norm and compare the two values.

8. Apply Hager's algorithm to estimate the condition number of the Hilbert $(n \times n)$ matrix defined by

$$h_{ij} = \frac{1}{i+j-2}$$

for n = 4, 16, 32. (Use the LAPACK routines or any other available routines). What do you observe?

- 9. In the parallel LU program, we did not implement row pivoting. Modify the code given in the text to accomplish row pivoting.
- 10. Using the parallel LU program as a guide, implement an MPI program to accomplish Cholesky Factorization.
- 11. Apply the *incomplete* Cholesky factorization to the tridiagonal matrix (-1, 4, -1). Compare the results with the results from the standard Cholesky factorization. Does your conclusion hold for any banded matrix?
- 12. In the cyclic reduction code presented in this chapter, the matrix \mathbf{A} is allocated as if it were a full matrix. Modify both the serial and the parallel code so that only the necessary amount of memory is used to store the tridiagonal matrix \mathbf{A} .
- 13. Consider the following system:

$$-2x_1 + x_2 = F_1$$

$$x_1 - 2x_2 + 2x_3 = F_2$$

$$x_2 - 2x_3 + 2x_4 = F_3$$

$$x_3 - 2x_4 + 2x_5 = F_4$$

$$x_4 - 2x_5 + 2x_6 = F_5$$

$$x_5 - 2x_6 + 2x_7 = F_6$$

$$x_6 - 2x_7 = F_7$$

Solve this system by hand using cyclic reduction. At what level does this system terminate, and with what equation? Is there sufficient information at the point of termination to obtain the solution? If so, accomplish the back substitution to obtain the answer.

- 14. Use the Householder transformation to show that if \mathbf{H}_e is a Hessenberg matrix and $\mathbf{H}_e = \mathbf{Q} \mathbf{R}$ then the matrix $\mathbf{H}_e^* = \mathbf{R} \mathbf{Q}$ is also a Hessenberg matrix.
- 15. The flop count for solving the overdetermined system $\mathbf{A}\mathbf{x} = \mathbf{b}$ where \mathbf{A} is of size $m \times n$ is (choose one)
 - (a) $\frac{mn^2}{2} + \frac{n^3}{6}$ for normal equations and $\frac{mn^2}{2} \frac{n^3}{6}$ for the Householder QR method.
 - (b) The reverse of (a).
 - (c) None of the above.

- 16. The flop count for QR factorization of an $n \times n$ matrix with column pivoting using the Householder method is (choose one)
 - (a) $\frac{5}{3}n^3$.
 - (b) $\frac{5}{4}n^3$.
 - (c) $\frac{4}{5}n^2$.
- 17. The number of additions and multiplications in Cholesky factorization is roughly half that of LU factorization.
 - (a) True.
 - (b) False.
- 18. The QR method conserves the bandwidth of a matrix (choose one)
 - (a) Always true.
 - (b) In some cases.
 - (c) Never.
- 19. The QR factorization of a matrix **A** is (choose one)
 - (a) Always unique.
 - (b) Unique if \mathbf{A} is non-singular.
 - (c) Not unique.
- 20. The most efficient way of distributing a matrix on a parallel machine is by (choose one)
 - (a) Rows.
 - (b) Columns.
 - (c) Depends on the problem one is trying to solve.
- 21. Define $h = \frac{1}{N}$, $\mu = h^2$, and $q[i][j] = (8\pi^2 + 1)h^2 sin(2\pi hi)sin(2\pi hj)$ where $i, j = 0, \ldots, N-1$. Let **A** be of the form given in figure 9.18.

Solve the matrix system $\mathbf{A}\mathbf{u} = \mathbf{q}$ for \mathbf{u} using the following for methods:

- (a) Serial Jacobi.
- (b) Parallel Jacobi.
- (c) Serial Conjugate Gradient.
- (d) Serial Preconditioned Conjugate Gradient, preconditioned using incomplete Cholesky.

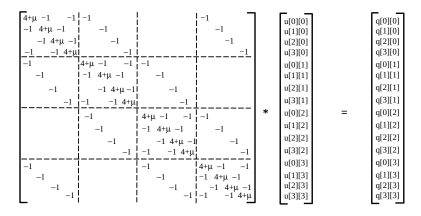


Figure 9.18: Matrix system $\mathbf{A}\mathbf{u} = \mathbf{q}$ for N = 4.

- (e) Parallel Conjugate Gradient.
- (f) Parallel Preconditioned Conjugate Gradient, preconditioned using incomplete Cholesky.

Solve for both N=4 and N=20. Here N denotes the number of points used in both the x and y directions (hence the total number of grid points is N^2 points, which corresponds to the rank of the matrix for which we are solving). For the serial algorithms, provide a graphical plot of the solution u[i][j] at the points (hi, hj) (either a contour or surface plot). For the parallel algorithms, show the parallel speed-up using different number of processors (this will require you to use the MPI_Wtime function to time your runs).

- 22. Solve the Poisson equation on a three-dimensional grid $N \times N \times N$ using a second-order finite difference discretization with Dirichlet boundary conditions. Employ the following (serial) algorithms:
 - (a) Conjugate Gradients.
 - (b) Conjugate Gradients with incomplete Cholesky as preconditioner.
 - (c) SSOR with Chebyshev acceleration.
 - (d) Jacobi.

Estimate the computational work in terms of $\mathcal{O}(n^{\alpha})$, that is find α . Verify these estimates by solving for $n = N^3 = 64^3$ and timing your codes on the same computer.

23. In the text we presented a QR factorization example using the Hilbert matrix. At the conclusion of the example, we had constructed the matrix \mathbf{R} . From the values of \mathbf{w} found in the example, compute the matrix \mathbf{Q} and show that indeed $\mathbf{A} = \mathbf{Q}\mathbf{R}$.

Hint: Recall that the Householder matrix is given by

$$\mathbf{H} = \mathbf{I} - 2 \frac{\mathbf{w} \mathbf{w}^T}{\mathbf{w}^T \mathbf{w}}.$$

Construct \mathbf{H}_1 and \mathbf{H}_2 and use them to construct \mathbf{Q} by the expressions given in the text.

- 24. Modify the QR routine in the text so that it computes the value of \mathbf{Q} . You will need to modify the input arguments of the function to accept a SCMatrix Q which you should fill in with the appropriate values.
- 25. Write a serial function which accomplishes QR decomposition of a tri-diagonal matrix using Givens rotations. The function should have at least the following three arguments: the matrix A as input and the matrices Q and R as output.
- 26. Create a function which solves the system $\mathbf{A}\mathbf{x} = \mathbf{b}$ for tridiagonal matrices \mathbf{A} , and uses the QR code you wrote previously. What property of both \mathbf{Q} and \mathbf{R} can we use to accomplish this efficiently?
- 27. Write a parallel program which accomplishes QR decomposition of a tri-diagonal matrix using Givens rotations. Partition the matrix **A** by rows across the processors. Design your program so that each processor has the rows of **Q** and **R** which correspond to the rows of **A** which reside on the processor.
- 28. Consider five symmetric Toeplitz matrices **A** with entries given by (k = 1, ..., n)

$$a_k^{(1)} = 1/k;$$
 $a_k^{(2)} = 1/\sqrt{k};$ $a_k^{(3)} = 1/k^2;$ $a_k^{(4)} = k;$ $a_k^{(5)} = \cos k/k,$

and an arbitrary non-zero vector **b**. Use PCGM to solve the systems $\mathbf{A}\mathbf{x} = \mathbf{b}$ with circulant preconditioners and experiment with different types (i.e., number of Toeplitz diagonals employed). Compare your results without CG preconditioning in terms of the number of iterations for sizes up to n = 100 and tolerance levels just above single machine accuracy.

- 29. Estimate the operation count for GMRES(k) for fixed k and compare it with GMRES assuming a large value of the order n of matrix \mathbf{A} . Does GMRES(k) converge for any value of k?
- 30. For the GMRES example problem given in the text, use the function provided to attempt GMRES(2). Add *cout* statements to keep report the residual. Try a variety of different initial guesses \mathbf{x}_0 and plot the residual versus iteration for each case. Is the convergence rate the same?
- 31. Consider the linear advection-diffusion equation

$$\frac{\partial u}{\partial t} + \frac{\partial u}{\partial x} = \nu \frac{\partial^2 u}{\partial x^2}$$

where ν is the diffusion coefficient, in the domain

$$x \in [0, 10]$$
 and $t \in [0, 5]$.

We also assume periodic boundary conditions and that the initial conditions are:

$$u(0 \le x \le 1; 0) = x(1-x)$$
 and $u(1 \le x \le 10; 0) = 0$.

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Employ a second-order upwind for the advection and a central finite difference scheme for the diffusion (with a Crank-Nicolson in time) to discretize this problem. Invert the resulted system using GMRES(k) and experiment with different values of k to minimize the solution time. Use n=128 points for the discretization. Does the value of optimum k depends on the time step Δt . Is this method unconditionally stable?

Chapter 10

Fast Eigensolvers

In this chapter we introduce methods for solutions of the standard eigenvalue problem

$$\mathbf{A}\mathbf{x} = \lambda\mathbf{x}$$

as well as for generalized eigenproblems; \mathbf{A} is a square $n \times n$ matrix. The main theory is based on the solvers of the previous chapter for linear systems. Unlike, however, the methods of the previous chapter where both direct and iterative approaches are effective, in eigenvalue problems only iterative solvers are efficient. We start with the simple power method and its variants, and we proceed with more sophisticated methods including a method for non-symmetric eigenproblems using the Arnoldi iteration. We classify the different eigensolvers as *local* or *global* if they are typically used to compute one or two eigenvalues or the entire spectrum, respectively.

We introduce one new MPI function, $MPI_Alltoall$, and demonstrate its use through some of the algorithms presented in this chapter. In addition, we reiterate the use of $MPI_Allgather$ and $MPI_Allreduce$ through example implementations of algorithms presented in this chapter.

10.1 Local Eigensolvers

We have already seen in chapter 2 that computing the eigenvalues accurately from the determinant may not always be possible. The Newton-Raphson method of chapter 4 is an accurate method of computing the roots of the characteristic polynomial, but it can be extremely inefficient for large systems. In the following, we present a simple method to compute iteratively the *maximum* and *minimum* eigenvalues and corresponding eigenvectors. We can also compute any other eigenvalue for which we have a good initial guess.

10.1.1 Basic Power Method

This is a very simple method to obtain the *maximum* eigenvalue. The main idea is to obtain iterates from

$$\mathbf{x}^{k+1} = c\mathbf{A}x^k.$$

where c is a normalization constant that prevents \mathbf{x}^{k+1} from becoming too large. After many iterations $(k \to \infty)$, \mathbf{x}^{k+1} will converge to the eigenvector \mathbf{v}_1 of \mathbf{A} corresponding to the maximum eigenvalue λ_1 . Here, we assume that there exists an eigenvalue λ_1 which dominates, i.e.,

$$|\lambda_1| > |\lambda_2| \ge |\lambda_3| \ldots \ge |\lambda_n|$$
.

We can initialize this iteration by an arbitrary (non-zero) vector \mathbf{x}^0 .

To see why this process converges and at what rate we project the initial guess \mathbf{x}^0 to the space spanned by all the eigenvector \mathbf{v}_i of \mathbf{A} , i.e.,

$$\mathbf{x}^0 = c_1 \mathbf{v}_1 + c_2 \mathbf{v}_2 \dots + \dots + c_n \mathbf{v}_n$$

and thus

$$\mathbf{x}^k = \mathbf{A}\mathbf{x}^{k-1} = \ldots = A^k\mathbf{x}^0 = c_1\lambda_1^k\mathbf{v}_1 + \ldots + c_n\lambda_n^k\mathbf{v}_n.$$

Now, we see that

$$\frac{\mathbf{x}^k}{c_1 \lambda_1^k} = \mathbf{v}_1 + \frac{c_2}{c_1} \left(\frac{\lambda_2}{\lambda_1}\right)^k \mathbf{v}_2 + \ldots + \frac{c_n}{c_1} \left(\frac{\lambda_n}{\lambda_1}\right)^k \mathbf{v}_n$$

converges to \mathbf{v}_1 because all factors

$$\left(\frac{\lambda_i}{\lambda_1}\right)^k, i \neq 1$$

are less than one and tend to zero for $k \to \infty$.

The *convergence rate* is determined by the relative magnitude of the second largest to the largest term, i.e., the ratio

$$\frac{|\lambda_2|}{|\lambda_1|}$$
,

which represents the most resistive contribution to the error. The smaller this ratio is the faster the convergence is.

The following pseudo-code summarizes the algorithm:

Initialize:
$$\mathbf{x}^0$$

Begin Loop: for $k = 1, 2, ...$
 $\hat{\mathbf{x}}^k = \mathbf{A}\mathbf{x}^{k-1}$
 $\mathbf{x}_k = \frac{\hat{\mathbf{x}}^k}{\max(\hat{\mathbf{x}}^k)}$
endfor

End Loop:

In the above pseudo-code, the operator $max(\mathbf{y})$ of a vector \mathbf{y} returns the entry of \mathbf{y} with the maximum modulus. Hence if $\mathbf{y} = (4.32, -9.88, 2.9)^T$, $max(\mathbf{y}) = -9.88$ since |-9.88| > |4.32| > |2.9|. This operation is used merely for normalization of the eigenvector. Upon termination of the loop, $max(\hat{\mathbf{x}}^k) \to \lambda_1$ and $\mathbf{x}^k \to \mathbf{v}_1$.

Example: Let us consider the 3×3 Hilbert matrix

$$\mathbf{A} = \begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ \frac{1}{2} & \frac{1}{3} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{4} & \frac{1}{5} \end{bmatrix}$$

to which we will apply the power method to find its maximum eigenvalue. Starting with an initial guess of $(1,1,1)^T$, we obtain the sequence of eigenvalue iterates listed in table 10.1. The iteration terminates when the absolute difference between successive iterates is less than 10^{-7} .

Iteration	Eigenvalue Iterate
0	1.833333333
1	1.437878788
2	1.410835968
3	1.408537172
4	1.408337881
5	1.408320573
6	1.40831907
7	1.40831894
8	1.408318928

Table 10.1: Sequence converging to the largest eigenvalue of the 3×3 Hilbert matrix.

In the algorithm above the value of λ_1 is estimated from the maximum component of \mathbf{x}^k .

However, any other norm can be used, e.g., the L_2 norm

$$\lambda_1 = \frac{\parallel x^{k+1} \parallel}{\parallel x^k \parallel} = \frac{\parallel \mathbf{A} \mathbf{x}^k \parallel}{\parallel x^k \parallel},$$

or alternatively the eigenvalue can be computed via the Rayleigh quotient. This is defined by

$$R(\mathbf{A}, x^k) = \frac{(\mathbf{x}^k)^T \mathbf{A} \mathbf{x}^k}{\parallel \mathbf{x}^k \parallel^2}.$$

If $\mathbf{x} = \mathbf{v}_1$, i.e., the first eigenvector, then obviously $R(\mathbf{A}, \mathbf{v}_1) = \lambda_1$. In general, if \mathbf{A} is a symmetric matrix and \mathbf{x} is a close approximation to the eigenvector \mathbf{v}_1 , the Rayleigh quotient is close to the corresponding eigenvalue λ_1 . This can be seen by projecting $\mathbf{x} = \sum_{i=1}^n c_i \mathbf{v}_i$, as before, where \mathbf{v}_i are orthonormal due to symmetry of \mathbf{A} , and $\mathbf{v}_i^T \mathbf{v}_i = 1$. Then

$$R(\mathbf{A}, x) = \frac{\mathbf{x}^T \mathbf{A} \mathbf{x}}{\mathbf{x}^T \mathbf{x}} = \frac{(c_1 \mathbf{v}_1 + \dots + c_n \mathbf{v}_n)^T \mathbf{A} (c_1 \mathbf{v}_1 + \dots + c_n \mathbf{v}_n)}{c_1^2 + c_2^2 + \dots + c_n^2}$$

$$= \frac{\lambda_1 c_1^2 + \lambda_2 c_2^2 + \dots + \lambda_n c_n^2}{c_1^2 + c_2^2 + \dots + c_n^2}$$

$$= \lambda_1 \left[\frac{1 + \left(\frac{\lambda_2}{\lambda_1}\right) \left(\frac{c_2}{c_1}\right)^2 + \dots + \left(\frac{\lambda_n}{\lambda_1}\right) \left(\frac{c_n}{c_1}\right)^2}{1 + \left(\frac{c_2}{c_1}\right)^2 + \dots + \left(\frac{c_n}{c_1}\right)^2} \right].$$

Due to our assumption that $\mathbf{x} \approx \mathbf{v}_1$ we have that $c_1 \gg c_i$, $\forall i \neq 1$, and thus, the quantity in brackets tends to 1, and correspondingly $R(\mathbf{A}, \mathbf{x})$ tends to λ_1 .

The convergence of the power method can be enhanced by *shifting* the eigenvalues, so instead of multiplying the initial guess by powers of \mathbf{A} we multiply by powers of $(\mathbf{A} - \sigma \mathbf{I})$ which has eigenvalues $(\lambda_i - \sigma)$ while the eigenvectors remain the same. The corresponding convergence rate is then estimated by

$$\left|\frac{\lambda_2 - \sigma}{\lambda_1 - \sigma}\right|.$$

We can exploit this to enhance convergence by carefully choosing σ to significantly reduce the fraction above. It turns out that if all the eigenvalues are real, the best shift is either

$$\sigma = \frac{1}{2}(\lambda_2 + \lambda_n)$$
 or $\sigma = \frac{1}{2}(\lambda_1 + \lambda_{n-1})$

if $(\lambda_1 - \lambda_2)$ or $(\lambda_{n-1} - \lambda_n)$ is larger, respectively.

Remark: If the eigenvalue λ_1 is complex the power method fails unless complex arithmetic is used with an initial complex vector \mathbf{x}^0 .

10.1.2 Inverse Shifted Power Method

In order to compute selectively the *minimum* eigenvalue we can apply again the power method by multiplying by powers of the inverse, i.e., A^{-1} . Thus, the iteration procedure here is

$$\mathbf{A}\mathbf{x}^{k+1} = c\,\mathbf{x}^k.$$

where again c is a normalization constant. This method is most effective with a proper shift, so the modified iteration is

$$(\mathbf{A} - \sigma I)\mathbf{x}^{k+1} = c\,\mathbf{x}^k,$$

and the solution will converge to the smallest shifted eigenvalue $|\lambda_n - \sigma|$ and the corresponding eigenvector, assuming that the shift σ is close to λ_n (the smallest eigenvalue). The rate of convergence is now

$$\frac{|\lambda_n - \sigma|}{|\lambda_{n-1} - \sigma|},$$

so if σ is close to λ_n , the above ratio is very small and convergence is very fast. Clearly, if σ is close to any other eigenvalue λ_i the corresponding contribution dominates and then we obtain the eigenpair $(\lambda_i, \mathbf{v}_i)$. Hence, if we have a collection of approximations σ_i to the true eigenvalues λ_i , we can use the inverse shifted power method to obtain all the eigenpairs $(\lambda_i, \mathbf{v}_i)$.

The following pseudo-code summarizes the algorithm:

Initialize: Choose
$$\mathbf{x}^0$$
 Choose σ Factorize $\mathbf{A} - \sigma \mathbf{I} = \mathbf{L} \mathbf{U}$

Begin Loop: for $k = 1, 2, ...$

$$\hat{\mathbf{x}}^k = \mathbf{U}^{-1} \mathbf{L}^{-1} \mathbf{x}^{k-1}$$

$$\mathbf{x}^k = \frac{\hat{\mathbf{x}}^k}{\max(\mathbf{x}^k)}$$
if $|R(\mathbf{A}, x^k) - R(\mathbf{A}, x^{k-1})| < \epsilon$
return endfor

Upon termination (after the tolerance ϵ is reached), \mathbf{x}^k converges to the desired eigenvector, and the corresponding eigenvalue can be computed from the Rayleigh quotient.

Remark 1: Note that we do not actually compute explicitly the inverse \mathbf{A}^{-1} or $(\mathbf{A} - \sigma \mathbf{I})^{-1}$ but we simply perform an LU factorization only once outside the loop. The computational complexity of this algorithm is $\mathcal{O}(n^2)$ times the number of iterations plus the initial $\mathcal{O}(2n^3/3)$ cost for the LU factorization.

Remark 2: If an iterative solver is used instead of the LU factorization for the linear solver, then a slightly more stringent tolerance can be used for the linear solver compared to the desired accuracy in eigenvalue. This idea can be used when **A** is sparse and may result in big savings.

Remark 3: To accelerate convergence, we can start with a few iterations using the standard power method, obtain a first good guess and corresponding shift σ via the Rayleigh quotient, and then switch to the inverse iteration method.

Remark 4: The matrix $(\mathbf{A} - \sigma \mathbf{I})$ is ill-conditioned, however in practice the error associated with this seems to favor the inverse iteration as it grows toward the direction of the desired eigenvector. Therefore, the inverse shifted power method is a *stable* method.

We can modify the inverse shifted power method and enhance convergence even more (to third-order) if we update the value of the shift adaptively using the **Rayleigh quotient**.

The following algorithm presents this modification:

Initialize: Choose
$$\mathbf{x}^0$$

Compute $\sigma_0 = \frac{(\mathbf{x}^0)^T \mathbf{A} \mathbf{x}^0}{(\mathbf{x}^0)^T \mathbf{x}^0}$

Begin Loop: for $k = 0, 1, \dots$

$$\mathbf{L}^k \mathbf{U}^k = \mathbf{A} - \sigma_k \mathbf{I}$$

$$\hat{\mathbf{x}}^{k+1} = (\mathbf{U}^k)^{-1} (\mathbf{L}^k)^{-1} \mathbf{x}^k$$

$$\mathbf{x}^{k+1} = \frac{\hat{\mathbf{x}}^{k+1}}{\max(\hat{\mathbf{x}}^{k+1})}$$

$$\sigma_{k+1} = \frac{(\mathbf{x}^{k+1})^T \mathbf{A} \mathbf{x}^{k+1}}{(\mathbf{x}^{k+1})^T \mathbf{x}^{k+1}}$$
if $|\sigma_{k+1} - \sigma_k| < \epsilon$
return
endfor
End Loop

Remark 5: While this algorithm triples the number of correct digits in each iteration it requires $\mathcal{O}(n^3)$ work at each iteration because the matrix $(\mathbf{A} - \sigma_k \mathbf{I})$ changes in each iteration. A more economical approach is to "freeze" σ_k for a few iterations so that the LU decomposition is not employed in each iteration. The resulted convergence rate is then *less than cubic* but overall this is a more efficient approach.

Software

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Putting it into Practice

Below we present an implementation of the inverse shifted power method with Rayleigh quotient adaptation. This function takes two inputs: the matrix \mathbf{A} for which we are interested in finding an eigenvalue and eigenvector, and a vector \mathbf{x} containing the initial guess \mathbf{x}^0 . This function has as its return value an eigenvalue of \mathbf{A} , and places the corresponding eigenvector associated with the eigenvalue in the vector \mathbf{x} .

```
double ISPowerMethod(SCMatrix &A, SCVector &x){
  const int maxit = 100;
  const double tol = 1.0e-7;
  double tmp, sigma1, sigma2;
  SCMatrix Amod(A.Rows());
  SCVector y(A.Rows());
  // Calculation of the Rayleigh Quotient
  y = A*x;
  sigma1 = dot(y,x)/dot(x,x);
  for(int k=0;k<maxit;k++){</pre>
    Amod = A;
    for(int i=0;i<Amod.Rows();i++)</pre>
      Amod(i,i) = Amod(i,i) - sigma1;
    // Use LU solver from Chapter 9 to obtain solution x
    GaussElimination(Amod, x, 1);
    // Normalize with the element of maximum modulus
    tmp = x.ElementofMaxMod();
    x = x/tmp;
    y = A*x;
    sigma2 = dot(y,x)/dot(x,x);
    if(fabs(sigma1-sigma2) < tol)</pre>
      return sigma2;
    sigma1 = sigma2;
```

```
}
cout << "ISPowerMethod - Max number of iterations reached\n";
return sigma2;
}</pre>
```

Remark 6: A downside of this implementation is that we require storage of an additional matrix MatAmod having the same size as **A**. We have chosen to do this because we want to be able to re-use the LU solver we implemented in Chapter 9. If memory were at a premium, we could re-write the LU solver to take into account the modification to the diagonal entry, and hence we would not need to store an additional matrix.

Example: Let us consider the 3×3 Hilbert matrix

$$\mathbf{A} = \begin{bmatrix} 1 & \frac{1}{2} & \frac{1}{3} \\ \frac{1}{2} & \frac{1}{3} & \frac{1}{4} \\ \frac{1}{3} & \frac{1}{4} & \frac{1}{5} \end{bmatrix}$$

and use the inverse shifted power method to solve for the maximum eigenvalue. Using the program above, if we take the initial eigenvector guess to

$$\left[\begin{array}{c} 1.0\\1.0\\1.0\end{array}\right],$$

we obtain the eigenvalue 1.40832 and corresponding eigenvector

$$\begin{bmatrix} 1.0 \\ 0.556033 \\ 0.390908 \end{bmatrix}.$$

Observe that this is the same maximum eigenvalue which we found before using the power method (as it should be!). If, however, we take the initial eigenvector guess to be

$$\begin{bmatrix}
0.4 \\
-0.4 \\
-0.4
\end{bmatrix}$$

we obtain the eigenvalue 0.122327 and corresponding eigenvector

$$\begin{bmatrix} -0.843517 \\ 0.813998 \\ 1.0 \end{bmatrix}.$$

As was pointed out earlier, the inverse shifted power method allows us to pick up different eigenvalues (other than just the minimum) based upon the initial eigenvector guess. In this case, our initial guess provided us with the second largest eigenvalue of the Hilbert matrix.

10.2 Householder Deflation

The Householder transformation is very useful in "deflating" a matrix, \mathbf{A} , i.e., reducing its size from $n \times n$ to $(n-1) \times (n-1)$. This is often needed in eigensolvers, especially in conjunction with local and semi-direct methods. The objective is to compute the subdominant eigenvalues after we compute the maximum or minimum eigenvalue with the power method or inverse power method, respectively.

Let us assume that we have computed the maximum eigenvalue λ_1 and the corresponding eigenvector \mathbf{v}_1 . Recall from section 9.3 that we can then obtain a Householder matrix \mathbf{H} by

$$\mathbf{H}\mathbf{v}_1 = \begin{bmatrix} \alpha \\ 0 \\ \vdots \\ 0 \end{bmatrix} = \alpha \mathbf{e}_1.$$

We now construct the matrix $\mathbf{G} \equiv \mathbf{H}\mathbf{A}\mathbf{H}^T$ and investigate its structure. Since $(\lambda_1, \mathbf{v}_1)$ is an eigenpair it satisfies

$$\mathbf{A}\mathbf{v}_1 = \lambda_1 \mathbf{v}_1 \Rightarrow \mathbf{H} \mathbf{A} \mathbf{v}_1 = \lambda_1 \mathbf{H} \mathbf{v}_1$$

$$\Rightarrow \mathbf{H} \mathbf{A} \cdot \mathbf{I} \cdot \mathbf{v}_1 = \lambda_1 \mathbf{H} \mathbf{v}_1 \Rightarrow \mathbf{H} \mathbf{A} \cdot (\mathbf{H}^T \mathbf{H}) \cdot \mathbf{v}_1 = \lambda_1 \mathbf{H} \mathbf{v}_1$$

$$\Rightarrow \mathbf{H} \cdot \mathbf{A} \cdot \mathbf{H}^T (\mathbf{H} \mathbf{v}_1) = \lambda_1 (\mathbf{H} \mathbf{v}_1),$$

where we recall that **H** is orthonormal and thus $\mathbf{I} = \mathbf{H}^T \cdot \mathbf{H}$. We also have by definition that

$$\mathbf{H}\mathbf{v}_1 = \alpha \mathbf{e}_1$$

so by substituting in the above, we obtain

$$\mathbf{H} \cdot \mathbf{A} \cdot \mathbf{H}^{T}(\alpha \mathbf{e}_{1}) = \lambda_{1} \alpha \mathbf{e}_{1} \Rightarrow \mathbf{G} \mathbf{e}_{1} = \lambda_{1} \mathbf{e}_{1} = \begin{bmatrix} \lambda_{1} \\ 0 \\ \vdots \\ 0 \end{bmatrix}.$$

Therefore, the matrix G has the form

$$\mathbf{G} = \left[egin{array}{ccccc} \lambda_1 & * & * & * & * & * \\ dots & & \mathbf{A}_{n-1} & & \\ 0 & & & \end{array}
ight]$$

where \mathbf{A}_{n-1} is a submatrix $(n-1) \times (n-1)$ which has the same (n-1) remaining eigenvalues of \mathbf{A} . This can be seen by computing

$$\det(\mathbf{G} - \lambda \mathbf{I}) = (\lambda_1 - \lambda) \cdot \det(\mathbf{A}_{n-1} - \lambda I) = 0.$$

To compute the first subdominant eigenvalue of **A** we then need to apply the power method to \mathbf{A}_{n-1} , repeat the process once more, and so on.

We now recapitulate the basic algorithm:

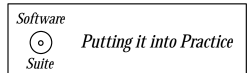
- 1. Compute the maximum eigenvalue λ_1 and the corresponding eigenvector \mathbf{v}_1 using the power method or shifted inverse power method.
- 2. Obtain the Householder matrix **H** using the eigenvector \mathbf{v}_1 .
- 3. Form the matrix $\mathbf{G} = \mathbf{H}\mathbf{A}\mathbf{H}^T$.
- 4. Extract the eigenvalue at G_{11} .
- 5. Define $\mathbf{A}_{n-1} \equiv G_{2:n,2:n}$.
- 6. Repeat process on the newly defined matrix \mathbf{A}_{n-1} .

The deflation procedure can be applied efficiently if we want to compute *two* to *three* eigenvalues. For a larger number of eigenvalues we need to switch to global eigensolvers, see section 10.3.

Remark: Another deflation procedure is to subtract off from the matrix **A** the contribution $\lambda_1 \mathbf{v}_1 \mathbf{v}_1^T$, so the new matrix

$$\mathbf{A}_1 \equiv \mathbf{A} - \lambda_1 \mathbf{v}_1 \mathbf{v}_1^T$$

has eigenvalues $0, \lambda_2, \lambda_3, \dots, \lambda_n$. However, this procedure could sometimes be unstable unlike the Householder deflation which is always stable.



We will now present a program for accomplishing one iteration of the algorithm just recapitulated above. To accomplish the algorithm presented, we must introduce a new MPI function, $MPI_Alltoall$. This function allows a "scatter" from each process simultaneously, see figure 10.1. We will now present the function call syntax, argument list explanation, usage example and some remarks.

MPI_Alltoall:

Function Call Syntax

```
int MPI_Alltoall(
        void*
                        sendbuf
                                       in
                        sendcount
        int
                                       in
        MPI_Datatype
                       sendtype
                                       in
        void*
                        recvbuf
                        recvcount
        int
                                       in
        MPI_Datatype
                       recvtype
                                       in
        MPI_Comm
                        comm
```

Understanding the Argument List

- sendbuf starting address of the send buffer.
- sendcount number of elements in the send buffer.
- sendtype data type of the elements in the send buffer.
- recvbuf starting address of the receive buffer.
- recvcount number of elements for any single receive.
- recvtype data type of the elements in the receive buffer.
- comm communicator.

Example of Usage

Remarks

- *MPI_Alltoall* is a collective operation (i.e., it should be called by all processes within the communicator).
- Both the *sendbuf* and *recvbuf* arrays are relevant on all processes in the communicator.
- In most cases the *sendtype* and *recvtype* are identical and the value of *sendcount* and the value of *recvcount* are identical. MPI requires that the amount of data sent (sendcount times the size in bytes of the datatype sendtype) equals the amount of data received (recvcount times the size in bytes of the datatype recvtype) per process/root pair.
- The allocated size of both the *sendbuf* and *recvbuf* arrays should be at least equal to the value of *recvtype* times the number of processes (*totalnodes*).

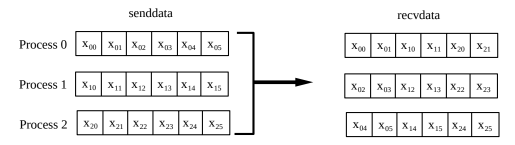


Figure 10.1: MPI_Alltoall schematic demonstrating data distribution to all processes of two data objects from each process.

We now present an MPI program to accomplish one iteration of the previously presented Householder deflation algorithm. In this program, we assume that the size of the matrix is divisible by the number of processes used. For the purpose of this example, we will once again use the Vandermonde matrix formed at the Chebyshev points. It will be distributed by rows across the processes (each process forming only those rows which it needs). Observe that we will once again use $MPI_Allreduce$ and $MPI_Allgather$ (as part of the power method algorithm), and that we introduce the use of $MPI_Alltoall$. We now present the source code for this program, and will provide some remarks following it.

```
#include <iostream.h>
#include <iomanip.h>
#include "SCmathlib.h"
#include "SCchapter3.h"
#include<mpi.h>

void ChebyVandermonde(int npts, double *A, int row);

// Global variable to set size of the system
const int size = 10;
```

```
int main(int argc, char *argv[]){
 int i,j,k,index,cnt;
 int mynode, totalnodes;
 double tmps1, tmps2, sum, total;
 const int maxit = 1000; //maximum number of iterations
                         //for the power method
 MPI_Init(&argc,&argv);
 MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
 MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
 int numrows = size/totalnodes;
 double **A_local = new double*[numrows];
 double **A_local_t = new double*[numrows];
 double * xold = new double[size];
 double * xnew = new double[numrows];
 double * tmparray = new double[size];
 for(i=0;i<numrows;i++){</pre>
   A_local[i] = new double[size];
   A_local_t[i] = new double[size];
   index = mynode*numrows + i;
   ChebyVandermonde(size, A_local[i], index);
 }
 /*****************/
 /* Use the power method to obtain the first eigenvector */
 // Initialize starting vector to 1.0
 for(i=0;i<size;i++)</pre>
   xold[i] = 1.0;
 for(int it=0;it<maxit;it++){</pre>
   // Matrix-vector multiplication
   for(i=0;i<numrows;i++)</pre>
     xnew[i] = dot(size,A_local[i],xold);
   // Compute Euclidian norm of new vector
   sum = 0.0;
   for(i=0;i<numrows;i++)</pre>
     sum += xnew[i]*xnew[i];
   MPI_Allreduce(&sum, &total, 1, MPI_DOUBLE,
                 MPI_SUM,MPI_COMM_WORLD);
   total = sqrt(total);
```

```
// Scale vector by its norm
 for(i=0;i<numrows;i++)</pre>
   xnew[i] = xnew[i]/total;
 // Gather (Allgather) new vector to all processors
 MPI_Allgather(xnew, numrows, MPI_DOUBLE, tmparray, numrows,
MPI_DOUBLE,MPI_COMM_WORLD);
 // Compute difference between old and new vector
 sum = 0.0;
 for(i=0;i<size;i++){</pre>
   sum += (xold[i]-tmparray[i])*(xold[i]-tmparray[i]);
   xold[i] = tmparray[i]; //replace old with new
 }
 if(sqrt(sum) < 1.0e-7) //termination condition</pre>
   break;
}
// Modify eigenvector per Householder transformation
xold[0] = xold[0] + Sign(xold[0]);
/****************/
/* Compute S = A*H (S is a temporary state
                                            */
/* stored in A_local)
                                            */
tmps1 = dot(size,xold,xold);
for(i=0;i<numrows;i++){</pre>
 tmps2 = dot(size,A_local[i],xold);
 for(j=0;j<size;j++)</pre>
   A_local[i][j] = A_local[i][j] - 2.0*tmps2*xold[j]/tmps1;
}
/************/
/* Transpose temporary state S */
/***********/
for(i=0;i<numrows;i++){</pre>
 MPI_Alltoall(A_local[i], numrows, MPI_DOUBLE, tmparray,
              numrows, MPI_DOUBLE,MPI_COMM_WORLD);
 cnt = 0:
 for(k=0;k<totalnodes;k++)</pre>
```

```
for(j=0;j<numrows;j++)</pre>
A_local_t[j][i+k*numrows] = tmparray[cnt++];
 }
 /***************************/
 /* Compute G = H*S = H*A*H */
 /********************/
 tmps1 = dot(size,xold,xold);
 for(i=0;i<numrows;i++){</pre>
   tmps2 = dot(size,A_local_t[i],xold);
   for(j=0;j<size;j++)</pre>
     A_{local_t[i][j]} = A_{local_t[i][j]} -
                      2.0*tmps2*xold[j]/tmps1;
 }
 /* Transpose G so that it is stored by rows across the
 /* processors, just as the original A was stored
                                                       */
 for(i=0;i<numrows;i++){</pre>
   MPI_Alltoall(A_local_t[i], numrows, MPI_DOUBLE, tmparray,
                numrows, MPI_DOUBLE,MPI_COMM_WORLD);
   cnt = 0;
   for(k=0;k<totalnodes;k++)</pre>
     for(j=0;j<numrows;j++)</pre>
A_local[j][i+k*numrows] = tmparray[cnt++];
 }
 for(i=0;i<numrows;i++){</pre>
   delete[] A_local[i];
   delete[] A_local_t[i];
 }
 delete[] A_local;
 delete[] A_local_t;
 delete[] tmparray;
 delete[] xold;
 delete[] xnew;
 MPI_Finalize();
}
```

```
void ChebyVandermonde(int npts, double *A, int row){
  int i,j;
  double * x = new double[npts];

  ChebyshevPoints(npts,x);

  for(j=0;j<npts;j++)
     A[j] = pow(x[row],j);

  delete[] x;
}</pre>
```

Remark 1: Because of the particular form of the Householder matrix, we do not have to form it explicitly. By taking advantage of this form, we can accomplish the matrix multiplication of $\mathbf{A} * \mathbf{H}$ in $\mathcal{O}(n^2)$ operations instead of the normal $\mathcal{O}(n^3)$ operations necessary for an arbitrary matrix-matrix multiplication.

Remark 2: Observe that we use the $MPI_Alltoall$ function to transpose the intermediary stage $\mathbf{A} * \mathbf{H}$ so that we can accomplish pre-multiplication by \mathbf{H} again. Examine the indexing carefully following the MPI call. After receiving the array tmparray, we must now partition it properly across the rows which a processor holds.

Remark 3: In general, the $MPI_Alltoall$ function call is one of the most taxing communication calls because it requires the exchange of information from each process to all other processes. Because of this fact, the $MPI_Alltoall$ function is often used in benchmarking the communication capability of parallel machines.

10.3 Global Eigensolvers

Unlike local eigensolvers global methods for eigenproblems do not require a good initial guess for an eigenvalue. In addition, they provide the entire eigenspectrum. We will discuss two such approaches that are based on two different concepts: the QR method and the Lanczos method. The first one is semi-direct and general while the second one is iterative and suitable for symmetric sparse eigensystems.

10.3.1 The QR Eigensolver

The main idea of this method is to construct a sequence of *similar* matrices $\mathbf{A}_1, \mathbf{A}_2, \dots, \mathbf{A}_k$, where $\mathbf{A}_1 = \mathbf{A}$, and \mathbf{A}_k approaches an upper triangular matrix, as $k \to \infty$, of the form

$$A_k \to \begin{bmatrix} \lambda_1 & * & * & \dots & * \\ & \lambda_2 & * & \dots & * \\ & & \lambda_3 & \dots & \vdots \\ & & & \ddots & \\ & & & & \lambda_n \end{bmatrix}.$$

The Schur triangulization theorem [46] guarantees that this triangular matrix exists if an orthonormal matrix \mathbf{Q} is involved in the similarity transformation. If the matrix \mathbf{A} is symmetric, we know that all the eigenvalues are real; the scalar entries in the diagonal correspond to those real eigenvalues. If the matrix is non-symmetric, however, we may encounter complex eigenvalues, which manifest themselves as 2×2 submatrices on the diagonal in the form

$$A_k \to \begin{bmatrix} \lambda_1 & * & * & * & \dots & \dots & * \\ 0 & a_{22} & a_{23} & * & \dots & \dots & * \\ 0 & a_{32} & a_{33} & * & \dots & \dots & * \\ 0 & 0 & 0 & \lambda_4 & & & \vdots \\ \vdots & & & & \lambda_5 & & \\ & & & & \ddots & \\ 0 & & & & & \lambda_n \end{bmatrix}.$$

These 2×2 submatrices, as shown above, appear also for double eigenvalues and they are sometimes referred to as *bumps*. The eigenvalues λ_2 and λ_3 can then be obtained by simply solving the second-order characteristic polynomial of the 2×2 submatrix.

We now present the basic QR iteration:

Initialize:
$$\mathbf{A}_0 = \mathbf{A} \ and \ \mathbf{A}_0 = \mathbf{Q}_0 \mathbf{R}_0$$

Begin Loop: $\mathbf{A}_1 = \mathbf{R}_0 \mathbf{Q}_0 \ and \ \mathbf{A}_1 = \mathbf{Q}_1 \mathbf{R}_1$
 $\mathbf{A}_2 = \mathbf{R}_1 \mathbf{Q}_1 \ and \ \mathbf{A}_2 = \mathbf{Q}_2 \mathbf{R}_2$

:

Remark: The key observation in the algorithm above is that we alternate between $\mathbf{Q}_i \mathbf{R}_i$ decompositions and $\mathbf{R}_i \mathbf{Q}_i$ products.

To prove that all matrices in the sequence are *similar*, we first examine

$$\mathbf{A}_1 = \mathbf{R}_0 \mathbf{Q}_0 = (\mathbf{Q}_0^T \mathbf{A}_0) \mathbf{Q}_0$$

and since $\mathbf{Q}_0^T \mathbf{Q}_0 = \mathbf{I}$ we conclude that \mathbf{A}_1 is similar to $\mathbf{A}_0 = \mathbf{A}$.

This applies to iteration k as well, i.e.

$$\mathbf{A}_k = \mathbf{R}_{k-1} \mathbf{Q}_{k-1} = (\mathbf{Q}_{k-1}^T \mathbf{A}_{k-1}) \mathbf{Q}_{k-1}.$$

The method *converges* if all eigenvalue have *different magnitude*, and the sequence of the diagonal will be

$$|\lambda_1>|\lambda_2|>\ldots>|\lambda_n|,$$

see [94]; the rate of convergence is

$$\frac{|\lambda_{i+1}|}{|\lambda_i|}.$$

We note here that a similar sequence can be generated with about the same or even less computational cost by using the LU decomposition, instead of the QR factorization, the so-called LR method. However, no pivoting can be employed as this will destroy the upper triangular structure and may change the eigenvalues (e.g., sign); thus the LU procedure will be unstable. We also note that the QR decomposition can be applied to a singular matrix unlike the LU factorization. So overall, the QR eigensolver is definitely the preferred solution. However, a case for which the LU factorization may be used over the QR is when we have a banded and diagonally dominant matrix A. The reason is that LU preserves the bandwidth unlike the QR approach, which fills in the (initially) zero off-diagonal elements at the rate of one additional diagonal on each side per iteration. The diagonal dominance of A guarantees that no pivoting is needed for stability, so overall the LR eigensolver will be a faster method in this case.

10.3.2 The Hessenberg QR Eigensolver

The basic QR method has two main disadvantages:

1. It requires QR factorization in each step which costs

$$\mathcal{O}\left(\frac{4n^3}{3}\right)$$

i.e., twice as much as an LU decomposition.

2. It also diverges if two eigenvalues have the same magnitude.

Fortunately, both problems have been solved!

With regards to the cost, the main idea is to *first transform* the matrix \mathbf{A} to an upper Hessenberg matrix, i.e.,

$$\mathbf{A} \longrightarrow \mathbf{H}$$

(see section 9.3.1), and subsequently perform QR factorization on \mathbf{H} . The method works because all subsequent matrices maintain the *Hessenberg* form as can be verified with an example! The initial cost to perform the Hessenberg reduction is

$$\mathcal{O}\left(\frac{5}{3}n^3\right)$$
,

however the QR factorization of a Hessenberg matrix is $\mathcal{O}(n^2)$, and this is the cost we incur in every iteration. Therefore, the Hessenberg QR eigensolver is an $\mathcal{O}(n)$ more efficient than the basic QR eigensolver.

10.3.3 Shifted QR Eigensolver

The convergence rate is at most *linear* for the QR eigensolver, i.e., the error in every iteration is

$$\epsilon_{k+1} \sim C\epsilon_k$$

where $\epsilon_k \equiv \lambda_n - (a_{nn})_k$.

This is dictated by the ratio

$$\frac{|\lambda_{i+1}|}{|\lambda_i|}$$
,

which we want to make as small as possible. We can employ the idea of a *shift* similar to what we did in the power method (see section 10.1.1).

Let us first assume that we have only *real* and *simple* eigenvalues. Instead of applying the QR factorization to \mathbf{H} we will apply it to $(\mathbf{H} - \sigma \mathbf{I})$, thereby shifting the entire eigenspectrum by σ to $(\lambda_i - \sigma)$.

Specifically, we choose σ such that the element in the position (n, n-1) converges to zero, i.e. $h_{n,n-1} \to 0$ as rapidly as possible. Upon convergence, the entry in the position (n, n) (after we subtract the shift σ) will be the desired eigenvalue. Clearly, the closest estimate to λ_n we have is $h_{nn}^{(k)}$ of the matrix \mathbf{H}_k .

The modified algorithm to generate the sequence is then

Perform QR factorization:

$$\mathbf{H}_k - h_{nn}^{(k)} \mathbf{I} = \mathbf{Q}_k \mathbf{R}_k$$

Obtain New Iterate:

$$\mathbf{H}_{k+1} = \mathbf{R}_k \mathbf{Q}_k + h_{nn}^{(k)} \mathbf{I}$$

and so on.

Then \mathbf{H}_{k+1} is similar to \mathbf{H}_k because

$$\mathbf{H}_{k+1} = \mathbf{Q}_k^T \mathbf{Q}_k \mathbf{R}_k \mathbf{Q}_k + \sigma \mathbf{I}$$

$$= \mathbf{Q}_k^T (\mathbf{Q}_k \mathbf{R}_k) \mathbf{Q}_k + \sigma \mathbf{I}$$

$$= \mathbf{Q}_k^T [\mathbf{Q}_k \mathbf{R}_k + \sigma I] \mathbf{Q}_k$$

$$= \mathbf{Q}_k^T \mathbf{H}_k \mathbf{Q}_k.$$

• The convergence of the shifted method is quadratic in general but increases to cubic if the original matrix A is symmetric.

For double or complex eigenvalues the shift strategy has to change in order to deal effectively with the "bumps" in the diagonal of the \mathbf{A}_k matrix, as discussed earlier. Specifically, we modify the algorithm so that each iteration has two substeps. We can use the eigenvalues λ_2 and λ_3 of the submatrix to obtain the iterates as follows

$$\mathbf{H}_1 - \lambda_2 \mathbf{I} = \mathbf{Q}_1 \mathbf{R}_1$$
$$\mathbf{H}_2 = \mathbf{R}_1 \mathbf{Q}_1 + \lambda_2 \mathbf{I}$$

¹A convergence criterion used in practice is to take $|h_{i,i-1}^{(k)}| < C[|h_{ii}^{(k)}| + |h_{i-1,i-1}^{k}|]$.

$$\begin{aligned} \mathbf{H}_2 - \lambda_3 \mathbf{I} &= \mathbf{Q}_2 \mathbf{R}_2 \\ \mathbf{H}_3 &= \mathbf{R}_2 \mathbf{Q}_2 + \lambda_3 \mathbf{I} \\ \vdots &\vdots \end{aligned}$$

and so on, for subsequent iterations.

This method requires complex arithmetic because λ_2 and λ_3 may be a complex pair - this is certainly a disadvantage. However, Wilkinson [94] has come up with another method that avoids complex arithmetic and is based on the sum $(\lambda_2 + \lambda_3)$ and the product $\lambda_2 \cdot \lambda_3$ of the complex pair.

Remark: After the last eigenvalue λ_n has been obtained, deflation of the Hessenberg matrix can be applied to find the next one at reduced computational cost. In general, it takes about two to three QR iterations to compute one eigenvalue. The *total cost* is about $\mathcal{O}(10n^3)$ but such an operation count is approximate and problem dependent.

10.3.4 The Symmetric QR Eigensolver: Wilkinson Shift

Symmetric matrices are special, first because they have only real eigenvalues but also because the Hessenberg transformation reduces the matrix to a symmetric tridiagonal matrix. The QR iteration maintains this property for the entire sequence. The other good news is that the QR factorization of a symmetric tridiagonal matrix costs only $\mathcal{O}(n)$, instead of $\mathcal{O}(n^2)$ for the Hessenberg matrix. Therefore, the computational cost in this case is $\mathcal{O}(n)$ times the number of iterations to converge to all eigenvalues.

Wilkinson [94] has suggested that instead of using the $h_{nn}^{(k)}$ as the shift σ to accelerate convergence, a more effective way is to use the eigenvalue of the 2 × 2 matrix

$$\begin{bmatrix} h_{n-1,n-1} & h_{n,n-1} \\ h_{n,n-1} & h_{nn} \end{bmatrix}$$

which is closer to the entry h_{nn} . Convergence of the symmetric QR eigensolver is *cubic* and the operation count is $\mathcal{O}(n^2)$ but there is an $\mathcal{O}(\frac{2}{3}n^3)$ cost associated with the reduction of **A** to a tridiagonal matrix.

10.3.5 Parallel QR Eigensolver: Divide-and-Conquer

A fully parallel algorithm for the symmetric eigenvalue problem was developed by Dongarra & Sorensen [32]. It is based on the QR iteration and the divide-and-conquer approach we have encountered many times in this book, i.e. we split the problem into two subproblems, each of which is further split into two subproblems, and so on.

First, by reducing **A** to its Hessenberg form (recall section 9.3.1), we tridiagonalize the full symmetric matrix $\mathbf{A} \to \mathbf{T}$, where the latter has the form

$$\mathbf{T} = \begin{bmatrix} a_1 & b_1 \\ b_1 & a_2 & b_2 & O \\ & \ddots & \ddots & \ddots \\ & & b_{n-2} & a_{n-1} & b_{n-1} \\ O & & b_{n-1} & a_n \end{bmatrix}.$$

We then divide **T** into two symmetric tridiagonal matrices of half size as follows:

$$\mathbf{T}_{1} = \begin{bmatrix} a_{1} & b_{1} \\ b_{1} & a_{2} & b_{2} & & O \\ & \ddots & \ddots & \ddots \\ & & b_{k-2} & a_{k-1} & b_{k-1} \\ O & & b_{k-1} & a_{k} & -b_{k} \end{bmatrix}$$

and

$$\mathbf{T}_{2} = \begin{bmatrix} (a_{k+1} - b_{k}) & b_{k+1} \\ b_{k+1} & a_{k+2} & b_{k+2} & O \\ \vdots & \vdots & \ddots & \vdots \\ O & b_{n-1} & a_{n} \end{bmatrix}$$

so that

$$\mathbf{T} = \begin{bmatrix} \mathbf{T}_1 & 0 \\ 0 & \mathbf{T}_2 \end{bmatrix} + b_k \mathbf{z} \mathbf{z}^T,$$

where

$$\mathbf{z} = \begin{bmatrix} 0 \\ \vdots \\ 0 \\ 1 \\ \vdots \\ 0 \end{bmatrix} \leftarrow k \\ \leftarrow k + 1$$

Next, we can diagonalize both T_1 and T_2 using appropriate orthogonal matrices Q_1 and Q_2 as follows

$$\mathbf{T}_1 = \mathbf{Q}_1 \mathbf{D}_1 \mathbf{Q}_1^T$$
 and $\mathbf{T}_2 = \mathbf{Q}_2 \mathbf{D}_2 \mathbf{Q}_2^T$

where

and

$$\mathbf{D} \equiv \left[\begin{array}{cc} \mathbf{D}_1 & \mathbf{0} \\ \mathbf{0} & \mathbf{D}_2 \end{array} \right] \, .$$

Based on this definition and by introducing

$$\boldsymbol{\xi} = \begin{bmatrix} \mathbf{Q}_1 & 0 \\ 0 & \mathbf{Q}_2 \end{bmatrix}^T \mathbf{z}$$

we can re-write the matrix T as follows

$$\mathbf{T} = \begin{bmatrix} \mathbf{Q}_1 & 0 \\ 0 & \mathbf{Q}_2 \end{bmatrix} \left\{ \begin{bmatrix} \mathbf{D}_1 & 0 \\ 0 & \mathbf{D}_2 \end{bmatrix} + b_k \boldsymbol{\xi} \boldsymbol{\xi}^T \right\} \begin{bmatrix} \mathbf{Q}_1 & 0 \\ 0 & \mathbf{Q}_2 \end{bmatrix}^T$$

which shows that the matrix T is *similar* to the matrix

$$\mathbf{G} \equiv \mathbf{D} + b_k \boldsymbol{\xi} \boldsymbol{\xi}^T.$$

It turns out that the eigenvalues λ_i of **G** are given by the following equation

$$1 + b_k \sum_{i=1}^n \frac{\xi_i^2}{d_i - \lambda} = 0, \tag{10.1}$$

where ξ_i are the elements of the vector $\boldsymbol{\xi}$. Observe that the problem has been reduced to the root finding problem:

• Find λ such that $f(\lambda) = 0$ where $f(\lambda) = 1 + b_k \sum_{i=1}^n \frac{\xi_i^2}{d_i - \lambda}$.

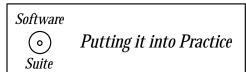
The equation for $f(\lambda)$ is known as the **secular equation**. In [26] several approaches in solving equation (10.1) are presented. Having computed the eigenvalues λ_i of \mathbf{G} , which are also eigenvalues of \mathbf{A} , we can now compute the eigenvectors. First, the corresponding eigenvectors of \mathbf{G} are (see [26, 32])

$$\mathbf{y}_i = (\mathbf{D} - \lambda_i \mathbf{I})^{-1} \boldsymbol{\xi}.$$

Finally, the eigenvectors of the original matrix **A** are given by

$$\mathbf{v}_i = \left[\begin{array}{cc} \mathbf{Q}_1 & 0 \\ 0 & \mathbf{Q}_2 \end{array} \right] \mathbf{y}_i.$$

Clearly, this is the first level of dividing the problem. We can now repeat *recursively* this process for \mathbf{T}_1 and \mathbf{T}_2 , which are also tridiagonal and half in size compared to the original matrix. These, in turn can be further divided into two, and so on.



We now present a parallel implementation of the parallel QR eigensolver just discussed. This program solves for a very simple symmetric tridiagonal matrix consisting of the values $0.5, 1.5, \ldots$ on the diagonal and the value 0.3 on the off diagonals. This matrix was chosen for two reasons: first, it is diagonally dominant similar to most matrices which we encountered when using finite difference discretizations, and secondly, all the eigenvalues of this matrix

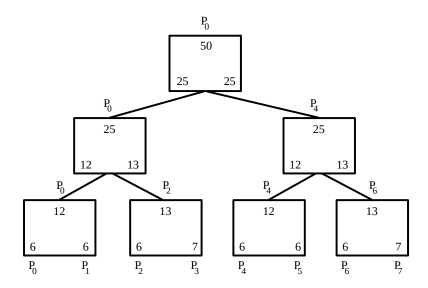


Figure 10.2: Computational partitioning of a matrix of size 50. This partitioning is explained in the text.

are simple (i.e., the multiplicity of each eigenvalue is exactly one). The rationale behind the second point will be presented in the remarks following the program.

For simplicity, we assume that the number of MPI processes used is a power of two. This being the case, an example partitioning of a matrix of size 50 is presented in figure 10.2.

Figure 10.2 is to be understood as follows. Suppose that we have eight processes P_0, \ldots, P_7 on which to solve for a matrix **A** of size 50. First, traversing down the tree, we see that using the algorithm described above, we continuously subdivide the problem until each processor has assigned to it some subproblem to solve. In this case, P_0 solves a subproblem of size 6, P_1 solves a subproblem of size 6, etc. Each of these subproblems is solved in parallel by each individual process. When all the subproblems have been solved, a fan-in style algorithm is employed to complete the computation. Traversing back up the partitioning tree, we see that process P_0 and P_1 combine their subproblems, each of size 6, into one subproblem of size 12 which is now solved on process P_0 . In this implementation, P_1 remains idle throughout the remainder of the computation. Similarly, other processes combine their subproblems pairwise until finally the original problem of size 50 is solved.

We now present a C++/MPI code to accomplish the parallel QR eigensolver procedure just described. We present the commented code, and then will conclude with some remarks about the code.

```
#include <iostream.h>
#include <iomanip.h>
#include "SCmathlib.h"
#include "SCchapter10.h"
#include<mpi.h>

// Global variable to set size of the system const int size = 50;
```

```
int main(int argc, char *argv[]){
  int i,j,k,ll,m,isum,ioffset,cnt,N1,N2;
  int mynode, totalnodes;
  MPI_Status status;
  double bn,*tmpd,**tmpdd;
  MPI_Init(&argc,&argv);
  MPI_Comm_size(MPI_COMM_WORLD, &totalnodes);
 MPI_Comm_rank(MPI_COMM_WORLD, &mynode);
  // Set up storage on each process
  double * a = new double[size];
  double * b = new double[size];
  double * lambda = new double[size];
  double ** Q = CreateMatrix(size,size);
  double ** Q1 = CreateMatrix(size, size);
  double ** Q2 = CreateMatrix(size, size);
  double * xi = new double[size];
  double * d = new double[size];
  int ** index = ICreateMatrix(totalnodes,totalnodes);
  double ** adjust = CreateMatrix(totalnodes, totalnodes);
  //Form the Matrix A
  for(i=0;i<size;i++){</pre>
   a[i] = i+1.5;
   b[i] = 0.3;
  }
  //Set up recursive partitioning of the matrix.
  //Each process will solve for some subset of the problem
  index[0][0] = size;
  for(i=0;i<log2(totalnodes);i++){</pre>
    isum = 0:
    for(j=0; j < pow(2,i); j++){
      index[i+1][2*j] = index[i][j]/2;
      index[i+1][2*j+1] = index[i][j] - index[i+1][2*j];
      isum += index[i+1][2*j];
      adjust[i][j] = b[isum-1];
      a[isum-1] = a[isum-1] - b[isum-1];
              = a[isum] - b[isum-1];
      a[isum]
      isum += index[i+1][2*j+1];
    }
```

```
}
// Each process solves recursively for its subpart of
// the problem.
ioffset = (int) log2(totalnodes);
isum = 0;
for(k=0;k<mynode;k++)</pre>
  isum += index[ioffset][k];
TDQREigensolver(index[ioffset][mynode],&a[isum],
                 &b[isum],d,Q1);
// Fan-in algorithm to finish solving system
for(i=0;i<log2(totalnodes);i++){</pre>
  isum = 0; cnt = 0;
  for(j=0;j<totalnodes;j+=(int)pow(2,i)){</pre>
    if(mynode == j){
      if(isum%2==0){
        MPI_Recv(d+index[ioffset][isum],index[ioffset][isum+1],
                  MPI_DOUBLE, j+(int)pow(2,i),1,MPI_COMM_WORLD,
                  &status);
        for(k=0;k<index[ioffset][isum+1];k++)</pre>
          MPI_Recv(Q2[k],index[ioffset][isum+1],MPI_DOUBLE,
                    j+(int)pow(2,i),1,MPI_COMM_WORLD, &status);
        N1 = index[ioffset][isum];
        N2 = index[ioffset][isum+1];
        bn = adjust[ioffset-1][cnt++];
      }
      else{
        MPI_Send(d,index[ioffset][isum],MPI_DOUBLE,
                  j-(int)pow(2,i),1,MPI_COMM_WORLD);
        for(k=0;k<index[ioffset][isum];k++)</pre>
          MPI_Send(Q1[k],index[ioffset][isum],MPI_DOUBLE,
                    j-(int)pow(2,i),1,MPI_COMM_WORLD);
        }
    }
    isum++;
  }
  for(j=0; j<totalnodes; j+=(int)pow(2,i+1)){</pre>
    if(mynode == j){
```

```
cnt = 0;
for(k=0;k<N1;k++)
  xi[cnt++] = Q1[N1-1][k];
for(k=0;k<N2;k++)
  xi[cnt++] = Q2[0][k];
// Solve for the secular equation to
// obtain eigenvalues
SolveSecularEq(bn,N1+N2,d,xi,lambda);
// Form the Q matrix from Q1 and Q2
for(k=0;k<N1;k++){
  for(l1=0;l1<N1+N2;l1++){
    Q[k][11] = 0.0;
    for(m=0; m<N1; m++)
      Q[k][11] += Q1[k][m]*xi[m]/(d[m]-lambda[11]);
  }
}
for(k=0;k<N2;k++){
  for(l1=0;l1<N1+N2;l1++){
    Q[N1+k][11] = 0.0;
    for(m=0; m<N2; m++)
      Q[k+N1][11] += Q2[k][m]*xi[N1+m]/
                      (d[N1+m]-lambda[l1]);
  }
}
// Normalize the Q matrix so that each eigenvector
// has length one
double sum;
for(k=0;k<N1+N2;k++){
  sum = 0.0;
  for(ll=0;ll<N1+N2;ll++)
    sum += Q[11][k]*Q[11][k];
  sum = sqrt(sum);
  for(ll=0;ll<N1+N2;ll++)
    Q[11][k] = Q[11][k]/sum;
}
// Swap d and lambda arrays for use in the
// next part of the fan-in algorithm
tmpd = d;
d = lambda;
lambda = tmpd;
```

}

```
// Swap Q and Q1 for use in the
      // next part of the fan-in algorithm
      tmpdd = Q1;
      Q1 = Q;
      Q = tmpdd;
    }
  ioffset = ioffset - 1;
}
if(mynode==0){
  cout << "The eigenvalues are: " << endl;</pre>
  for(k=0;k<size;k++)</pre>
    cout << d[k] << endl;</pre>
}
MPI_Barrier(MPI_COMM_WORLD);
delete[] a;
delete[] b;
delete[] lambda;
delete[] xi;
delete[] d;
DestroyMatrix(Q,size,size);
DestroyMatrix(Q1,size,size);
DestroyMatrix(Q2,size,size);
DestroyMatrix(adjust, totalnodes, totalnodes);
IDestroyMatrix(index,totalnodes,totalnodes);
MPI_Finalize();
```

Remark 1: Each process initially solves its assigned subproblem using the recursive routine TDQREigensolver, which can be found in the $software\ suite$. Observe that the parallel code resembles the serial TDQREigensolver solver – understanding the serial code will greatly help in interpreting the parallel code.

Remark 2: To solve the secular equation we use a routine SolveSecularEq which can be found in the software suite. This routine implements a very simple bisection algorithm for finding the roots of the secular equation. One assumption of this routine is that all the roots are distinct. More complicated methodologies which handle a wider variety of cases

can be found in [26]. Commonly, a modified Newton-Raphson method is used for solving such problems.

Remark 3: It should be pointed out that solving for the eigenvectors using the equation

$$\mathbf{y}_i = (\mathbf{D} - \lambda_i \mathbf{I})^{-1} \boldsymbol{\xi},$$

is not always numerically stable because two values of λ_i very close to each other will produce nearly parallel eigenvectors. In [26], several alternative method which are more numerically stable are presented.

Remark 4: In [32], a discussion is provided as to how one might keep all processors busy during the fan-in procedure. In the example code given above, when a process has completed its subproblem and if it is not involved in the fan-in process, it will remain idle. However, as pointed out in [32], the idle processes could be called back into service to work on solving the secular equation or for accomplishing parts of the matrix multiplications that are required. This will enhance the parallel efficiency.

10.3.6 The Lanczos Eigensolver

This algorithm, developed by Lanczos in the 1950s, is particularly useful for large *symmetric* matrices which are *sparse*. The QR method can also be used for such systems, but it does not maintain the sparsity of the new matrices in each iteration and thus becomes inefficient.

The key idea in Lanczos's approach is to make use of all the previous iterates

$$\mathbf{v}, \mathbf{A}\mathbf{v}, \mathbf{A}^2\mathbf{v}, \dots \mathbf{A}^{k-1}\mathbf{v}$$

produced in a power method iteration. These k vectors form the so-called Krylov space:

$$\mathcal{K}_k(\mathbf{A}, \mathbf{v}) = \text{span} \{\mathbf{v}, \mathbf{A}\mathbf{v}, \mathbf{A}^2\mathbf{v}\dots, \mathbf{A}^{k-1}\}$$

of dimension k. In principle, after n iterations assuming that all

$$\mathbf{v}, \mathbf{A}\mathbf{v}, \dots, \mathbf{A}^{n-1}\mathbf{v}$$

are linearly independent, we can express any vector \mathbf{x} in terms of the basis formed by the vectors generated by that sequence. However, this sequence is not computationally friendly, and there is a bias towards the maximum eigenvalues of \mathbf{A} . The idea is then to orthogonalize these vectors but do it fast, that is do not use Gram-Schmidt or QR but something new! The new orthonormalization will be based on the magic three-term recurrence formula! We have already seen how to use recurrence relations in the context of orthogonal polynomials (see section 3.1) and also in the conjugate gradient method (see 4.1.7).

Assuming that we have produced the orthonormal vectors from the above sequence

$$\mathbf{q}_1, \, \mathbf{q}_2, \, \ldots, \, \mathbf{q}_k$$

then we can construct the orthornormal matrix \mathbf{Q} by employing \mathbf{q}_i as its columns. We also recall that

$$\mathbf{Q}^T\mathbf{Q} = \mathbf{I},$$

which is the condition of orthonormality.

We can then transform \mathbf{A} to its *similar* matrix \mathbf{T} using the similarity transformation, i.e.

$$\mathbf{Q}^T \mathbf{A} \mathbf{Q} = \mathbf{T} \text{ or } \mathbf{A} \mathbf{Q} = \mathbf{Q} \mathbf{T}$$

Clearly, the eigenvalues of A are the same as the eigenvalues of the matrix T. The latter is a tridiagonal matrix of the form

$$T_k = \begin{bmatrix} & & \alpha_1 & & \beta_1 & & & \\ & \beta_1 & & \alpha_2 & & \beta_2 & & & O \\ & & \ddots & & \ddots & & \ddots & \\ O & & & \beta_{k-1} & & \alpha_k & & \beta_k & \end{bmatrix}.$$

The tridiagonal structure of T is a consequence of the three-term magic formula, so the columns of Q satisfy

$$\mathbf{A}\mathbf{q}_i = \beta_{i-1}\mathbf{q}_{i-1} + \alpha_i\mathbf{q}_i + \beta_i\mathbf{q}_{i+1}, \quad i = 1, 2, \dots$$

We can compute now the coefficients α_i and β_i from the orthonormality conditions. First, multiplying the above equation by \mathbf{q}_i^T we obtain

$$\mathbf{q}_i^T \mathbf{A} \mathbf{q}_i = \beta_{i-1} \mathbf{q}_i^T \mathbf{q}_{i-1} + \alpha_i \mathbf{q}_i^T \mathbf{q}_i + \beta_i \mathbf{q}_i^T \mathbf{q}_{i+1},$$

where $\mathbf{q}_i^T \mathbf{q}_i = 1$; also $\mathbf{q}_i^T \mathbf{q}_{i-1} = 0$ and $\mathbf{q}_i^T \mathbf{q}_{i+1} = 0$, due to the orthonormality condition. Therefore, from this equation we obtain

$$\alpha_i = \mathbf{q}_i^T \mathbf{A} \mathbf{q}_i.$$

Conversely, assuming that α_i is obtained from the above formula we can prove that $\mathbf{q}_{i+1} \perp \mathbf{q}_i$ by induction.

The coefficient β_i is also obtained from the recurrence formula

$$\mathbf{r}_i \equiv \beta_i \mathbf{q}_{i+1} = \mathbf{A} \mathbf{q}_i - \alpha_i \mathbf{q}_i - \beta_{i-1} \mathbf{q}_{i-1}$$

if we further assume that

$$\beta_i \neq 0 \Rightarrow \beta_i = \parallel \mathbf{r}_i \parallel_2$$

and

$$\mathbf{q}_{i+1} = \frac{\mathbf{r}_i}{\beta_i} \,.$$

All vectors \mathbf{q}_{i+1} generated by the three-term sequence are orthogonal to all \mathbf{q}_k , k < i given the symmetry of the matrix \mathbf{A} . This can be proved by induction as follows. First, we assume that

$$\mathbf{q}_{i+1} \perp \mathbf{q}_i$$
 and $\mathbf{q}_{i+1} \perp \mathbf{q}_{i-1}$

and also that from the previous step (of the induction process) we have already that the condition

$$\mathbf{q}_i \perp \mathbf{q}_k, \quad k < i$$

is valid. We will then prove that $\mathbf{q}_{i+1} \perp \mathbf{q}_k$ but first we prove that the vectors \mathbf{q}_k and \mathbf{q}_i are A-orthogonal. For this, we multiply the three-term recurrence formula by \mathbf{q}_k $(k \leq i-2)$

$$\mathbf{q}_{k}^{T} \mathbf{A} \mathbf{q}_{i} = (\mathbf{q}_{k}^{T} \mathbf{A}^{T}) \mathbf{q}_{i} = (\mathbf{A} \mathbf{q}_{k})^{T} \mathbf{q}_{i} \text{ (due to symmetry)}$$

$$= [\beta_{k-1} \mathbf{q}_{k-1} + \alpha_{k} \mathbf{q}_{k} + \beta_{k} \mathbf{q}_{k+1}]^{T} \mathbf{q}_{i}$$

$$= \beta_{k-1} \mathbf{q}_{k-1}^{T} \mathbf{q}_{i} + \alpha_{k} \mathbf{q}_{k}^{T} \mathbf{q}_{i} + \beta_{k} \mathbf{q}_{k+1}^{T} \mathbf{q}_{i}$$

$$= 0 + 0 + 0$$

$$\Rightarrow \mathbf{q}_{j}^{T} (\mathbf{A} \mathbf{q}_{i}) = 0.$$

Now, multiplying the three-term formula again we have

$$\beta_i \mathbf{q}_k^T \mathbf{q}_{i+1} = \mathbf{q}_k^T \mathbf{A} \mathbf{q}_i - \alpha_i \mathbf{q}_k^T \mathbf{q}_i - \beta_i \mathbf{q}_k^T \mathbf{q}_{i-1} = 0 + 0 + 0,$$

so we have proved by induction that \mathbf{q}_{i+1} is orthogonal to all previous \mathbf{q}_i . We note here that each induction step has two substeps, as presented above, first to show the A-orthogonality and second to obtain the final result.

In practice, the problem is that the orthogonality is not preserved. In fact, as soon as one eigenvalue converges all the basis vectors \mathbf{q}_i pick up perturbations biased toward the direction of the corresponding eigenvector and orthogonality is lost. This has been analyzed in detail by Paige [74] who found that a "ghost" copy of the eigenvalue will appear again in the tridiagonal matrix \mathbf{T} . The straightforward remedy is to fully re-orthonormalize the sequence by using Gram-Schmidt or even QR. However, either approach would be expensive if the dimension of the Krylov space is large, so instead a selective re-orthonormalization is pursued. More specifically, the practical approach is to orthonormalize half-way i.e., within half machine-precision $\sqrt{\epsilon_M}$, see [4].

The following algorithm presents an orthogonalized version of the basic Lanczos algorithm for symmetric matrices.

Lanczos Algorithm

Initialize: choose
$$\mathbf{q}_0 = 0$$
; arbitrary $\mathbf{r} \neq 0$; and $\beta_0 = ||\mathbf{r}||_2$
Begin Loop: for $j = 1, \dots$

$$\mathbf{q}_j = \frac{\mathbf{r}}{\beta_{j-1}}$$

$$\mathbf{r} = A\mathbf{q}_j$$

$$\mathbf{r} = \mathbf{r} - \mathbf{q}_{j-1}\beta_{j-1}$$

$$\alpha_j = \mathbf{q}_j^T\mathbf{r}$$

$$\mathbf{r} = \mathbf{r} - \mathbf{q}_j\alpha_j$$
Orthogonalize if necessary
$$\beta_j = ||\mathbf{r}||_2$$
Compute approximate eigenvalues of \mathbf{T}_j
Test for convergence endfor

End Loop

Remark 1: The algorithm can be implemented using a single matrix-vector product per iteration and BLAS1 routines, i.e. ddot and daxpy. Only three vectors need to be stored at any given time, that is \mathbf{r} , \mathbf{q}_j and \mathbf{q}_{j-1} although there are also algorithms that require only two stores but are more difficult to implement.

Remark 2: We note that if we have n iterations in the above loop we will obtain the entire eigenspectrum of \mathbf{T} and therefore of \mathbf{A} . This can be done with a standard QR method operating on a tridiagonal matrix, which requires *linear* work only. However, we can stop at any iteration j < n and the resulted matrix \mathbf{T}_j will have approximate values for the first j eigenvalues of the matrix \mathbf{A} . Hence, this method is both *local* in that we can obtain a subset of the eigenvalues but also *global* since we can obtain the entire eigenspectrum.

Remark 3: If the eigenvalues of A are not well separated, then we can use a *shift* and employ the matrix

$$(\mathbf{A} - \sigma \mathbf{I})^{-1}$$
,

following the shifted inverted power method to generate the appropriate Krylov subspaces.

Remark 4: The Lanczos algorithm can be extended to non-symmetric systems – see modified algorithms in references [79], [4].

10.4 Generalized Eigenproblems

So far we have dealt with linear eigenvalue problems but in applications we often encounter problems of the form

$$\mathbf{A}(\lambda)\mathbf{v} = 0,$$

where $\mathbf{A}(\lambda)$ is a matrix function of λ . The general problem can be, in theory, solved using the Newton-Raphson method for systems. However, we have (n+1) unknowns $\lambda, \mathbf{x}_i, \ldots, \mathbf{x}_n$ and only n equations. To close the system we introduce the condition of normalization which provides the $(n+1)^{th}$ equation.

Other special forms of the generalized eigenproblem can be handled with more efficient approaches. We demonstrate a few examples next.

In its simplest form the generalized eigenproblem may be written as

$$\mathbf{A}\mathbf{v} = \lambda \mathbf{B}\mathbf{v}$$
.

Assuming that **B** is not singular, then the above can be turned to

$$(\mathbf{B}^{-1}\mathbf{A})\mathbf{v} = \lambda \mathbf{v},$$

which is similar to the standard eigenproblem presented earlier. However, there is still concern if \mathbf{B} is not singular but *nearly singular*. The algorithm in the next section deals with this case.

10.4.1 The QZ Eigensolver

When the matrix \mathbf{B} is almost singular, it is not accurate to construct $\mathbf{B}^{-1}\mathbf{A}$ explicitly to find its eigenvalues. Instead, we transform \mathbf{A} and \mathbf{B} simultaneously to simpler forms and then compute the eigenvalues. This is accomplished in two steps.

In the *first step*, we pre- and post-multiply $\bf A$ and $\bf B$ by two orthonormal matrices $\bf Q$ and $\bf Z$ as follows to achieve:

$$\mathbf{A}^* = \mathbf{Q}^T \mathbf{A} \mathbf{Z}$$
 upper Hessenberg matrix $\mathbf{B}^* = \mathbf{Q}^T \mathbf{B} \mathbf{Z}$ upper triangular matrix

In the *second step*, we apply QR to the matrix pair A^* and B^* to reduce them further to an upper triangular form.

The eigenvalues of the generalized problem $\mathbf{A}\mathbf{v}=\lambda\mathbf{B}\mathbf{v}$ after triangulization of \mathbf{A} are given by

$$\lambda_i = \begin{cases} a_{ii}^*/b_{ii}^* & \text{if } b_{ii}^* \neq 0\\ \infty & \text{if } b_{ii}^* = 0 \text{ and } a_{ii}^* \neq 0. \end{cases}$$

For details on the algorithm we refer the reader to [70] and [21, 79].

10.4.2 Singular Eigenproblems

Many interesting applications involve a singular matrix **B**; for example, the Orr-Sommerfeld equation in fluid dynamics [80]. In this case we can employ a local iteration procedure similar to the inverse power method. To this end, we solve the original system along with its adjoint, i.e.,

$$[\mathbf{A} - (\lambda_k - \sigma)\mathbf{B}]\mathbf{u}^{k+1} = \sigma \mathbf{B}\mathbf{u}^k$$
$$[\mathbf{A} - (\lambda_k - \sigma^*)\mathbf{B}]^T\mathbf{v}^{k+1} = \sigma \mathbf{B}^T\mathbf{v}^k,$$

where σ and σ^* are normalization constants. We assume that we have some good initial guesses λ_1 , \mathbf{u}^0 and \mathbf{v}^0 . If \mathbf{A} and \mathbf{B} are symmetric the above two equations are identical. For non-symmetric systems this formulation with the *adjoint* accelerates convergence.

The eigenvalue is obtained through a generalized Rayleigh quotient of the form

$$\lambda_{k+1} = \frac{\mathbf{v}_k^T \mathbf{A} \mathbf{u}^k}{\mathbf{v}_k^T \mathbf{B} \mathbf{u}^k},$$

which will also be used in the next iteration. The convergence rate of this local method is third-order.

10.4.3 Polynomial Eigenproblems

We now consider the second-order polynomial eigenproblem of the form

$$[\mathbf{A}\lambda^2 + \mathbf{B}\lambda + \mathbf{C}]\mathbf{v} = 0,$$

where **A** is non-singular. By setting $\mathbf{x} = \mathbf{v}$ and $\mathbf{y} = \lambda \mathbf{v}$, the equation above can be re-written as

$$\mathbf{A}\lambda\mathbf{y} + \mathbf{B}\mathbf{y} + \mathbf{C}\mathbf{x} = 0,$$

which can be recast into

$$\mathbf{G}_2\mathbf{y} + \mathbf{G}_1\mathbf{x} = \lambda\mathbf{y}$$

with $G_2 = -A^{-1}B$ and $G_1 = -A^{-1}C$. Finally, we put it in block matrix form

$$\left[\begin{array}{cc} 0 & \mathbf{I} \\ \mathbf{G}_1 & \mathbf{G}_2 \end{array}\right] \left[\begin{array}{c} \mathbf{x} \\ \mathbf{y} \end{array}\right] = \lambda \left[\begin{array}{c} \mathbf{x} \\ \mathbf{y} \end{array}\right].$$

The solution of the above eigenproblem is a linear $2n \times 2n$ system and can be solved using any local or global eigensolvers from the previous section.

10.5 Arnoldi Method:

Non-Symmetric Eigenproblems

In the previous chapter we presented the Arnoldi iteration which transforms a non-symmetric matrix \mathbf{A} to a Hessenberg matrix \mathbf{H} . In fact, the Arnoldi iteration can go part way; if the matrix \mathbf{A} has order n then the matrix \mathbf{H} can have order m with m < n, and we denote it by \mathbf{H}_m .

We now address the question of how exactly we locate the eigenvalues of \mathbf{A} ? We need to find first the eigenvalues of \mathbf{H}_m – that is why we bothered with this Hessenberg reduction. To this end, we can employ a QR iteration for \mathbf{H}_m to find the eigenpairs $(\lambda_i^{(m)}, \mathbf{y}_i^{(m)})$. However, these eigenvalues are only approximations to eigenvalues of \mathbf{A} . The accuracy increases as m increases, and the convergence rate is (approximately) linear for this method although not always; rigorous analysis is difficult to perform, see [79].

To assess the convergence of the method we need the residual. A good and cheap way to compute the residual is to consider the magnitude of the last component of the eigenvector $\mathbf{y}_i^{(m)}$, or more specifically the quantity

$$h_{m+1,m}|\mathbf{e}_m^T \mathbf{y}_i^{(m)}| = \| (\mathbf{A} - \lambda_i^{(m)} \mathbf{I}) \mathbf{r}_i^{(m)} \|_2$$

$$(10.2)$$

where $\mathbf{r}_{i}^{(m)}$ is the so-called Ritz approximate eigenvector defined by

$$\mathbf{r}_i^{(m)} = \mathbf{V}_m \mathbf{y}_i^{(m)}$$
 .

The above equality is based on the equation

$$(\mathbf{A} - \lambda_i^{(m)} \mathbf{I}) \mathbf{r}_i^{(m)} = h_{m+1,m} \mathbf{e}_m^T \mathbf{y}_i^{(m)} \mathbf{v}_{m+1},$$

which is derived from the fundamental Arnoldi relationship, namely

$$\mathbf{A}\mathbf{V}_m = \mathbf{V}_m \mathbf{H}_m + h_{m+1,m} \mathbf{v}_{m+1} \mathbf{e}_m^T.$$

Multiplying the above by $\mathbf{y}_i^{(m)}$ and utilizing the fact that $\mathbf{H}_m \mathbf{y}_i^{(m)} = \lambda_i^{(m)} \mathbf{y}_i^m$, we have that

$$\mathbf{A} \underbrace{\mathbf{V}_m y_i^{(m)}}_{\mathbf{r}_i^{(m)}} - \lambda_i^{(m)} \underbrace{\mathbf{V}_m \mathbf{y}_i^{(m)}}_{\mathbf{r}_i^{(m)}} = h_{m+1,m} \mathbf{v}_{m+1} \mathbf{e}_m^T \mathbf{y}_i^{(m)}.$$

Taking the L_2 norm of both sides and using the fact that $\|\mathbf{v}_{m+1}\|_2 = 1$ we obtain equation (10.2).

10.6 Available Software for Eigensolvers

The basic algorithms we have presented in this chapter are relatively easy to program but more sophisticated versions with respect to preconditioning, restarts, orthogonalization, and parallel implementation are available in free software in

• www.netlib.org

Here we briefly mention some of the most popular packages.

- For reduction of a dense matrix to a tridiagonal form using Householder transformations, the routines **xSYTRD** and **HETRD** for real and complex matrices are available in LAPACK. The corresponding routines in ScaLAPACK are **PxSYTRD** and **PxHETRD**.
- For various QR eigensolvers the routines **xSTEQR** and **xSTERF** are available in LAPACK.
- The parallel divide-and-conquer eigensolver we presented in this chapter is implemented in the routine **xSTEVD** of LAPACK. It is much faster than xSTEQR but requires more memory.
- The Lanczos algorithm is available in packages implemented by various authors. For example, the code **LANCZOS** presented in [19] implements the version without orthogonalization; the code **LANZ** in [61] involves partial orthogonalization; and the codes **LANSO** and **PLANSO** for the serial and parallel version, respectively produced by researchers at UC Berkeley involve periodic orthogonalization.

- For Arnoldi eigensolvers the package **ARPACK**, which stands for ARnoldi PACKage, is available in both Fortran as well as in C++; the latter is called ARPACK++. Specifically, **ARPACK++** has a good C++ interface to the original ARPACK codes so that the C++ users do not have to deal with the reverse communication interface problem that characterize the Fortran routines of ARPACK.
- For generalized eigenproblems, a simple driver routine **xSYGV** exists in LAPACK that is based on the QR method. A more sophisticated version **xSYGVX** chooses from a number of eigensolvers based on efficiency for a particular problem. For the parallel divide-and-conquer method the routine **xSYGVD** exists which is based on the simple driver of LAPACK.

10.7 Homework Problems

- 1. Show that the matrix $\mathbf{A} = \mathbf{L}\mathbf{U}$, where \mathbf{L} is a unit lower triangular and \mathbf{U} is upper triangular, has the same eigenvalues with the matrix $\mathbf{B} = \mathbf{U}\mathbf{L}$.
- 2. The power method to finding eigenvalues is applicable to (choose one)
 - (a) Symmetric matrices.
 - (b) Symmetric, positive-definite matrices.
 - (c) General matrices.
- 3. Write a C++ code to test the convergence rate of the inverted shifted power method applied to the matrix resulted from the two-dimensional discretization of:
 - (a) $\nabla^2 u = \lambda u$ (symmetric)
 - (b) $\nabla^2 u u_x u_y = \lambda u$ (non-symmetric)

on a square domain $[0,1] \times [0,1]$ with second-order finite differences. We assume that the boundary conditions are all homogeneous. Plot the error in predicting the minimum eigenvalue versus the iteration number to estimate the convergence rate.

- 4. Prove that the QR iteration maintains the Hessenberg structure, which is a key step in applying efficiently the QR algorithm.
- 5. Show that the eigenvalues of the tridiagonal matrix defined by

$$d_{ii} = 2, \quad i = 1, \dots, n$$

 $d_{i,i-1} = -1, \quad i = 2, \dots, n$
 $d_{i-1,i} = -1, \quad i = 2, \dots, n$

are given by

$$\lambda_j = 2 - 2\cos(j\pi/(n+1))$$

with corresponding eigenvectors

$$\mathbf{v}_j = [\sin(j\pi/(n+1), \sin(2j\pi/(n+1), \dots, \sin(nj\pi/(n+1))]^T.$$

Then, use the power method and the inverse power method to compute the maximum and minimum eigenvalues for n = 50. What is the convergence rate of the two methods for this problem?

6. Construct a QR eigensolver for a symmetric positive definite matrix **A** using the Cholesky decomposition. Estimate the operation count and write a C++ code for the matrix resulted from the discretization of

$$\nabla^2 u - u = \lambda u$$
, in $[0, 1] \times [0, 1]$.

Is your theoretical estimate consistent with what you find experimentally by timing your solver. (Use a sufficiently large number of grid points, e.g. a grid of 100×100 .)

- 7. Consider a symmetric Toepliz matrix with entries $a_k = 1/k$, k = 1, ..., n. We want to apply Rayleigh quotient iteration to it. First, show that the shifted matrix $(\mathbf{A} \sigma_k \mathbf{I})$, where σ_k is the Rayleigh quotient remains a Toepliz matrix. Then compute the largest three and the smallest three eigenvalues of \mathbf{A} for n = 25, 50 and 100. What is the convergence rate you observe?
- 8. Consider five symmetric Toeplitz matrices **A** with entries given by (k = 1, ..., n)

$$a_k^{(1)} = 1/k;$$
 $a_k^{(2)} = 1/\sqrt{k};$ $a_k^{(3)} = 1/k^2;$ $a_k^{(4)} = k;$ $a_k^{(5)} = \cos k/k,$

and construct corresponding circulant matrices \mathbf{C} employing different number of Toeplitz diagonals. Compute the three largest and three smaller eigenvalues of the matrix $\mathbf{C}^{-1}\mathbf{A}$ and compare with the corresponding eigenvalues of \mathbf{A} . Use $n=20,30,\ldots,100$. What do you observe?

9. Consider the finite difference discretization of the Helmholtz eigenvalue problem

$$\nabla^2 u - 4u = \lambda u$$

in the domain $[0,1] \times [0,1]$. Apply the QR and LR (based on LU) eigensolvers and compute the CPU time required to obtain the entire eigenspectrum. Consider a grid with $N \times N$ and N = 128,256 and 512.

10. Let **T** be a block-tridiagonal matrix of order $n^2 \times n^2$ with **A** of order $n \times n$ in the diagonal and **B** also of order $n \times n$ in the two off-diagonals. Assuming that we decompose **A** and **B** as follows

$$\mathbf{Q}\mathbf{A}\mathbf{Q}^T = diag(a_1, a_2, \dots, a_n)$$

 $\mathbf{Q}\mathbf{B}\mathbf{Q}^T = diag(b_1, b_2, \dots, b_n)$,

then find analytical expressions for the n^2 eigenpairs of **T** in terms of a_i , b_i and **Q**.

11. Consider the $n \times n$ matrix **A** the entries of which are computed from

$$a_{ij} = \frac{ij}{(i+j)^2} \,.$$

Compute the eigenvalues and eigenvectors of this matrix using the Lanczos algorithm for n = 5, 10 and 20, at different levels of accuracy.

- 12. Prove that the Householder matrix is symmetric.
- 13. Show how a Householder matrix **H** of size $n \times n$ times a vector **x** of size n can be accomplished in $\mathcal{O}(n)$ operations. Explain how this implies that an $n \times n$ matrix **A** can be pre-multiplied by **H** in $\mathcal{O}(n^2)$ operations.
- 14. Using the MPI program presented in this chapter, write an MPI program which accomplishes the full Householder deflation to obtain all the eigenvalues.

- (a) First explain how the rows of the matrix **A** should be partitioned to maximize parallel performance (block partitioning, cyclic partitioning, etc)? Show diagrammatically why one partitioning is preferable over another.
- (b) For a 6×6 system and 2 processes, show by diagram the communication pattern that you would use for the full deflation.

Hint: You will need to use the variable length versions of $MPI_Allgather$ and $MPI_AlltoAll$ ($MPI_Allgatherv$ and $MPI_Alltoallv$).

- 15. Write a simple MPI program using $MPI_Alltoall$ to transpose a matrix **A** of size N. Using the MPI_Wtime function, perform the following timings:
 - (a) For a fixed number of processes, P=2 and P=4, plot the time to transpose matrices of size n=10,20,30,40 and 100. What scaling do you see? Document the machine on which you were running, the number of processors the machine has, etc.
 - (b) For a fixed matrix size n = 100, plot the time to transpose a matrix on P = 2, 4 and 8 processors. What scaling do you encounter?
- 16. In the parallel QR eigensolver presented in the text, each process requires storage for full Q, Q1 and Q2 arrays. Modify the program so that only the minimal amount of memory is used.
- 17. Write an C++/MPI program which solves the secular equation in parallel.
- 18. In the parallel QR eigensolver presented in the text, a fan-in algorithm was used to combine the solved subproblems. Suppose, however, that instead of using a fan-in algorithm, we solve the problem as follows: Let a master node partition the problem into P subproblems, having each process solve its subproblem (just as in the program we presented). Instead of fan-in, have the master node be in charge of the combination effort, using the other processes to solve in parallel the secular equation and the matrix multiplications.
 - (a) Implement the algorithm just described.
 - (b) Which algorithm is faster, the fan-in or the newly proposed algorithm? For matrices of size 50 and 100, perform timing tests on 2,4, and 8 processors. What can you conclude?

Appendix A

A. C++ Basics

A.1 Compilation Guide

For the purposes of this compilation example, we will assume that we are using the GNU g++ compiler to compile a C++ program we have written contained within the file *myprog.cpp*. In the following examples, the argument following the '-o' flag designates the file name to be used for the output. If no '-o' option is specified, most compilers default to using the name "a.out". We now present several different programming scenarios:

• No user-defined libraries or user-defined header files are needed, and no special system libraries (such as those associated with *math.h* are needed):

```
g++ -o myprog myprog.cpp
```

• No user-defined libraries or user-defined header files are needed, but the special system library corresponding to *math.h* is needed:

```
g++ -o myprog myprog.cpp -lmath
```

• User-defined libraries, user-defined header files, and the special system library corresponding to *math.h* are needed:

```
g++ -o myprog myprog.cpp -I/users/kirby/includes -L/users/kirby/libs -lSCmathlib -lmath
```

The string following the '-I' flag designates the location of the user-defined header files to be included. The string following the '-L' flag designates the location of the user-defined libraries to be included. The string '-lSCmathlib' links the program with the user-defined library we created, and the string '-lmath' links the program with the system math library corresponding to math.h.

A.2 C++ Basic Data Types

C++ has a set of standard data types representing the most common basic storage units of the language. They can be partitioned as follows:

- 1. A Boolean type (bool)
- 2. Integer types (such as *int*)
 - (a) int
 - (b) unsigned int
 - (c) signed int
 - (d) short int (short-handed declaration is short)
 - (e) long int (short-handed declaration is long)
- 3. Floating-point types (such as float)
 - (a) float (single precision)
 - (b) double (double precision)
 - (c) long double (extended double precision)
- 4. Character types (such as *char*)
 - (a) char
 - (b) unsigned char
 - (c) signed char
- 5. A type void which represents the absence of information
- 6. Types constructed from the above types
 - (a) Pointer types (such as int*)
 - (b) Reference types (such as int&)
 - (c) Array types (such as *int*[])

We refer the reader to [86] for a complete description and usage explanation of the data types listed above.

A.3 C++ Libraries

A.3.1 Input/Output Library – iostream.h

Below we present three additional operations found within the C++iostream system library which were not discussed previously. For more information about the contents of iostream.h, we refer the reader to [86].

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• **cin.get(ch)** - allows the user to obtain from standard input the next character (including white space and the end-of-line character). This function takes in its argument list a variable of type *char*, into which it places the character obtained from standard input (i.e. this variable is passed by reference).

- cin.eof() returns non-zero (true) if end-of-file has been encountered in the standard input stream.
- **cout.put(ch)** allows the user to print to standard output a single character. This function takes in its argument list a variable of type *char*, the contents of which are printed to standard output.

A.3.2 Input/Output Manipulation Library – iomanip.h

Below is a list of a few of the input/output manipulation operations which can be found in the C++iomanip system library. For more information about the contents of iomanip.h, we refer the reader to [86].

- **cout** << **setiosflags(ios::flag)** sets/resets the I/O flag. The designator *flag* may be one of the following:
 - 1. fixed floating-point output will be printed in fixed format (not using scientific notation).
 - 2. scientific floating-point output will be printed in scientific notation.
 - 3. showpoint enforces that decimal points and trailing zeroes are displayed.
- cout << setprecision(i) sets the number of decimal places of accuracy to be used when printing floating point (float/double) values. This function takes as input an integer variable representing the number of digits to be used.
- **cout** << **setw(i)** sets the field width to be used for the next item printed. This function takes as input an integer variable which designates the number of characters used to determine the field width.
- **cout** << **endl** indicates the ending of one line of output, and the initiation of a new line of output.

A.3.3 Mathematics Library – math.h

Below is a list of some of the mathematical operations which can be found in the C++ math library. For more information about the contents of math.h, we refer the reader to [86].

```
sine of x
\sin(x)
\cos(x)
                         cosine of x
                         tangent of x
tan(x)
                         \sin^{-1}(x) in range [-\pi/2, \pi/2], x \in [-1, 1].
asin(x)
                         \cos^{-1}(x) in range [0, \pi], x \in [-1, 1].
acos(x)
                         \tan^{-1}(x) in range [-\pi/2, \pi/2].
atan(x)
                         [\tan^{-1}(y/x) \text{ in range } [-\pi, \pi].
atan2(y, x)
\sinh(x)
                         hyperbolic sine of x
                         hyperbolic cosine of x
\cosh(x)
                         hyperbolic tangent of x
tanh(x)
                         exponential function e^x
\exp(x)
\log(x)
                         natural logarithm ln(x), x > 0.
\log 10(x)
                         base 10 logarithm \log_{10}(x), x > 0.
pow(x, y)
                         x^{y}. A domain error occurs if x=0 and y<0, or if
                         x < 0 and y is not an integer.
                         \sqrt{x}, x > 0.
\operatorname{sqrt}(x)
                         smallest integer not less than x, as a double.
cdil(x)
floor(x)
                         largest integer not greater than x, as a double.
fabs(x)
                         absolute value |x|
                         x \cdot 2^n
dexp(x, n)
frexp(x, int *exp)
                         splits x into a normalized fraction in the interval
                         [1/2, 1], which is returned, and a power of 2, which is
                         stores in *exp. If x is zero, both parts of the result
                         are zero.
modf(x, double *ip)
                         splits x into integral and fractional parts, each with the
                         same sign as x. It stores the integral part in *ip, and
                         returns the fractional part.
fmod(x, y)
                         floating-point remainder of x/y, with the same sign as
                         x. If y is zero, the result is implementation-defined.
```

A.4 Operator Precedence

Table A.1 provides the C++ convention used for operator precedence. This list is ordered from highest to lowest precedence.

$A.5 \quad C++ \text{ and BLAS}$

Many of the C++ programs presented in this book can be re-programmed to use BLAS for almost all of the mathematics operations. We did not originally program things using BLAS for pedagogical reasons; we attempted to use every programming example to expand the reader's familiarity with C++ constructs.

When combining C++ and BLAS, the reader should consult the following website:

http://www.netlib.org/blas/

OPERATIONS	ASSOCIATIVITY
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	left to right
! \sim ++ + - * & (type) size of	right to left
* / %	left to right
+ -	left to right
<< >>	left to right
< <= > >=	left to right
== !=	left to right
&	left to right
^	left to right
	left to right
& &	left to right
	left to right
?:	right to left
$= \ += \ -= *= /= \%= \& = \hat{\ } = = <<= >>= $	right to left
,	left to right

Table A.1: Unary +, -, and * have higher precedence than the binary forms.

which contains explicit documentation for linking C programs to BLAS. One modification that C++ requires is that BLAS function declarations should be done as follows:

```
extern "C" {
  void daxpy_(int*, double*, double *, int*, double *, int*);
}
```

which notifies the compiler to use the C programming language function name convention.

Appendix B

B. MPI Basics

B.1 Compilation Guide

For the purposes of this compilation example, we will assume that we are using the GNU g++ compiler to compile a C++ program we have written contained within the file *myprog.cpp*. We will also assume that the machine on which you are trying to compile is a parallel machine with some version of MPI installed. You will need to contact your system administrator to find out the exact version of MPI that is available and the paths on your local architecture. In the following examples, the argument following the '-o' flag designates the file name to be used for the output. If no '-o' option is specified, most compilers default to using the name "a.out". We now present several different programming scenarios:

No user-defined libraries or user-defined header files are needed, and no special system libraries (such as those associated with math.h are needed) other than the MPI libraries:

```
g++ -o myprog myprog.cpp -lmpi
```

• No user-defined libraries or user-defined header files are needed, but the special system library corresponding to *math.h* is needed along with the MPI libraries:

```
g++ -o myprog myprog.cpp -lmath -lmpi
```

• User-defined libraries, user-defined header files and the special system library corresponding to *math.h* are needed along with the MPI libraries:

```
g++ -o myprog myprog.cpp -I/users/kirby/includes -L/users/kirby/libs -lSCmathlib -lmath -lmpi
```

The string following the '-I' flag designates the location of the user-defined header files to be included. The string following the '-L' flag designates the location of the user-defined libraries to be included. The string '-ISCmathlib' links the program with the user-defined library we created, and the string '-lmath' links the program with the system math library corresponding to math.h. The string '-lmpi' links the MPI libraries.

You will need to contact your system administrator to verify the exact command used on your computing architecture to run an MPI program. In general, most architectures execute MPI programs in a fashion similar to the following:

mpirun -np 4 myprog

where mpirun is a special program used for starting execution on the parallel machine, '-np 4' indicates the number of processes requested, and *myprog* denotes the program executable compiled as discussed above.

B.2 MPI Commands

Below we present an extended list of the MPI datatypes, MPI reduction operations, and MPI function declarations.

B.2.1 Predefined Variable Types in MPI

MPI datatype	$C\ datatype$
MPI_CHAR	signed char
MPI_SHORT	singed short int
MPI_INT	signed int
MPI_LONG	singed long int
MPI_UNSIGNED_CHAR	unsigned char
MPI_UNSIGNED_SHORT	unsigned short int
MPI_UNSIGNED	unsigned int
MPI_UNSIGNED_LONG	unsigned long int
MPI_FLOAT	float
MPI_DOUBLE	double
MPI_LONG_DOUBLE	long double
MPI_BYTE	
MPI_PACKED	

B.2.2 Predefined Reduction Operators in MPI

Operation Name	Meaning
MPI_MAX	Maximum
MPI_MIN	Minimum
MPI_SUM	Sum
MPI_PROD	Product
MPI_LAND	Logical and
MPI_BAND	Bitwise and
MPI_LOR	Logical or
MPI_BOR	Bitwise or
MPI_LXOR	Logical exclusive or
MPI_BXOR	Bitwise exclusive or
MPI_MAXLOC	Maximum and location of maximum
MPI_MINLOC	Minimum and location of minimum

B.2.3 MPI Function Declarations

```
    MPI_Get_count(
    MPI_Status* status /* in */...
    MPI_Datatype datatype /* in */...
    int* count /* out */...
```

• MPI_Recv

```
int MPI_Recv(
        void*
                       message
                                      out
        int
                        count
                                      in
        MPI_Datatype
                       datatype
                                      in
                                            */,
        int
                       source
                                      in
        int
                                      in
                        tag
        MPI_Comm
                                      in
                        comm
        MPI\_Status*
                                      out
                       status
```

• MPI_Send

```
int MPI_Send(
                                                 */,
         void*
                            message
                                            in
                                                 */,
         int
                            count
                                            in
         MPI_Datatype
                                            in
                           datatype
                                            in
         int
                            \operatorname{dest}
         int
                                            in
                            tag
         MPI_Comm
                                            in
                            comm
```

• MPI_Bsend

```
int MPI_Bsend(
                                               */,
           void*
                            message
                                          in
           int
                            count
                                          in
           MPI_Datatype
                           datatype
                                          in
                                              */,
           int
                            dest
                                          in
           int
                                          in
                            tag
           MPI_Comm
                                          in
                            comm
• MPI_Rsend
   int MPI_Rsend(
           void*
                            message
                                          in
           int
                            count
                                          in
           MPI_Datatype
                            datatype
                                          in
                            dest
                                          in
           int
           int
                                          in
                            tag
           MPI_Comm
                            comm
                                          in
• MPI_Ssend
   int MPI_Ssend(
           void*
                            message
                                          in
           int
                            count
                                          in
           MPI_Datatype
                            datatype
                                          in
           int
                            dest
                                          in
                                              */,
           int
                            tag
                                          in
           MPI\_Comm
                                          in
                            comm
• MPI_Buffer_attach
   int MPI_Buffer_attach(
           void*
                                       in */,
                           buffer
           int
                           size
                                       in
• MPI_Buffer_detach
   int MPI_Buffer_detach(
           void*
                           buffer\_address
                                               out
           int
                           size_ptr
                                               out
• MPI_lbsend
   int MPI_Ibsend(
           void*
                            message
                                          in
           int
                            count
                                          in
           MPI_Datatype
                           datatype
                                          in
           int
                            dest
                                          in
           int
                                          in
                            tag
                                                */,
           MPI_Comm
                                          in
                            comm
           MPI_Request*
                            request
                                          out
```

• MPI_Irecv

```
int MPI_Irecv(
                                             */,
        void*
                         message
                                        out
        int
                         count
                                        in
                                             */,
        MPI_Datatype
                         datatype
                                        in
                         dest
                                        in
        int
                         tag
                                        in
        MPI_Comm
                                        in
                         comm
        MPI_Request*
                         request
                                        out
                                             */)
```

• MPI_Irsend

```
int MPI_Irsend(
        void*
                                              */,
                         message
                                        in
        int
                         count
                                        in
        MPI_Datatype
                         datatype
                                        in
                                              ^*/,
        int
                         dest
                                        in
        int
                         tag
                                        in
        MPI_Comm
                                        in
                         comm
                                              */)
        MPI_Request*
                                        out
                         request
```

• MPI_Isend

```
int MPI_Isend(
                                              */,
        void*
                         message
                                        in
                                              */,
        int
                         count
                                        in
        MPI_Datatype
                         datatype
                                        in
        int
                         dest
                                        in
                                              ^*/,
        int
                                        in
                         tag
        MPI_Comm
                                        in
                         comm
                                              */)
        MPI_Request*
                         request
                                        out
```

• MPI_Issend

```
int MPI_Isend(
        void*
                         message
                                        in
        int
                         count
                                        in
                                              */,
        MPI_Datatype
                         datatype
                                        in
        int
                         dest
                                        in
        int
                                        in
                         tag
                                              */,
        MPI_Comm
                         comm
                                        in
                                              */)
        MPI_Request*
                         request
                                        out
```

ullet MPI_Request_free

• MPI_Test

• MPI_Testall

• MPI_Testany

```
int MP_Testany(
                        array_size
         int
                                                in
         MPI_Request
                        requests[]
                                                in/out
                                                        */,
         int*
                        completed_index
                                                out
         int*
                        flag
                                                out
         MPI_Status*
                        status
                                                out
```

• MPI_Testsome

```
int MP_Testsome (
         int
                        array_size
                                                in
         MPI\_Request
                        requests[]
                                                in/out
         int*
                        completed_count
                                                         */,
                                                out
                        indices[]
         int
                                                out
         MPI_Status
                        statuses[]
                                                out
```

• MPI_Wait

• MPI_Waitall

MPI_Waitany

```
int MP_Waitany(

int array_size /* in */,

MPI_Request requests[] /* in/out */,

int* completed_index /* out */,

MPI_Status* status /* out */)
```

• MPI_Waitsome

```
int MP_Waitsome(
                                                        */,
         int
                        array_size
                                               in
         MPI_Request
                        requests[]
                                               in/out
         int*
                                                        */,
                        completed_count
                                               out
                        indices[]
         int
                                               out
                                                        */)
         MPI_Status
                        statuses[]
                                               out
```

• MPI_Cancel

• MPI_Iprobe

```
int MP_Iprobe(
        int
                       source
                                    in
        int
                       tag
                                    in
        MPI_Comm
                       comm
                                    in
                                /*
        int*
                       flag
                                    out
        MPI_Status*
                       status
                                    out
```

• MPI_Probe

• MPI_Test_cancelled

```
int MPI_Test_cancelled(

MPI_Status* status /* in */,
int* flag /* out */)
```

• MPI_Bsend_init

```
int MPI_Bsend_init(
        void*
                         message
                                        in
        int
                         count
                                        in
        MPI_Datatype
                         datatype
                                        in
        int
                         dest
                                        in
        int
                                        in
                         tag
                                             */,
        MPI_Comm
                                        in
                         comm
        MPI_Request*
                                             */)
                         request
                                        out
```

• MPI_Recv_init

```
int MPI_Recv_init(
                                                   */,
            void*
                              message
                                              out
                                                   */,
            int
                              count
                                              in
                                                    */,
            MPI_Datatype
                              datatype
                                              in
                                                   */,
            int
                              source
                                              in
                                                   */,
            int
                              tag
                                              in
                                                    */,
            MPI_Comm
                              comm
                                              in
                                                   \dot{*/})
            MPI_Request*
                              request
                                              out
• MPI_Rsend_inti
   int MPI_Rsend_init(
                                                   */,
            void*
                              message
                                              in
            int
                              count
                                                    (/,
                                              in
                                                   */,
            MPI_Datatype
                              datatype
                                              in
            int
                              dest
                                              in
            int
                              tag
                                              in
            MPI\_Comm
                                                    */,
                                              in
                              comm
                                                   */)
            MPI_Request*
                                              out
                              request
• MPI_Send_int
   int MPI_Send_init(
                                                    */,
            void*
                              message
                                              in
            int
                              count
                                              in
            MPI_Datatype
                              datatype
                                              in
                                                   ^{\ast ^{\prime }}/,
            int
                              dest
                                              in
                                                    */,
            int
                              tag
                                              in
                                                    */,
            MPI_Comm
                                              in
                              comm
            MPI_Request*
                                                    */)
                                              out
                              request
• MPI_Ssend_init
   int MPI_Ssend _init(
                                                   */,
            void*
                              message
                                              in
                                                   */,
            int
                              count
                                              in
                                                   ^{\ast ^{\prime }/,}
            MPI_Datatype
                              datatype
                                              in
            int
                              dest
                                              in
                                                    */,
            int
                                              in
                              tag
                                                   */,
            MPI_Comm
                              comm
                                              in
                                                   */)
            MPI_Request^*
                              request
                                              out
• MPI_Start
```

```
• MPI_Startall
```

```
int MPI_Startall(
    int array_size /* in */,
    MPI_Request requests[] /* in/out */)
```

• MPI_Sendrecv

```
int MPI_Sendrecv(
        void*
                         sendbuf
                                         in
        int
                         sendcount
                                         in
                         sendtype
        MPI_Datatype
                                         in
                         dest
                                         in
        int
        int
                         sendtag
                                         in
        void*
                         recvbuf
                                         out
        int
                         recvcount
                                         in
        MPI_Datatype
                         recvtype
                                         in
        int
                         source
                                         in
        MPI_Datatype
                         recvtag
                                         in
                                              */,
        MPI_Comm
                         comm
                                         in
                                               */)
        MPI_Status*
                         status
                                         out
```

• MPI_Address

```
int MPI_Address(
void* location /* in */,
MPI_Aint* address /* out */)
```

• MPI_Get_elements

```
int MPI_Get_elements(

MPI_Status* status /* in */

MPI_Datatype datatype /* in */

int* count /* out */)
```

• MPI_Type_commit

• MPI_Type_contiguous

```
int MPI_Type_contiguous(
    int count /* in */,
    MPI_Datatype oldtype /* in */
    MPI_Datatype* newtype /* out */)
```

• MPI_Type_extent

```
int MPI_Type_extent(

MPI_Datatype datatype /* in */

MPI_Aint* extent /* out */
```

```
• MPI_Type_free
```

MPI_Type_hindexed

```
int MPI_Type_hindexed(
int count /* in */,
int blocklengths[] /* in */,
MPI_Aint displacements[] /* in */,
MPI_Datatype oldtype /* in */
MPI_Datatype* newtype /* out */)
```

• MPI_Type_hvector

```
int MPI_Type_hvector(
        int
                          count
                                           in
        int
                          blocklength
                                           in
        MPI_Aint
                          stride
                                           in
        MPI_Datatype
                          oldtype
                                           in
        MPI_Datatype*
                         newtype
                                           out
```

• MPI_Type_indexed

```
int MPI_Type_indexed(
                                                      */,
        int
                          count
                                                      */,
                          blocklength[]
        int
                                                in
                          displacements[]
        int
                                                in
        MPI_Datatype
                          oldtype
                                                in
        MPI_Datatype*
                          newtype
                                                out
```

• MPI_Type_1b

• MPI_Type_size

```
int MPI_Type_size(

MPI_Datatype datatype /* in */
int* size /* out */)
```

• MPI_Type_struct

```
int MPI_Type_stuct(

int count /* in */,

int blocklengths[] /* in */,

MPI_Aint displacements[] /* in */,

MPI_Datatype types[] /* in */

MPI_Datatype* newtype /* out */)
```

• MPI_Type_ub

• MPI_Type_vector

```
int MPI_Type_vector(
        int
                          count
                                            in
        int
                          blocklength
                                            in
        int
                          stride
                                            in
        MPI_Datatype
                          oldtype
                                            in
        MPI_Datatype*
                          newtype
                                            out
```

MPI_Pack

```
int MPI_Pack(
        void*
                         inbuf
                                            in
        int
                         incount
                                            in
        MPI_Datatype
                         datatype
                                            in
        void*
                         pack_buf
                                            out
        int
                         pack_buf_size
                                            in
        int*
                         position
                                            in/out
        MPI_Comm
                         comm
                                            in
```

• MPI_Pack_size

```
int MPI_Pack_size(

int incount /* in */,

MPI_Datatype datatype /* in */

MPI_Comm comm /* in */,

int* size /* out */)
```

• MPI_Unpack

```
int MPI_Unpack(
                                                    */,
        void*
                         pack_buf
                                            in
        int
                         pack_buf_size
                                            in
        int*
                         position
                                            in/out
        void*
                         outbuf
                                            out
        int
                         outcount
                                            in
        MPI_Datatype
                         datatype
                                            in
        MPI_Comm
                         comm
                                            in
```

• MPI_Barrier

```
int MPI_Barrier(

MPI_Comm comm /* in */)
```

• MPI_Bcast

```
int MPI_Bcast(
                                                */,
        void*
                         buffer
                                        in/out
        int
                         count
                                        in
                                                */,
                         datatype
        MPI_Datatype
                                        in
                         root
                                        in
        MPI_Comm
                                                */)
                         comm
                                        in
```

• MPI_Allgather

```
int MPI_Allgather(
        void*
                         sendbuf
                                         in
        int
                         sendcount
                                         in
        MPI_Datatype
                        sendtype
                                         in
        void*
                         recvbuf
                                         out
                                              */
        int
                         recvcount
                                         in
                                              */,
        MPI_Datatype
                        recvtype
                                         in
                                              */)
        MPI_Comm
                                         in
                         comm
```

• MPI_Allgatherv

```
int MPI_Allgatherv(
                                                        */,
         void*
                          sendbuf
                                                  in
         int
                          sendcount
                                                  in
         MPI_Datatype
                          sendtype
                                                  in
         void*
                          \operatorname{recvbuf}
                                                  out
         int
                          recvcounts[]
                                                  in
         int
                          displacements[]
                                                  in
         MPI_Datatype
                                                  in
                          recvtype
                                                        */)
         MPI_Comm
                                                  in
                           comm
```

• MPI_Alltoall

```
int MPI_Alltoall(
        void*
                        sendbuf
                                         in
        int
                        sendcount
                                         in
                        sendtype
        MPI_Datatype
                                         in
        void*
                        recvbuf
                                         out
        int
                        recvcount
                                              */
                                         in
        MPI_Datatype
                        recvtype
                                         in
        MPI_Comm
                                              */)
                         comm
                                         in
```

• MPI_Alltoallv

```
int MPI_Allgatherv(
        void*
                         sendbuf
                                                    in
                         sendcounts[]
        int
                                                    in
                         send_displacements[]
        int
                                                    in
        MPI_Datatype
                         sendtype
                                                    in
        void*
                         recvbuf
                                                    out
        int
                         recvcounts[]
                                                    in
                         recv_displacements[]
        int
                                                    in
        MPI_Datatype
                         recvtype
                                                    in
                                                          */)
        MPI_Comm
                         comm
                                                    in
```

• MPI_Gather

```
int MPI_Gather(
                                               */,
        void*
                         sendbuf
                                         in
        int
                         sendcount
                                               */,
                                         in
        MPI_Datatype
                        sendtype
                                         in
        void*
                                               */,
                         recvbuf
                                         out
        int
                         recvcounts
                                         in
                                               */,
        MPI_Datatype
                        recvtype
                                         in
        MPI\_Comm
                         comm
                                         in
```

• MPI_Gatherv

```
int MPI\_Gatherv(
        void*
                         sendbuf
                                               in
                         sendcount
        int
                                               in
        MPI_Datatype
                         sendtype
                                               in
        void*
                         recvbuf
                                               out
        int
                         recvcounts[]
                                               in
                                                     */,
                         displacements[]
        int
                                               in
        MPI_Datatype
                         recvtype
                                               in
                         root
                                               in
        MPI_Comm
                         comm
                                               in
```

• MPI_Scatter

```
int MPI_Scatter(
        void*
                         sendbuf
                                         in
                         sendcount
        int
                                         in
        MPI_Datatype
                        sendtype
                                         in
        void*
                         recvbuf
                                         out
        int
                         recvcount
                                         in
        MPI_Datatype
                        recvtype
                                         in
                                              */,
        int
                         root
                                         in
        MPI_Comm
                                              */)
                         comm
```

• MPI_Scatterv

```
int MPI_Scatterv(
           void*
                           sendbuf
                                                 in
                           sendcounts[]
           int
                                                 in
                           displacements[]
           int
                                                 in
           MPI_Datatype
                           sendtype
                                                 in
           void*
                           recvbuf
                                                 out
           int
                           recvcount
                                                 in
                                                      */,
           MPI_Datatype
                           recvtype
                                                in
                                                      */,
           int
                                                 in
                           root
           MPI_Comm
                           comm
                                                in
• MPI_Allreduce
   int MPI_Allreduce(
           void*
                           operand
                                          in
           void*
                           result
                                          out
           int
                           count
                                               */,
                                          in
           MPI_Datatype
                           datatype
                                          in
           MPI_Op
                           operator
                                          in
           MPI_Comm
                           comm
                                          in
• MPI_Op_create
   int MPI_Op_create(
           MPI_User_function*
                                function
                                               in
                                                    */,
           int
                                commute
                                               in
           MPI_Op
                                operator
                                               out
• MPI_Op_free
   int MPI_Op_free(
           MPI_Op*
                      operator
                                /* in/out
                                            */)
• MPI_Reduce
   int MPI_Reduce(
                                            */,
           void*
                         operand
                                       in
           void*
                         result
                                       out
           int
                                       in
                         count
           MPI_Op
                         operator
                                       in
           int
                         root
                                       in
           MPI_Comm
                        comm
                                       in
• MPI_Reduce_Scatter
   int MPI_Reduce_scatter(
           void*
                            operand
                                              in
           void*
                            recvbuf
                                              out
           int
                            recvcounts[]
                                              in
           MPI_Datatype
                            datatype
                                              in
           MPI_Op
                            operator
                                              in
           MPI_Comm
                            comm
                                              in
```

```
• MPI_Scan
```

```
int MPI_Scan(
                                             */,
        void*
                        operand
                                       in
        void*
                        result
                                       out
                                             */,
        int
                        count
                                       in
        MPI_Datatype
                        datatype
                                       in
        MPI_Op
                        operator
                                       in
        MPI_Comm
                        comm
                                       in
```

• MPI_Comm_group

```
int MPI_comm_group(

MPI_Comm comm /* in */,

MPI_Group* group /* out */)
```

• MPI_Group_compare

```
int MPI_Group_compare(

MPI_Group group1 /* in */,

MPI_Group group2 /* in */,

int* result /* out */)
```

• MPI_Group_difference

```
int MPI_Group_difference(

MPI_Group group1 /* in */,

MPI_Group group2 /* in */,

MPI_Group* newgroup /* out */)
```

• MPI_Group_excl

• MPI_Group_free

• MPI_Group_incl

• MPI_Group_intersection

```
int MPI_Group_intersection(

MPI_Group group1 /* in */,

MPI_Group group2 /* in */,

MPI_Group* newgroup /* out */)
```

• MPI_Group_range_excl

• MPI_Group_range_incl

```
int MPI_Group_range_incl(
                                                   */,
        MPI_Group
                            group
                                              in
                                                   */,
        int
                            n
                                              in
                                                   */,
        int
                            ranges [][3]
                                              in
        MPI_Group*
                            newgroup
                                              out
```

• MPI_Group_rank

• MPI_Group_size

• MPI_Group_translate_ranks

```
int MPI_Group_translate_ranks(
        MPI_Group
                                 group1
                                                in
        int
                                                in
                                 n
        int
                                 ranks[]
                                                in
                                                      */,
                                                     */,
        MPI_Group*
                                 group2
                                                in
        int
                                 ranks2[]
                                                out
```

• MPI_Group_union

```
int MPI_Group_union(

MPI_Group group1 /* in */,

MPI_Group* group2 /* in */,

MPI_Group* newgroup /* out */)
```

```
• MPI_Comm_compare
```

• MPI_comm_create

```
int MPI_Comm_create(

MPI_Comm comm1 /* in */,

MPI_Group new_group /* in */,

MPI_Comm* new_comm /* out */)
```

• MPI_Comm_dup

```
int MPI_Comm_dup(

MPI_Comm comm /* in */,

MPI_Comm* new_comm /* out */)
```

• MPI_Comm_free

• MPI_Comm_rank

```
\begin{array}{cccc} \mathrm{int} \ \mathrm{MPI\_Comm\_rank}( \\ \mathrm{MPI\_Comm} & \mathrm{comm} & /* & \mathrm{in} & */, \\ \mathrm{int}^* & \mathrm{result} & /* & \mathrm{out} & */) \end{array}
```

• MPI_Comm_size

```
int MPI_Comm_size(

MPI_Comm comm /* in */;

int* size /* out */
```

• MPI_Comm_split

```
int MPI_Comm_split(

MPI_Comm old_comm /* in */,
int split_key /* in */,
int rank_key /* in */,
MPI_Comm* new_comm /* out */)
```

• MPI_Comm_remote_group

```
int MPI_Comm_remote_group(

MPI_Comm comm /* in */,

MPI_Group* group /* out */)
```

• MPI_Comm_remote_size

```
int MPI_Comm_remote_size(

MPI_Comm comm /* in */,

int* size /* out */)
```

• MPI_Comm_test_inter

```
int MPI_Comm_test_inter(

MPI_Comm comm /* in */,
int* flag /* out */)
```

• MPI_Intercomm_create

```
int MPI_Intercomm_create(
        MPI_Comm
                           loca_comm
                                              in
        int
                           local\_leader
                                              in
        MPI_Comm
                           peer_comm
                                              in
        int
                           remote_leader
                                              in
        int
                           tag
                                              in
        MPI_Comm*
                                              out
                           intercomm
```

• MPI_Intercomm_merge

```
int MPI_Intercomm_merge(

MPI_Comm intercomm /* in */,
int high /* in */,
MPI_Comm* intracomm /* out */)
```

• MPI_Attr_delete

• MPI_Attr_get

• MPI_Attr_put

```
int MPI_Attr_put(

MPI_Comm comm /* in */,
int keyval /* in */,
void* attribute /* in */)
```

• MPI_Keyval_create

```
int MPI_Keyval_create(

MPI_Copy_function* copy_fn /* in */,

MPI_Delete_function* delete_fn /* in */,

int* keyval /* out */,

void* extra_arg /* in */)
```

```
• MPI_Keyval_free
```

```
int MPI_Keyval_free(
int* keyval /* in/out */,
```

• MPI_Topo_test

```
int MPI_Topo_test(

MPI_Comm comm /* in */,

int* top_type /* out */)
```

• MPI_Cart_coords

• MPI_cart_create

```
int MPI_Cart_create(
        MPI_Comm
                       old\_comm
                                        in
                       ndims
        int
                                        in
                       dims[]
        int
                                        in
                       periods[]
        int
                                        in
        int
                       reorder
                                        in
        MPI_Comm*
                       cart_comm
                                        out
```

• MPI_Cartdim_get

```
int MPI_Cartdim_get(

MPI_Comm comm /* in */,
int* ndims /* out */)
```

• MPI_Cart_get

```
int MPI_Cart_get(
                                                 */,
         MPI_Comm
                       comm
                                           in
                       \max\_dims
         int
                                           in
                       \dim S[]
         int
                                           out
                       periods[]/*
         int
                                      out
         int
                       coords[]
                                           out
```

• MPI_Cart_map

```
int MPI_Cart_map(
                                           */,
        MPI_Comm
                      comm
                                     in
                      ndims
        int
                                     in
                      dims[]
        int
                                     in
                      periods[]
        int
                                     in
        int*
                      newrank
                                           */)
```

• MPI_Cart_rank

• MPI_Cart_shift

• MPI_Cart_sub

```
int MPI_Cart_sub(

MPI_Comm comm /* in */,

int free_coords[] /* in */,

MPI_Comm* newcomm /* out */)
```

• MPI_Dims_create

• MPI_Graph_create

```
int MPI_Graph_create(
        MPI_Comm
                       old\_comm
                       nnodes
        int
                                         in
        int
                       index[]
                                         in
        int
                       edges[]
                                         in
        int
                       reoder
                                         in
        MPI_Comm
                       graph_comm
```

• MPI_Graphdims_get

```
int MPI_Graphdims_get(

MPI_Comm comm /* in */,

int* nnodes /* out */,

int* edges /* out */,
```

• MPI_Graph_get

```
int MPI_Graph_get(
           MPI_Comm
                        comm
                                        in
                        max_index
           int
                                        in
           int
                        max_edges
                                        in
                        index[]
           int
                                        out
                        edges[]
           int
                                        out

    MPI_Graph_map

   int MPI_Graph_map(
           MPI_Comm
                        comm
                                      in
           int
                        nnodes
                                      in
           int
                        index[]
                                      in
           int
                        edges[]
                                      in
           int*
                        newrank
                                      out
• MPI_Graph_neighbors
   int MPI_Graph_neighbors(
           MPI_Comm
                             comm
           int
                             rank
                                                in
           int
                             max_neighbors
                                                in
                             neighbors[]
           int
                                                out
• MPI_Graph_neighbors_count
   int MPI_Graph_neighbors_count(
                                                        */,
           MPI_Comm
                                   comm
                                                        */,
           int
                                   rank
                                                   in
           int*
                                   nneighbors
                                                   out
• MPI_Get_processor_name
   int MPI_Get_processor_name(
           char*
                                name
                                                 out
           int*
                                resultlen /*
                                            out
• MPI_Errhandler_create
   int MPI_Errhandler_create(
           MPI_Handler_function*
                                  function
                                                  in
           MPI_Errhandler*
                                  errhandler
                                                  out */)
• MPI_Errhandler_free
   int MPI_Errhandler_free(
           MPI_Errhandler*
                             errhandler /* in/out */)
• MPI_Errhandler_get
   int MPI_Errhandler_get(
```

comm

errhandler

out

MPI_Comm

MPI_Errhandler*

• MPI_Errhandler_set

• MPI_error_class

```
int MPI_Error_class(
    int errorcode /* in */,
    int* errorclass /* out */)
```

• MPI_Error_string

```
int MPI_Error_string(
    int errorcode /* in */,
    char* string /* out */,
    int* resultlen /* out */)
```

• MPI_Wtick

doube MPI_Wtick(void)

• MPI_Wtime

doube MPI_Wtime(void)

• MPI_Abort

```
doube MPI_Abort(

MPI_comm comm /* in */,

int error_code */ in */)
```

• MPI_Finalize

int MPI_Finalize(void)

• MPI_Init

• MPI_Initialized

```
int MPI_Initialized
( int^* \qquad flag \quad /^* \quad out \quad ^*/)
```

B.2.4 MPI Constants and Definitions

• Error Classes

MPI_SUCCESS

MPI_ERR_BUFFER

MPI_ERR_COUNT

MPI_ERR_TYPE

MPI_ERR_TAG

MPI_ERR_COMM

MPI_ERR_RANK

MPI_ERR_REQUEST

MPI_ERR_ROOT

MPI_ERR_GROUP

MPI_ERR_OP

MPI_ERR_TOPOLOGY

MPI_ERR_DIMS

MPI_ERR_ARG

MPI_ERR_UNKNOWN

MPI_ERR_TRUNCATE

MPI_ERR_OTHER

MPI_ERR_INTERN

MPI_PENDING

MPI_ERR_IN_STATUS

MPI_ERR_LASTCODE

• Assorted Constants

MPI_BOTTOM

MPI_PROC_NULL

MPI_ANY_SOURCE

MPI_ANY_TAG

MPI_UNDEFINED

MPI_BSEND_OVERHEAD

MPI_KEYVAL_INVALID

• Error Handling Specifiers

MPI_ERRORS_ARE_FATAL MPI_ERRORS_RETURN

• Maximum Sizes for Strings

MPI_MAX_PROCESSOR_NAME MPI_MAX_ERROR_STRING

• Basic Datatypes

MPI_CHAR

MPI_SHORT

MPI_INT

MPI_LONG

MPI_UNSIGNED_CHAR

MPI_UNSIGNED_SHORT

MPI_UNSIGNED

MPI_UNSIGNED_LONG

MPI_FLOAT

MPI_DOUBLE

MPI_LONG_DOUBLE

MPI_BYTE

MPI_PACKED

MPI_LONG_LONG_INT /* optional */

• Datatypes for Reduction Functions

MPI_FLOAT_INT

MPI_DOUBLE_INT

MPI_LONG_INT

MPI_2INT

MPI_SHORT_INT

MPI_LONG_DOUBLE_INT

• Datatypes for Building Derived Types

MPI_UB

MPI_LB

• Predefined Communicators

MPI_COMM_WORLD

MPI_COMM_SELF

• Results of Communicator and Group Comparisons

MPI_IDENT

MPI_CONGRUENT

MPI_SIMILAR

MPI_UNEQUAL

• Attrbute Keys for Implementation Information

MPI_TAG_UB

MPI_IO

MPI_HOST

MPI_WTIME_IS_GLOBAL

• Collective Reduction Operations

MPI_MAX

MPI_MIN

MPI_SUM

 $\mathrm{MPI} \underline{\hspace{0.1cm}} \mathrm{PROD}$

MPI_MAXLOC

MPI_MINLOC

MPI_BAND

MPI_BOR

MPI_BXOR

MPI_LAND

MPI_LOR

MPI_LXOR

• Null Handles

MPI_GROUP_NULL

MPI_COMM_NULL

MPI_DATATYPE_NULL

MPI_REQUEST_NULL

MPI_OP_NULL

MPI_ERHANDLER_NULL

• Empty Group

MPI_GROUP_EMPTY

• Topologies

MPI_GRAPH

MPI_CART

• Type Definitions

The following type definitions are in the file mpi.h.

• Opaque Types

MPI_Aint

MPI_Status

• Handles to Assorted Structures

MPI_Group

 MPI_Comm

MPI_Datatype

MPI_Request

MPI_Op

• Prototypes for User-Define Functions

```
typedef int MPI_Copy_function(
          \mathrm{MPI}\_\mathrm{Comm}
                                     oldcomm,
                                     keyval,
          int
          void*
                                     extra_arg,
          void*
                                     attribute_val_in,
                                     attribute_val_out,
          \mathrm{void}^*
                                     flag)
          int
typedef int MPI_Delete_function(
          MPI_Comm
                                      comm,
          int
                                      keyval,
          void*
                                      attribute\_val
          void*
                                      extra_arg)
typedef void MPI_Handler_function(
          MPI_Comm*
                                          comm,
          int*
                                          error_code,
                                          ...)
typedef void MPI_User_function(
          \mathrm{void}^*
                                      invec,
          void*
                                      inoutvec,
          int*
                                      len,
          {\rm MPI\_Datatype}^*
                                      datatype)
```

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